Structure of the Vesicular Stomatitis Virus L Protein in Complex with Its Phosphoprotein Cofactor

Highlights

- Cryo-EM structure of VSV L protein with bound P cofactor at 3.0 Å resolution
- P interacts with multiple domains of L to lock it in an initiation competent state
- Conserved tryptophan in priming loop supports transcription-initiating nucleotide
- L structure shows potential RNA exit channel for full-length replication product

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In Brief

Jenni et al. describe a 3.0 Å resolution cryo-EM structure of vesicular stomatitis virus L protein, bound with its P-protein cofactor, suggesting molecular features of RNA-synthesis initiation, transcript capping, and replication-product encapsidation.
Structure of the Vesicular Stomatitis Virus L Protein in Complex with Its Phosphoprotein Cofactor

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SUMMARY

The large (L) proteins of non-segmented, negative-strand RNA viruses are multifunctional enzymes that produce capped, methylated, and polyadenylated mRNA and replicate the viral genome. A phosphoprotein (P), required for efficient RNA-dependent RNA polymerization from the viral ribonucleoprotein (RNP) template, regulates the function and conformation of the L protein. We report the structure of vesicular stomatitis virus L in complex with its P cofactor determined by electron cryomicroscopy at 3.0 Å resolution, enabling us to visualize bound segments of P. The contacts of three P segments with multiple L domains show how P induces a closed, compact, initiation-competent conformation. Binding of P to L positions its N-terminal domain adjacent to a putative RNA exit channel for efficient encapsidation of newly synthesized genomes with the nucleoprotein and orients its C-terminal domain to interact with an RNP template. The model shows that a conserved tryptophan in the priming loop can support the initiating 5’ nucleotide.

INTRODUCTION

The large (L) protein encoded by the genomes of nonsegmented, negative-sense (NNS) RNA viruses carries out all the various catalytic steps associated with transcription and replication. A virally encoded phosphoprotein (P) is an essential cofactor, both for the incorporation of L into virions and for the regulation of replication and transcription. In addition to its RNA-dependent RNA polymerase (RdRp) activity, L caps and methylates the 5’ ends of transcripts. In the order of N, P, M, G, and L. The template for transcription is a full-length ribonucleoprotein (RNP)—that is, a genome-sense RNA fully coated with protein N. Each N subunit accommodates nine nucleotides of RNA in a groove along the waist of an elongated, two-lobe protein. Transcription of successive genes (with about 70% efficiency) initiates upon the termination and polyadenylation of the upstream transcript, produced by stuttering on a U7 sequence at the end of each gene (Iverson and Rose, 1981).

We described five years ago the structure of a VSV L-P complex from a cryo-electron microscopy (EM) reconstruction at 3.8 Å resolution (Liang et al., 2015). The multifunctional L protein—a single, 2109-amino-acid-long polypeptide chain—folds into three catalytic and two structural domains (Figure 1A; Data S1). The N-terminal RdRp domain and a capping (Cap) domain, which follows in primary sequence and has polynucleotidyl transferase (PRNTase) activity, form the core of the structure (Figure 1B). Negative-stain EM images had shown previously that in the absence of P, the three remaining domains—the connector domain (CD), the methyltransferase (MT), and the C-terminal domain (CTD)—have no fixed position with respect to the RdRp-Cap core structure, and the full molecular model from the cryo-EM showed particularly long linker segments between the Cap domain and CD and between the CD and MT (Liang et al., 2015; Rahmeh et al., 2010).

The polymerase catalytic site faces a cavity at the center of the RdRp domain (Liang et al., 2015). Channels between this cavity and the surface of the molecule allow entry and exit of the template RNA strand and entry of nucleoside triphosphates (NTPs). A loop from the Cap domain, projecting into the RdRp catalytic cavity, appears to be a priming loop to support the initiating nucleotide. Analogous loops are present in other viral RdRps that do not initiate on a polynucleotide primer (Lu et al., 2008; Tao et al., 2002). The priming loop must shift out of the way to allow elongation, as seen in the transition from initiation to elongation in the reovirus
polymerase (Tao et al., 2002). In VSV L, reconfiguration of the priming loop would allow the leading end of the elongating product strand to move into a cavity in the Cap domain that includes its active-site residues.

The capping reaction catalyzed by the L proteins of NNS RNA viruses proceeds through intermediates different from those of the mammalian-cell capping process (and different from the related and well-characterized capping mechanism of double-strand RNA viruses). Instead of an attack by the 5' end of the RNA on a guanosine monophosphate (GMP) covalently attached to a lysine side chain, the monophosphorylated 5' end of the transcript attaches covalently to a histidine side chain; resolution of this intermediate through attack by guanosine diphosphate (GDP) or guanosine triphosphate (GTP) results in a guanylated 5' terminus. The Cap domain is thus a PRNTase; 2'-O methylation and 7-N methylation, reactions carried out by the single active site on the MT domain, complete the modification.

Our previous L-P reconstruction resulted in a few regions of uncertain chain trace and did not clearly identify features corresponding to P (Liang et al., 2015). It was clear, however, that the binding of a fragment covering P residues 35–106, the sequence between the conserved N-terminal segment (PNTD) and the oligomerization domain (POD) (Figure 1A; Data S1), induced a "closed" structure of L in which the three terminal domains docked in well-defined positions onto the RdRp-Cap domain (Figure 1B). We interpreted the observed conformation as a preinitiation complex. Poor density in the region between the active sites of the RdRp and Cap domains prevented analysis of the coupling of transcription and capping.

We have now extended the resolution of the VSV L-P structure to 3.0 Å, from new images recorded on a Titan Krios microscope. The new reconstruction has allowed us to trace three segments of P bound to L and to correct a few small errors in the previous model. The molecular L-P interactions explain how P binding leads to a closed structure. Improved definition of several parts of the Cap domain and a clearer trace of the connections into and out of the connector domain have allowed us to identify a channel leading from the polymerase active site to a potential genome and anti-genome exit site at a position appropriate for acquisition of N, held close to that site by interaction with the N-terminal segment of P. Direct deposition of N onto the emerging RNA could afford prompt encapsidation and maximal protection. Better definition of the Cap domain active-site cavity also allows us to consider possible conformational changes that accompany formation of the covalent 5'-monophosphate-RNA intermediate and its transfer to GDP.

Figure 1. Structure of the VSV L Protein in Complex with the Phosphoprotein
(A) Linear domain organization of VSV L and P. Domain boundary numbers are shown. L-protein domains are colored as follows: RNA-dependent RNA polymerase domain (RdRp), cyan; capping domain (Cap), green; connector domain (CD), yellow; methyltransferase (MT), orange; C-terminal domain (CTD), red. Catalytic residues (CRs) are shown above. CRI–VI: conserved regions in L proteins of NNS RNA viruses. The L-protein scheme is adapted from Liang et al. (2015). P-protein N-terminal (PN), oligomerization (POD), and C-terminal (PCD) domains are in gray; its L-protein-binding domain (PL) is in magenta.
(B) Cryo-EM structure of the L-P complex determined at 3.0 Å resolution. Full-length L was incubated with a P fragment comprising residues 35–106. Domains colored as in (A). Dashed lines show flexible PL segments, connecting the three L-bound segments.
(C) L-protein-binding motif 1 of P shown as sticks (carbon, magenta; nitrogen, blue; oxygen, red). The density map (EMD-20614, B sharpened map) is shown as gray mesh around P (carving radius = 2.0 Å).
(D) L-protein-binding motif 2 of P, colored and with density map as shown in (C). Absence of density for almost all acidic amino acid side chains (Asp and Glu, indicated by asterisks) is characteristic of cryo-EM density maps.
(E) L-protein-binding motif 3 of P, colored and with density map shown as described in (C) and (D).
See also Figures S2 and S3 and Table S1.
RESULTS AND DISCUSSION

Structure of VSV L-P from Cryo-EM Reconstruction at 3.0 Å Resolution

We prepared the VSV L-P complex for cryo-EM imaging as described previously (Liang et al., 2015). We expressed full-length VSV L in insect cells and a P fragment (residues 35–106) in bacteria. We mixed L and P in vitro and purified the complex by gel filtration. We recorded images (Figures S1A and S1B) and carried out the image processing steps as described in the Method Details. The final map used for the modeling and interpretation reported here (Figure 1C) had an overall resolution of 3.0 Å, as determined by the spatial frequency at which the Fourier shell correlation (FSC) between half maps dropped below 0.143 (Figure S1C). The data quality allowed us not only to improve the overall resolution of the reconstruction, but also to classify out a more homogeneous set of particle images, yielding a density map in which all domains were equally well resolved.

We could place our previous model (PDB: 5A22) into the density without any major adjustments. We traced connections into and out of the connector domain more clearly than with the previous, lower-resolution reconstruction; corrected a local chain trace error associated with these connections; corrected a few errors in assigning sequence register to segments poorly defined in the previous map; and adjusted many side-chain rotamers, which were generally well defined in the new map. Data collection, model refinement, and validation statistics are in Table S1. The positions of various differences between the updated and the original coordinates are shown in Figures S2 and S3.

Interactions of the P Subunit with L

The most important new feature was interpretable density for parts of P (Figure S4), enabling us to model three ordered segments from residues 49 to 56, 82 to 89, and 94 to 105 (Figures 2A and 2B). The first segment binds in a shallow pocket on the outward-facing surface of the CTD. The interactions are mostly hydrophobic around a conserved tyrosine residue P Tyr53 that also hydrogen bonds to the side chain of L Asp1981 (Figure 2C).

P then wraps around the CTD with a stretch of flexible residues that we can detect in low-pass-filtered density maps (Figure S4B). The second segment binds between L Arg1419 and L Tyr1438. A conserved amino nitrogen or carbonyl oxygen. These interactions, which promise the end initiation, but it appears not to affect the internal initiation of transcripts, and it does not prevent capping (Ogino et al., 2019). Mutation of the homologous tryptophan in RABV L also compromises the end initiation, but it appears not to affect the internal initiation of transcripts, and it does not prevent capping (Ogino et al., 2019).

Further polymerization requires retraction of the priming loop or, potentially, displacement of the entire Cap domain. The HMPV (Pan et al., 2019) and RSV (Gilman et al., 2019) polymerase structures show a fully retracted priming loop with no
substantial displacement of the rest of the Cap domain. The priming loops of those two enzymes and the catalytic cavities of their Cap domains are similar, and we can show by superposition that the VSV priming loop can also retract without any major rearrangements in the rest of the Cap domain (Figure 4B). Moreover, when we model the VSV L priming loop based on the HMPV L protein structure, we find that a twisting tunnel connects the RdRp catalytic site with an opening where the CTD, RdRp, and MT all meet (Figure 4C). We suggest that this tunnel might be the exit pathway for uncapped products (i.e., the full-length antigenome and genome during replication and the uncapped leader during transcription), which bypass the active sites of the Cap and MT domains.

The N-terminal segment of P (PNTD) binds a single subunit of N, forming the so-called N0-P complex. The likely function of this complex is to deliver N to the nascent genome and antigenome as they emerge from the polymerase. About 15 residues connect the N-binding element (residues 6–34) with the segment of P docked onto the outer face of the CTD. The N0 passenger could therefore lie close to the proposed exit site for uncapped products. A possible explanation for the defect in Y53 mutants is the release of the docked N-terminal region of P from its site.
on the CTD, causing inefficient delivery of N to the replication products. This defect would not affect transcription.

At the other end of the P polypeptide chain, a small C-terminal domain (P\textsubscript{CTD}) interacts with the RNP. The binding site is at the C-terminal lobe of the N proteins (Green and Luo, 2009). Together with a presumptive interaction of the RNP with L at the template entry site, the contact with P could facilitate the passage of the template strand through the enzyme. We have proposed previously that the subunits of N (probably three) that must separate from the RNA template as it threads through the polymerase do not dissociate from each other and instead rebind the RNA as polymerization proceeds. The N-N interaction creates a chain-like array, with an N-terminal arm and a C-proximal loop contacting the two neighboring subunits in the chain (Ge et al., 2010; Green et al., 2006). Interaction with the C terminus of L-bound P could then ensure the correct reassociation of this putative chain of N subunits with exiting RNA (see Figure 3G).

Transcription of all products except the uncapped leader requires a different sequence of events. One possibility is that the immediately upstream transcript leaves, and the priming loop snaps back into the position seen in our current structure. Internal initiation does not appear to depend on T\textsubscript{Trp1167}; however, one possibility is that the 3’ end of the upstream gene, supports the initiating NTP instead. The interaction would need to account for the intergenic GA sequence on the template. Further polymerization would then displace the upstream transcript (Shuman, 1997).

Acquisition of the guanylate cap occurs when the transcript length has reached at least 31 nucleotides (Tekes et al., 2011). If we suppose that the formation of the covalent intermediate with T\textsubscript{His1227} occurs before the addition of the 31st nucleotide to the transcript, then approximately 30 nucleotides of transcript would need to fit into the combined active-site cavities of the RdRp and Cap domain. The volume of the Cap active-site cavity, marked as the transfer of N from N\textsuperscript{2}-P to emerging RNA. The C-terminal domain of P (P\textsubscript{CTD}) interacts with the RNP; it may guide the polymerase along the template. P dimerization through the oligomerization domain (P\textsubscript{OD}) would lead to additional RNP and N\textsuperscript{2}-P interactions.

See also Data S2.
with an asterisk in Figure 4C, is $1.2 \times 10^4 \text{ Å}^3$. Assuming a volume occupied per Dalton (Da) of RNA ($V_m$) of 2.1 to 4.6 Å$^3$ and an average mass of 340 Da per nucleotide (Speir and Johnson, 2012), the interior of the Cap domain could accommodate between 8 and 17 transcript nucleotides, in addition to the 10 nucleotides still base paired with the template in the RdRp catalytic cavity. These relatively generous estimates suggest that the addition of the guanylate to the 5' end of the transcript requires an opening of the multi-domain assembly from the closed conformation present in our structure. A plausible transition would be the release of the CD, MT, and CTD from their docked positions, as indeed occurs in the absence of P. If that transition also enabled a guanine nucleotide to access the RNA-protein covalent linkage, release of the capped 5' end would allow it to diffuse efficiently into the MT active site, since both the transcript (still anchored to the RdRp at its growing, 3' end) and the MT (linked to RdRp-Cap through the CD) cannot move far from each other.

The VSV L-P structure described here illustrates how the VSV P cofactor regulates L function by locking the domain organization into a single, pre-initiation configuration. It allows an explicit comparison with three other NNS viral L-P complexes shown in Figure S4C. In the VSV and RABV L-P complexes, an extended segment of P contacts several domains, thereby fixing the relative positions of the RdRp-Cap and CD-MT-CTD modules; the oligomerization domain of P does not participate in the contact with L. In the pneumovirus complexes, the oligomerization domain of P interacts directly with L, and the extended segment binds only the RdRp, allowing the three C-terminal domains to move freely about the linkers that connect them. A more complete analysis will require structures with RNP templates, such as transcribing intermediates, and studies of viral L proteins in complex with inhibitors.

STAR+METHODS
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.celrep.2019.12.024.
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AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES


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# STAR METHODS

## KEY RESOURCES TABLE

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Further information and requests for reagents may be directed to Stephen C. Harrison (harrison@crystal.harvard.edu). All unique reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement. Requests for reagents can be addressed to spjwhelan@wustl.edu.

Expression plasmids contained cDNA sequences from vesicular stomatitis virus (VSV) Indiana strain. Recombinant VSV-L protein was overexpressed in Sf21 insect cells. VSV-P protein (residues 35–106) was overexpressed in Escherichia coli Rosetta BL21 (DE3) cells (Novagen) (Liang et al., 2015).

Cryo-EM Structure Determination
The VSV L-P protein complex was prepared for structural analysis as previously described (Liang et al., 2015). Briefly, 3 μL of protein at ~0.35 mg/mL were applied to a C-flat grid (CF-1.2/1.3-4C, Protochips) that had been glow discharged at 40 mA for 30 s. Grids were plunge-frozen with an Vitrobot Mark II (FEI Company) after blotting for 3 s at 4°C and ~85% relative humidity. 3307 movies of vitrified protein solution were collected using SerialEM (Mastronarde, 2005) on a Titan Krios electron microscope (ThermoFisher) at 300 kV and 59,000 nominal magnification, on a K2 Summit detector (Gatan, Inc.) operated in counting mode, resulting in a calibrated pixel size of 0.485 Å. Each movie contained 50 frames, collected in a 10 s exposure at 2 electrons/Å²/frame. Most micrographs showed well-dispersed particles (Figure S1A). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015). CTF fringes on many micrographs were fitted to better than 3.0 Å resolution (Figure S1B). Movies were aligned and exposure-filtered using Unblur (Grant and Grigorieff, 2015). 3235 movies were selected for further processing based on the frame motion traces, excluding movies with traces suggesting large and zig-zag motion. The contrast transfer function (CTF) was determined using CTFFIND4 (Rohou and Grigorieff, 2015).
class at 4.0 Å resolution resulted in the final 3.0 Å reconstruction. The reconstruction was sharpened using cisTEM’s sharpening tool, amplifying signal at high resolution with a B factor-equivalent of 75 Å².

**Model Building and Refinement**

We used the program O (Jones et al., 1991) to adjust our previously determined VSV L protein structure (PDB-ID 5A22, determined at 3.8 Å resolution) (Liang et al., 2015) to fit the 3.0 Å resolution cryo-EM map. We modeled the P protein residues de novo. We refined coordinates and B factors with phenix.real_space_refine (version 1.16-3549) (Afonine et al., 2018). In addition to the standard chemical and grouped B factor restraints, we used secondary structure, rotamer, and Ramachandran restraints (weight = 1.0, nonbonded_weight = 500, rama_weight = 6.0). We used phenix.mtriage (Afonine, 2017) and MolProbity (Chen et al., 2010) to calculate the validation results shown in Figure S1C and Table S1. The final VSV L-P model comprises L protein residues 35–1211, 1217–1332, 1339–1590, and 1596–2109; P protein residues 49–56, 82–89, and 94–105; and two zinc atoms. Improved model regions between updated (PDB-ID 6U1X) and original (PDB-ID 5A22) coordinates (Figures S2 and S3) are as follows. RdRp residues: 61–64 (loop remodeled, sequence register shift of 60–64), 414–430 (sequence register shift of 414–430), 587–593 (loop remodeled, sequence register shift of 586–593). Cap residues: 958 (loop adjusted), 1035–1046 (sequence register shift of 1035–1046), 1072–1075 (loop remodeled), 1100 (loop adjusted), 1122–1125 (loop remodeled, sequence register shift of 1124–1126), 1164–1170 (loop remodeled, sequence register shift of 1161–1170), 1195–1232 (sequence register shift of 1195–1227), 1262 (loop adjusted), 1314 (loop adjusted), 1321–1332 (region adjusted). CD residues: 1380–1395 (region remodeled, sequence register shift of 1379–1396), 1498 (loop adjusted), 1535–1539 (loop remodeled, sequence register shift of 1535–1538). MT residues: 1718 (loop adjusted), 1806–1851 (sequence register shift of 1805–1841). CTD residues: 1952–1955 (loop remodeled, sequence register shift of 1950–1955), 2006–2008 (shift in β hairpin tip), 2066 (loop remodeled, 2086–2094 (loop remodeled, sequence register shift of 2084–2094), 2109 (Cz in main chain density instead side chain density of 2108).

**Projection Matching of Negative-Stain Class Averages**

We extracted 20 negative-stain class averages of L-P dimers from a screenshot of a previously published figure (Figure S4C in Rahmeh et al., 2010) using RELION-3 (Zivanov et al., 2018). From those dimer images we manually extracted 40 subparticles, one of each protomer, and masked them with a soft spherical mask. We obtained rotation angles for reference projections of our reconstruction with healpy (http://healpy.readthedocs.io), the Python implementation of the HEALPix library of routines for equal area sampling of a sphere (Górski et al., 2005). Because the L-P complex has no symmetry and we did not allow mirroring of reference projections in the subsequent alignment step, we sampled angles on the entire sphere. We aligned the subparticle images to the reference projections with e2simmm.py from EMAN2 (Bell et al., 2016). Alignments were scored by calculating a cross-correlation coefficient (CCC). Because information in the negative-stain class averages is restricted to low resolution, reference projections from very different angles resulted in similar CCCs, and the “correct” solution was not always the top scoring projection. We therefore looked at the top five alignment peaks, which we adjusted with the CCP4 program PEAKMAX (Winn et al., 2011), and then manually selected for further analysis the one whose reference projections most closely matched the particle image as judged by eye (Data S2). To calculate the distance between the P protein C termini of the two promoters, we transformed the model coordinates according to the determined alignment parameters. The analysis results for all class averages can be found in Data S2.

**Functional Assays**

**Cells**

BSR-T7 cells (a kind gift from K. Conzelmann) (Buchholz et al., 1999) were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Cominco Inc., 10-013-CV) containing 10% fetal bovine serum (FBS; Tissue Culture Biologicals, TCB 101) at 37°C and 5% CO₂.

**Plasmids**

For mammalian expression, plasmids expressing Y53 mutants were generated by site-directed mutagenesis using the Q5 polymerase-nese (New England Biolabs) on the parental P expressing plasmid (pMB-NS) (Pattnaik and Wertz, 1990) with dedicated primers (for Y53A: TGGGCCCTTGCCTTTCAGGCACG and GTATGCTTTTACCACT, for Y53D: TAGGCCCTTGGTTCGCGAC and GTATGCTTTTACCACT, for Y53F: CATACTAGGCCCTTTTTTTTAGAGAC and AGAGGGCTATATGCTG, for deltaY53: TTTCAGGCAGCGATGATC and AGAGGGCTATATGCTG). For bacterial expression and purification, fusion proteins made of the HRV_3C-P fragment was amplified from pET16b-P (Rahmeh et al., 2012) using GAAGTACTGTTCCAGGGTCCTATGGATAATCT and CTGGAASGGTTTTACCATATTTT, for Y53A: TAGGCCCTTGGTTCGCGAC and GTATGCTTTTACCACT, for deltaY53: TTTCAGGCAGCGATGATC and AGAGGGCTATATGCTG. For bacterial expression and purification, fusion proteins made of the HRV_3C-P fragment was amplified from pET16b-P (Rahmeh et al., 2012) using GAAGTACTGTTCCAGGGTCCTATTTT, for Y53A: TAGGCCCTTGGTTCGCGAC and GTATGCTTTTACCACT, for deltaY53: TTTCAGGCAGCGATGATC and AGAGGGCTATATGCTG. DeltaY53 mutations were introduced in pET16b-MBP-3Cc-P by site-directed mutagenesis with primers described above. To rescue viruses expressing P_Y53 mutants, Y53 mutations were introduced into pVSV1(+)-eGFP backbone (Whelan et al., 2000).
Protein purification

L and P proteins were purified as previously described (Rahme et al., 2012). MBP-P was purified from Rosetta cells (EMD Millipore) after induction with 0.3 mM IPTG overnight at 18°C. Cells were then pelleted and resuspended with 5 mL cold lysis buffer per gram of pellet (50 mM Hepes (pH 7.4), 100 mM NaCl, 5% glycerol, 5 mM imidazole, 5 mM 2-mercaptoethanol and 1X protease inhibitor cocktail [CompleteTM, Roche Cat. # 4693116001]). After sonication, 40 mL cell lysate were clarified by centrifugation at 20,000 x g for 20 min, and the supernatant was incubated with 2 mL of Ni-NTA beads (QIAGEN) overnight at 4°C. Beads were washed three times with lysis buffer (without protease inhibitor), and MBP-P proteins were eluted in 5 mL elution buffer (50 mM Hepes (pH 7.4), 100 mM NaCl, 5% glycerol, 5 mM imidazole and 5 mM 2-mercaptoethanol). The eluate was concentrated with a centrifugal filter (Amicon) and passed through a Superdex S200 size exclusion chromatography column (GE Healthcare) in storage buffer (20 mM Hepes (pH 7.4), 100 mM NaCl, 5% glycerol, and 5 mM 2-mercaptoethanol). MBP-P proteins eluted in a single peak and were kept at –80°C in storage buffer.

Virus

Attempts to rescue viruses expressing P_Y53 mutants, were performed as previously described (Whelan et al., 1995) using pSVV1 (+)-eGFP-P_Y53x plasmids. VSV-eGFP-P_Y53F was the only mutant virus forming plaques. It was amplified and titered on BSR-T7 cells.

N-RNA purification

Nucleocapsid template was purified essentially as described previously (Ongrádi et al., 1985). To limit background activity due to residual P bound to N-RNA, nucleocapsids were extracted from a recombinant VSV expressing a monomeric P protein whose oligomerization domain has been deleted (VSV-P<sub>Δolig</sub>). Briefly, 10 mg of gradient purified virions were disrupted on ice for 1 h in 10 mL of virion disruption buffer (VDB: 20 mM Tris-HCl (pH 7.4), 0.1% Triton X-100, 5% glycerol, 1 mM EDTA, 2 mM dithioerythritol and 600 mM NH<sub>4</sub>Cl). Lysozyme virions were centrifuged at 240,000 x g for 3.5 h through a glycerol step gradient of 0.25 mL each of 40, 45, and 50% glycerol in VDB. Pellets were resuspended overnight in 0.5 mL of 10 mM Tris-HCl (pH 7.4), 100 mM NaCl, 1 mM EDTA and 1 mM DTT, and disrupted again on ice for 1 h after mixing with 0.5 mL of 2X high salt VDB (1X HSVDB: 20 mM Tris-HCl (pH 7.4), 0.1% Triton X-100, 5% glycerol, 1 mM EDTA, 2 mM dithioerythritol and 1.5 M NH<sub>4</sub>Cl). Nucleocapsids were isolated by banding on a CsCl gradient. In a 5 mL tube, 1 mL of disrupted virions was added at the top of a step gradient of 2 mL of each 20 and 40% (w/v) CsCl in HSVDB. After centrifugation at 190,000 x g for 3 h, N-RNA were recovered by side puncture and diluted fivefold in NaCl-Tris-EDTA-DTT buffer (NTE: 10 mM Tris-HCl (pH 7.4), 100 mM NaCl, 1 mM EDTA and 2 mM DTT). N-RNA was then centrifuged through a 0.5 mL cushion of 50% glycerol in NTE buffer, resuspended overnight in NTE buffer and stored in aliquots at –80°C.

Production of minigenome particles

Production of transcription competent minigenome particles was as described previously with minor modifications (Pattnaik and Wertz, 1990; Wertz et al., 1994). BSR-T7 cells were plated in a 60mm dish and infected the next day with a vaccinia virus expressing T7 polymerase (vTF7-3) (Fuerst et al., 1986) at multiplicity of infection (MOI) 3 for 1 h in Dulbecco’s Phosphate Buffered Saline liquid (DBPS; Sigma Cat# 58300C). Cells were then transfected using Lipofectamine 2000 with plasmids expressing N (5.5 μg), P (1.6 μg), M (1.25 μg), G (2.5 μg), L (0.5 μg) and a minigenome containing an eGFP reporter gene (7.5 μg). Four hours later, the medium was replaced with 3 mL DMEM containing 2% FBS. The cell supernatant containing the DI particles (DI stock) was harvested 2 days after transfection.

Cell-based gene expression assay

BSR-T7 cells were plated in a 96-well plate and infected the next day with a vaccinia virus expressing T7 polymerase (vTF7-3) at MOI 3 for 1 h in DBPS. Cells were then transfected with plasmids expressing N (91 ng), P (26 ng) and L (7.5 ng) using Lipofectamine 2000. Four hours later, cells in each well were infected with 3 μL DI stock in 30 μL DMEM for 1 h. Two days after transfection, GFP signal was measured using a Typhoon FLA 9500 scanner (GE Healthcare).

In vitro transcription assays

RNA synthesis assay on naked RNA was described previously (Morin et al., 2012). Briefly, purified L (100 nM) was incubated with purified P or MBP-P (250 nM) and a 19nt-long RNA corresponding to the first nucleotides of the genomic promoter (1.25 μg). Reactions were performed in 10 μL at 30°C for 5 h in presence of radiolabeled GTP (32P-γGTP, PerkinElmer) and analyzed on a 20% polyacrylamide/7 M urea gel. The gel was exposed overnight to a phosphor screen (GE Healthcare).
Superposition of Atomic Models
To compare the VSV L-P structure to other viral polymerases, we used the program LSQMAN (Kleywegt and Jones, 1997) for superposition, except for the structure of the reovirus λ3 polymerase initiation complex, for which we also used the program O (Jones et al., 1991) to focus the alignment on the RdRp active site. Superposition statistics are summarized in Table S2.

Cavity Volume Calculation
We used the program VOIDOO (Kleywegt and Jones, 1994) to calculate the volume of the Cap active site cavity (Figure 4C, marked by an asterisk). The radius of the probe sphere was 1.4 Å. The calculated volume corresponds to the probe-occupied volume. We closed off the cavity of interest at the constrictions leading to the RdRp and MT active sites cavities, respectively, with dummy atoms.

Figure Preparation
We used MAFFT to calculate the amino acid multiple sequence alignments (Katoh et al., 2002). We prepared the figures with PyMol (The PyMOL Molecular Graphics System, Version 2.1 Schrödinger, LLC), Pov-Ray (www.povray.org), matplotlib (Hunter, 2007), and ESPript (Robert and Gouet, 2014). Secondary structure elements were assigned using DSSP (Kabsch and Sander, 1983).

QUANTIFICATION AND STATISTICAL ANALYSIS
In the plot of Figure 3B, the box extends from the lower to upper quartile values of the data. The line is at the median. The whiskers extending from the box show the range of the data. In Figure 3D, data are represented as mean ± SEM.

DATA AND CODE AVAILABILITY
The accession number for the EM 3-D reconstruction maps reported here is EMDB: 20614; the accession number for the atomic coordinates is PDB: 6U1X.
Supplemental Information

Structure of the Vesicular Stomatitis Virus L Protein in Complex with Its Phosphoprotein Cofactor

Simon Jenni, Louis-Marie Bloyet, Ruben Diaz-Avalos, Bo Liang, Sean P.J. Whelan, Nikolaus Grigorieff, and Stephen C. Harrison
Figure S1. Fourier Shell Correlation (FSC). Related to STAR Methods.

(A) Micrograph of vitrified VSV L-P, showing well-dispersed particles. The scale bar corresponds to 100 Å.

(B) Contrast transfer function (CTF) fringes calculated from the movie frames of the micrograph shown in A, with good match of the fitted pattern.

(C) FSC curves for the VSV L-P cryo-EM reconstruction and refined model calculated with phenix.mtriage (Afonine, 2017). Correlations for the two half maps are shown in blue after applying a loose spherical mask (dashed line) and a tight mask based on the model (solid line), respectively. The correlation between the refined model and final map is shown as red solid line.
**Figure S2.** Cα Distances between PDB-ID 6U1X and PDB-ID 5A22. Related to Figure 1 and STAR Methods.

The plot shows the Cα distances for each of the L protein domains after superposing the 3.0 Å resolution structure reported here (PDB-ID 6U1X) onto our previously determined structure at 3.8 Å resolution (PDB-ID 5A22). Domains are colored as in Figure 1.
Figure S3. Cryo-EM Density of Improved Model Regions of the VSV L-P Complex. Related to Figure 1 and STAR Methods.
The density map (EMD-20614, B sharpened map) is shown as gray mesh. Carbon atoms of the protein domains are colored as in Figure 1. Nitrogen atoms are blue; oxygen, red; sulfur, orange. Acidic amino acid side chains (Asp and Glu) with characteristic absence of density in the cryo-EM reconstruction are labeled with asterisks.
Figure S4. Cryo-EM Density of the Phosphoprotein and Comparison of Phosphoprotein-Binding to L Proteins from Different Viruses. Related to Figure 2.

(A) Multiple sequence alignment as shown and described in Figure 2A.

(B) A low-pass filtered map to 5 Å resolution of the VSV L-P reconstruction shows additional density for some less well ordered segments of the P protein (highlighted with arrows), corresponding to residues connecting the L protein-binding motifs. These residues are not above reasonable contour in the high-resolution map, and we therefore did not include them in the model of the complex.

(C) Structures were superimposed as described in the Methods section. The polymerase structures are from vesicular stomatitis virus (VSV), rabies virus (RABV), respiratory syncytial virus (RSV), and human metapneumovirus (HMPV). Protein Data Bank (PDB) accession identifiers for the structures are given. See Table S2 for details of the structures and superpositions. The P protein is colored in magenta.
Table S1. Cryo-EM Data Collection and Model Statistics. Related to STAR Methods.

<table>
<thead>
<tr>
<th>VSV L-P</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMD-20614</td>
</tr>
<tr>
<td>PDB-ID 6U1X</td>
</tr>
</tbody>
</table>

Indiana strain

**Data collection**
- Electron microscope: Titan Krios
- Magnification: 103092
- Voltage (kV): 300
- Defocus range (µm): 0.5–2.0
- Pixel size (Å): 0.97

**Reconstruction**
- Number of images: 83979
- Box size (pixels): 400
- Symmetry imposed: $C_1$
- Map resolution (Å): 3.0
- Estimated $B$ factor (Å²): 75

**Model statistics**
- Refinement resolution (Å): 3.0
- CC (mask): 0.82
- Non-hydrogen atoms: 16761
- Protein residues: 2059
- Mean $B$ factor (Å²): 70.7
- R.m.s. deviations
  - Bond lengths (Å): 0.004
  - Bond angles (°): 0.548
- Validation
  - MolProbity clash score: 3.7
  - Poor rotamers (%): 0.1
- Ramachandran plot
  - Favored (%): 99.7
  - Allowed (%): 0.3
  - Disallowed (%): 0.0

---

*a* Approximate range of underfocus.

*b* Resolution where Fourier shell correlation (FSC) between half-maps drops below 0.143 after applying a loose spherical shell mask and scaling of the FSC to estimate what it would be for a tight mask (Grant, T., Rohou, A., and Grigorieff, N. (2018). *cis*TEM, user-friendly software for single-particle image processing. Elife 7).

*c* The map was sharpened using *cis*TEM, with simple flattening of the power spectrum. Applying a sharpening $B$ factor of -75 Å² to the unsharpened map would approximately reproduce the *cis*TEM-sharpened map.

*d* Ramachandran restraints were used in the refinement target.
## Table S2. Superposition Statistics of Viral Polymerase Structures. Related to Figure 4 and STAR Methods.

<table>
<thead>
<tr>
<th>Structure</th>
<th>PDB-ID</th>
<th>Program</th>
<th>Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>VSV L protein</td>
<td>6U1X</td>
<td>N.A.</td>
<td>N.A.</td>
</tr>
<tr>
<td>Initiation-competent state</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reovirus λ3 polymerase</td>
<td>1N1H</td>
<td>LSQMAN</td>
<td>r.m.s. fit = 2.278 Å with 411 Cα atoms</td>
</tr>
<tr>
<td>Initiation complex</td>
<td></td>
<td>a</td>
<td></td>
</tr>
<tr>
<td>Elongation complex</td>
<td>1N35</td>
<td>LSQMAN</td>
<td>r.m.s. fit = 2.263 Å with 242 Cα atoms</td>
</tr>
<tr>
<td>Rotavirus VP1</td>
<td>6OJ6</td>
<td>LSQMAN</td>
<td>r.m.s. fit = 2.134 Å with 381 Cα atoms</td>
</tr>
<tr>
<td>Elongation complex</td>
<td></td>
<td>a</td>
<td></td>
</tr>
<tr>
<td>Rabies virus L protein</td>
<td>6UEB</td>
<td>LSQMAN</td>
<td>r.m.s. fit = 1.752 Å with 1729 Cα atoms</td>
</tr>
<tr>
<td>Initiation-competent state</td>
<td></td>
<td>a</td>
<td></td>
</tr>
<tr>
<td>RSV L protein</td>
<td>6PZK</td>
<td>LSQMAN</td>
<td>r.m.s. fit = 1.946 Å with 884 Cα atoms</td>
</tr>
<tr>
<td>Initiation-competent state</td>
<td></td>
<td>a</td>
<td></td>
</tr>
<tr>
<td>HMPV L protein</td>
<td>6U5O</td>
<td>LSQMAN</td>
<td>r.m.s. fit = 1.971 Å with 872 Cα atoms</td>
</tr>
<tr>
<td>Initiation-competent state</td>
<td></td>
<td>a</td>
<td></td>
</tr>
</tbody>
</table>

*a We obtained global alignments in LSQMAN with these commands:
  fast m1 A m2 A 50 25 100 ! initial brute-force structural alignment on a coarse grid
  im m1 * m2 *                       ! intermediate-fit
  dp m1 a m2 a sq 3.5 10           ! final fit with Needleman–Wunsch algorithm
*b We obtained the alignment in O with these commands:
  lsq_exp                               ! initial superposition of active site residues (VSV L708–L719)
  lsq_imp                               ! improvement of the alignment by considering the whole structures