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PII: S1877-959X(18)30345-5

DOI: https://doi.org/10.1016/j.ttbdis.2018.11.004

Reference: TTBDIS 1131

To appear in:

Received date: 16 August 2018 Revised date: 14 October 2018 Accepted date: 8 November 2018

Please cite this article as: Safonova MV, Shchelkanov MY, Khafizov K, Matsvay AD, Ayginin AA, Dolgova AS, Shchelkanov EM, Pimkina EV, Speranskaya AS, Galkina IV, Dedkov VG, Sequencing and genetic characterization of two strains Paramushir virus obtained from the Tyuleniy Island in the Okhotsk Sea (2015), *Ticks and Tick-borne Diseases* (2018), https://doi.org/10.1016/j.ttbdis.2018.11.004

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Sequencing and genetic characterization of two strains Paramushir virus

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Abstract

Paramushir virus belongs to Sakhalin virus genogroup within Orthonairovirus genus and is one of

the poorly studied viruses with unknown pathogenicity. At the moment, only one nearly complete

sequence of Paramushir virus genome, isolated in 1972, is available. Two new strains of PARV

were isolated in 2015 from a sample collected at the Tyuleniy Island in the Okhotsk Sea and

sequenced using a combination of high throughput sequencing and specific multiplex PCR. Both

strains are closely related to the early sequenced PARV strain LEIV-1149K. The signs of

intersegment reassortment and probable recombination were revealed, which point to a high

variability potential of Paramushir virus and may lead to the formation of strains with novel

properties, different from those of the predecessors.

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The new data regarding Paramushir virus can promote a better understanding of the diversity and relations within *Orthonairovirus* genus and help define intragenic demarcation criteria, which have not yet been established.

Key words: broad-range PCR, high throughput sequencing, Paramushir virus, orthonairoviruses, Russia

#### Introduction

At present, Orthonairovirus genus (formerly Nairovirus genus) contains 34 species; of them, 22 have not yet been classified (ICTV, 2016). All members of the Orthonairovirus genus have a negative-sense tripartite RNA genome (small [S], medium [M], and large [L] segments) with a high genetic diversity. Some of the orthonairoviruses were isolated from Eurasia, including the Crimean-Congo hemorrhagic fever virus (CCHFV), which is the causative agent of severe fever illness in humans (Deyde et al., 2006). Despite this fact, the information regarding biodiversity, spreading, and epidemiological potential of orthonairoviruses (excluding CCHFV, due to its medical significance) is insufficient. Paramushir virus (PARV) is one of the orthonairoviruses, for which the taxonomy has not yet been defined. After Dr. D. K. Lvov found two strains of PARV in 1972, no new strain has been identified until now (Lvov et al., 1976). Those strains (LEIV-2238 and LEIV-1149K) were isolated from *Ixodes signatus* ticks. The ticks were collected from seabird nests at Paramushir Island (the Kuril Islands in the Okhotsk Sea) (Figure 1a, c). At the moment, PARV, along with Avalon virus (AVAV), Tillamook virus (TILV), Sakhalin virus (SAKV), Taggert virus (TAGV), and Clo Mor virus (CMV) belong to the Sakhalin genogroup within Orthonairovirus genus (Walker et al., 2016). A partial sequence of the PARV L gene, strain LEIV-2238, is available in the NCBI GenBank under accession no KF801657 from March 2014, whereas the whole CDS sequence of PARV strain LEIV-1149K was submitted in 2017 and is now available in the NCBI GenBank under the accession nos. KP792717 – KP792719.

In this work, we describe two new strains of Paramushir orthonairovirus, which were isolated in Tyuleniy Island, located in the Far Eastern part of Russia in the Okhotsk Sea (see Figure 1a,b). Both strains were isolated in 2015 from *Ixodes uriae* White, 1852 (Acari: Ixodidae) ticks, and this was the first case of isolation of PARV after 43 years of the PARV discovery in the Far East of Russia. Tyuleniy Island is known to be the source of many viruses isolated from *I. uriae* ticks. These viruses include Tyuleniy virus (TYUV), belonging to *Flavivirus* genus in the *Flaviviridae* family (Lvov et al., 2001, 2014a, 2015); Sakhalin virus, (SAKV), belonging to *Orthonairovirus* genus in the *Nairoviridae* family (Lvov et al., 2001, 2014b, 2015); Zaliv Terpeniya virus (ZTV), Komandory virus (KOMV) and Rukutama virus (RUKV), which belong to *Phlebovirus* genus in the

Phenuiviridae family (Lvov et al., 2001, 2014b, 2014c, 2015, Alkhovsky et al., 2013); Okhotskiy virus (OKHV) and Aniva virus (ANIV), which belong to Orbivirus genus within the *Reoviridae* family (Lvov et al., 2001, 2014d, 2015). Thereby, this island is of great interest to the scientists who study the biodiversity and ecology of arboviruses. However, due to a large distance from the mainland and harsh climate, the accessibility to this area is limited. Nevertheless, in recent years, the interest in studying the island territories of the far Eastern part of Russia has increased, and some expeditions have been organized to Tyuleniy Island, including the one in August 2015 after a 25-year break. (Shchelkanov et al., 2017). In the course of the expedition, two strains of PARV were isolated, then sequenced using high throughput sequencing techniques, and subsequently characterized.

#### 2. Materials and methods

# Sample preparation and RNA extraction

During an arbovirus surveillance and control program, *I. uriae* ticks were collected from the nests of birds located in Tyuleniy Island (48°29′ N., 144° 38′ E) in the Okhotsk Sea near Sakhalin island (Figures 1a, 1b, 1c).

The identity and sex of the ticks were determined based on the morphology following earlier recommendations (Filippova, 1977). The ticks were classified according to species, sex, and developmental stages, and later processed in groups of 10 (immature ticks) and 25 individuals (adults). All the ticks were stored alive in a wet chamber until separation. The grouped tick samples (n=30) were homogenized using Tissue Lyser LT (Qiagen, Germany) for 5 min in 100 µL of phosphate buffered saline (pH 7.0, Dako, Denmark) before extraction, centrifuged at 10,000 g for 10 min, and the supernatant was used for nucleic acid extraction.

Total nucleic acids were extracted and purified using the RIBO-prep DNA/RNA extraction kit (k2-9-Et-100CE, AmpliSens, Russia), according to the recommendations of the manufacturer. DNA/RNA was eluted with 50  $\mu$ L of the elution buffer (AmpliSens, Russia) and stored at -70 °C until evaluation.

## Broad-range reverse transcriptase-polymerase chain reaction (RT-PCR)

Broad-range RT-PCR was performed using a set of broadly reactive degenerate oligonucleotides designed to target each viral species that could be transmitted by ticks within the following genera: *Flavivirus*, *Orthonairovirus*, *Phlebovirus*, *Orthobunyavirus*, *Orthoreovirus*, and *Orbivirus* (Table 1).

Reverse transcription and touch-down amplification were performed using the SuperScript III One-Step RT-PCR System with Platinum Taq DNA Polymerase (Invitrogen, USA) in a reaction mixture containing 2  $\mu$ L of RNA, 0.4  $\mu$ M sense primer, 0.4  $\mu$ M antisense primer, 12.5  $\mu$ L of 2X reaction

buffer, 1  $\mu$ L of enzyme mix, 0.2  $\mu$ L of 50 mM Mg<sub>2</sub>SO<sub>4</sub>, 1  $\mu$ L of bovine serum albumin (BSA; 1 ng/ $\mu$ L), and H<sub>2</sub>O in a total volume of 25  $\mu$ L.

The following thermal cycling parameters were used: 45 °C for 20 min and 94 °C for 3 min, followed by 5 cycles of 94 °C for 20 s, 55 °C to 45 °C (-2 °C per cycle) for 20 s, and 72 °C for 30 s; and then 40 cycles of 94 °C for 20 s, 45 °C for 20 s, 72 °C for 30 s; and finally 72 °C for 3 min. All amplification steps were performed using Maxy Gene gradient thermocycler (Axygen, USA). The products of the amplification reactions were analyzed by 1.2% agarose gel electrophoresis.

All PCR products of the expected size were cut out from the gel under UV illumination, extracted using a MinElute gel extraction kit (Qiagen, Germany) and ligated into the pGEM-T plasmid vector (Promega, USA). These plasmids were transformed into *Escherichia coli* (XL1 blue strain) (Maniatis et al., 1989) and 10 white colonies for each construct were picked and sequenced using standard M13R primers and ABI Prism 3500 XL sequencer (Applied Biosystems, USA). The obtained sequences were examined using the Basic Local Alignment Search Tool (BLAST, Altschul et al., 1990) NCBI (https://blast.ncbi.nlm.nih.gov/Blast.cgi).

## Primer design for the complete genome sequencing

In order to obtain the complete genome sequences of two PARV strains (excluding the 5' and 3' ends), a total of 60 primer pairs were designed (Table 2) using a web-based primer design tool – Primal Scheme (<a href="http://primal.zibraproject.org">http://primal.zibraproject.org</a>). This tool offers a complete pipeline for the development of efficient multiplex primer schemes for generation of the overlapping products; the size of which is determined by the target genome length, amplicon length, and required overlap (Quick et al., 2017). For PARV, we used the amplicon lengths of about 400 nts with a 50-nt overlap. The sequences of PARV strain LEIV–1149K (GenBank NCBI ac. nos. KP792717 – KP792719) were used as reference genomes.

# Library preparation and near complete genome sequencing

Reverse transcription reaction was performed using random hexanucleotide primers and Reverta-L kit (AmpliSens, Russia) according to the manufacturer's instructions. The cDNA was stored at -70 °C and used subsequently as a template for amplification. The designed primers were assorted into three groups, each containing twenty primer pairs. As a result, the three groups of 200–400 bp DNA fragments were amplified that were suitable for the subsequent 400-bp sequencing on an Ion S5 System (Table 2).

Hot-start multiplex PCR amplification reactions were performed in a 25-μL total volume containing 2 μL of the template cDNA, 0.1 μM of each sense primer, 0.1 μM of each antisense primer, 2.5 μL of dNTPs (1.76 mM, AmpliSens, Russia), and 10 μL of PCR buffer blue-2 with 7.5 mM MgCl<sub>2</sub> (AmpliSens, Russia). The following thermal cycling parameters were employed: 94 °C for 2 min;

40 cycles at 94 °C for 10 s, 55 °C for 15 s, 72 °C for 60 s; and a final extension at 72 °C for 5 min. The reactions were performed in a MaxyGene gradient thermocycler (Axygen, USA). The products were analyzed by 1.2% agarose gel electrophoresis (Figure 2a).

The library preparation included 5' phosphorylation and incorporation of barcoded adapters followed by the amplification of the final library. For this purpose, T4 Polynucleotide Kinase and T4 DNA Ligase (New England Biolabs, USA) were used with a slightly modified manufacturer's protocol. Amplification was performed using the PCR-mix 2 FEP/FRT kit (AmpliSens, Russia).

The concentrations of the fragments were measured with a Qubit 2.0 fluorimeter (Invitrogen, USA) using the Qubit dsDNA H Assay Kit (Invitrogen, USA). The quality assessment of the final libraries was carried out on an Agilent 2100 Bioanalyzer instrument (Agilent Genomics, USA), employing the Agilent High Sensitivity DNA Kit (Agilent Genomics, USA) (Figure 2b). The size selection of the final library was carried out on 2% E-Gel<sup>TM</sup> SizeSelect<sup>TM</sup> II Agarose Gels (Thermo Fisher Scientific, USA) on an E-Gel electrophoresis system (Thermo Fisher Scientific, USA).

Sequencing was performed using an Ion S5 System (Thermo Fisher Scientific, USA) with Ion 530 chips (Thermo Fisher Scientific, USA).

# Genetic analysis

The overall quality of the Ion S5 (Thermo Fisher Scientific) reads was confirmed using the FastQC software (Andrews, 2017), and low-quality sequences were filtered by the PRINSEQ program (Schmieder and Edwards 2011). The filtered reads were aligned to the reference viral protein sequences retrieved from the NCBI database (NCBI Resource Coordinators, 2017) using the BLASTx program (Altschul et al, 2017) to remove all non-target sequences. All the remaining reads were then mapped to the reference genome of Paramushir virus (GenBank NCBI ac. nos. KP792717 – KP792719) to confirm the coverage of the target regions using bwa mem tool (Li and Durbin, 2009). The individual sets of contigs were assembled for both samples using SPAdes (Bankevich et al, 2012) in a single cell mode and the resulting sequences were mapped to the reference Paramushir virus genome.

A dendrogram analysis was performed using the Jukes-Cantor substitution model and the trees were reconstructed using the Neighbor-Joining (NJ) tree algorithm in MEGA 6.0 software (Hall, 2013). Definition of SNV and estimation of identity were performed using Vector NTI v.7.0 software (Invitrogen, USA).

A test for probable recombinations was performed with SimPlot 3.5.1.0 software (Lole et al., 1999) and with the Recombination Detection Program (RDP) 4 beta 80 using eight methods provided by this software.

The availability and the location of signal peptides in translated nucleotide sequences were predicted using SignalP 4.1 software (<a href="http://www.cbs.dtu.dk/services/SignalP/">http://www.cbs.dtu.dk/services/SignalP/</a>)(Nielson, 2017). The

secondary structures of glycoproteins were predicted using PSIPRED v3.3 software (Buchan et al., 2013).

All molecular graphics were produced using PyMOL software (PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC).

#### 3. Results

## Sequencing

In two out of thirty analyzed samples (IDs 133 and 137), single bands of expected length for the Orthonairovirus genera were found using a broad range PCR. These bands were purified, sequenced by the Sanger method and examined using BLAST. Both sequences were of 400 bp in length and shared 97% nucleotide identity with the L segment of PARV (strain LEIV–1149K, ac. no KP792717). Thereby, both strains were identified as PARV. The strains were designated as Paramushir virus/Ixodes uriae/Russia/Tyuleniy Island/133/2015 and Paramushir virus/Ixodes uriae/Russia/Tyuleniy Island/137/2015 (briefly PARV strain 133 and PARV strain 137).

A total of 60 primer pairs were newly designed and used for the high-throughput sequencing library preparation for genomes of the two PARV strains. As a result, in total 489,436 and 358,075 sequencing reads were obtained for PARV strain 133 and PARV strain 137, respectively, and the nearly complete genomes of both strains were assembled (except for the 5' and 3' ends). The obtained reads were distributed unequally along the reference genome sequences. The S segments were 100% covered for both strains, the L segments were 98.4% and 95.7% covered for PARV strain 137 and PARV strain 133, respectively, and the M segments were 72.8% and 23.0% covered (for PARV strain 137 and PARV strain 133, respectively). The gaps were additionally sequenced using Sanger's sequencing technique by means of the ABI Prism 3500 XL (Applied Biosystems, USA); PARV strain 133 and PARV strain 137 were submitted to the GenBank NCBI under accession numbers MH124637, MH638289, MH124638, and MH124634-MH124636, respectively.

## Genetic analysis

The comparative analysis of PARV sequences showed the existence of SNVs in the sequence of the PARV strain 133 (L segment: n=260, including 19 nonsynonymous substitutions; M segment: n=861, 49 nonsynonymous; S segment: n=37, one nonsynonymous) as well as in the sequence of PARV strain 137 (L segment: n=232, 25 nonsynonymous substitutions; M segment: n=367, 49 nonsynonymous; S segment: n=35, 3 nonsynonymous). In addition, the PARV strain 133 had 114 unique SNVs in the L segment, 174 unique SNVs in the M segment and 6 unique SNVs in the S segment, whereas PARV strain 137 had 86 unique SNVs in the L segment, 11 unique SNVs in the M segment, and 4 unique SNVs in the S segment.

The two novel strains of PARV shared 98% and 97–98% identities with the L and S segment sequences of PARV LEIV–1149K (ac. nos. KP792717, KP792719), and differed from each other only by 1–2% within the L and S segments' nucleotide sequences. The comparisons of the M segment sequences demonstrated a 90%-nucleotide identity between PARV LEIV–1149K (ac. no KP792718) and PARV strain 137, whereas the nucleotide identity between strains PARV LEIV–1149K and PARV strain 133 was much lower (67%). Two newly sequenced strains shared 68% of nucleotide identity in the sequence of the M segment (Table 3). Thus, based on the nucleotide sequences of the new PARV strains, the M segment was found to be the most divergent.

The dendrogram analysis of nucleotide sequences of the L, M and S segments of PARV strain 133 and PARV strain 137, along with the appropriate sequences of other orthonairoviruses, supports their affiliation with *Orthonairovirus* genus (Figure 3). The location of PARV strain 133 and PARV strain 137 on the trees indicates their membership with the Sakhalin virus subgroup and a close relationship with the PARV strain LEIV–1149K, which was sequenced previously.

However, the dendrograms of different genomic segments revealed distinct genetic relationships among PARV LEIV-1149K, PARV strain 133, and PARV strain 137 (Figure 3). Regarding segments L and S, PARV strain 133 and PARV strain 137 form the reliably supported clade related to PARV strain LEIV-1149K. In the segment M, PARV strain LEIV-1149K and PARV strain 137 form the reliably supported clade related to PARV strain 133. Moreover, the different phylogenetic positions of SAKV strain LEIV-71S, ARTV strain LEIV-2366Arm, HAZV strain JC280, and TAMV strain LEIV–1308Uz sequences were observed. Regarding the L and S segments, the SAKV strain LEIV-71S grouped reliably with the SAKV strain LEIV-71C and TILV strain RML 86, but grouped with the AVAV strain CanAr173 and all known PARV strains in the M segment. In the L and M segments, the ARTV strain LEIV-2366Arm was an outgroup for the strains of PARV, SAKV, AVAV, TILV, TAGV and C (L)MV, whereas, in the S segment, TAFV strains AnD 11411 and ERVEV were reliably clubbed in the group, but the ARTV strain LEIV-2366Arm was an outgroup as well as the strains: ISKV LEZ 86-787 and UZAV LEIV-Kaz155. The CCHFV strain Shch 1 2016 was an outgroup for the reliably grouped HAZV strain JC280, NSHDV strain Ganjam G619, DUGV, KUPV strain K611 in the L and M segments, but in the S segment, it changed its place with HAZV strain JC280. In addition the TAMV strain LEIV–1308Uz was an outgroup for all analyzed strains in the L and S segments, but it reliably formed an internal group with the strains of PARV, SAKV, AVAV, TILV, TAGV, C (L)MV, ARTV strain LEIV-2366Arm, ISKV strain LEZ 86-787, and UZAV strain LEIV-Kaz155.

A Bootscan analysis with Simplot version 3.5.1 software showed the absence of probable recombination events in the L and S segments but revealed one potential recombination event at the nucleotide position 1,366–2,897 in the M segment (Figure 4a). An analysis provided by the RDP 4

beta 80 software showed two probable recombination events with recombinant fragments from the nucleotide position 1,523–1,910 and recombinant fragments from the nucleotide position 2,210–2,722 (according to genome alignment of Sakhalin virus genogroup); the PARV strain 133 is a recombinant strain, whose probable major parent is an unknown strain closely related to the AVAV. The PARV strain LEIV–1149K is a potential minor parent in the first probable recombination event, and the PARV strain 137 is a potential minor parent in the second probable recombination event (Figure 4 b).

# Analysis of putative proteins in the M segment

The glycoprotein precursors of orthonairoviruses form a number of smaller proteins after posttranslational modifications (Kuhn et al., 2016). The glycoprotein precursor of PARV strain 137 has a molecular weight of 153.4 kDa and contains 1376 aa, which is the same as that of PARV strain LEIV-1149K (AKC89338.1) (Figure 5 a). The cleavage site in the signal peptide was predicted at the N-terminal of the amino acid sequence of PARV strain 137 at position 28/29 aa, whereas in the amino acid sequences of the closely related viruses, its positions were 23/24 aa for PARV strain LEIV-1149K (AKC89338.1), 21/22 aa for AVAV strain CanAr 173 (AMT75378.1), 30/31 aa for SAKV strain LEIV-71C (AKC89335.1). This peculiarity has been described for CCHFV strains, whose cleavage sites are located from 20/21 aa to 27/28 aa of amino acid sequences (Goedhals et al., 2015). However, in contrast to CCHFV, two newly sequenced strains of PARV have no furindependent cleavage site R-X-[K/R]-R1. Therefore, instead of two proteins (Mucin-like protein and GP38), only one protein is formed after proteolysis from the N-terminal of the polypeptide (G1) with a molecular weight of 37 kDa. However, this peculiarity is common for a number of species belonging to Orthonairovirus genus (Lenz et al., 2001). The N-terminus of glycoprotein (Gn), for both PARV strain 133 and PARV strain 137, is cleaved off from the G1 by the means of subtilisin kexin isozyme 1 (SKI-1)/site 1 protease (S1P), which recognizes the conservative domain RRLL. As shown above, this protease is crucial in the surface glycoprotein processing for some pathogenic viruses, such as Lassa virus (LASV) (Lenz et al., 2001), lymphocytic choriomeningitis virus (LCMV) (Beyer et al., 2003), and CCHFV (Vincent et al., 2003).

The two smaller glycoproteins (Gn) contain 378 aa for the both PARV strain 133 and PARV strain 137, and have molecular weights of 42.3 kDa and 42.7 kDa, respectively. The C-terminal end of Gc is cleaved at site RRPL↓ by means of SKI−1/S1P-like protease. The 733RRPL736 motif is identical to that of the previously described PARV strain LEIV−1149K, but differs from that of the closely related species AVAV strain CanAr 173 (730RKPL733) and SAKV strain LEIV−71C (730RKPL733). The C-terminal Gc protein of PARV strain 137 highly conserved, with only 7 aa substitutions (98.9% aa identity) in comparison with PARV strain LEIV−1149K (with 8

substitutions in Gn (95.2%) and 24 in G1 (91.3%)). Moreover, in comparison to the closely related viruses, this protein is also highly conserved with 87.8% aa identity with Gc of AVAV strain CanAr 173 and 74.7% aa identity with Gc of SAKV strain LEIV–71C. Sixty-six amino acid substitutions in the sequenced fragment of PARV strain 133 Gc were found in comparison with PARV strain LEIV–1149K (86.4% aa identity) and 55 (85.4%) substitutions in Gn. The N-terminus of the G1 protein was the most variable fragment. The sequenced fragment of PARV strain 133 G1 had 41 substitutions in comparison with PARV strain LEIV–1149K, but only 30 substitutions in comparison with AVAV strain CanAr 173. The four transmembrane domains of the polyprotein M were predicted by the TMHMM server, which form two regions outside of the membrane, and two regions in the cytoplasm. This is most likely similar to the structure observed in Qalyub virus (QYBV) (Walker et al., 2013).

The M segment of PARV strain 137 contains 63 cysteine residues, whereas the sequenced fragment of M segment in PARV strain 133 contains 50 cysteine residues located mainly in the conservative positions in comparison with PARV strain LEIV–1148K (AKC89338.1), AVAV strain CanAr 173 (AMT75378.1), and SAKV strain LEIV–71C (AMT75420.1). This suggests the presence of a large number of disulfide bonds and a complex spatial structure of proteins similar to the related viruses. For instance, by comparing the C-terminus of Gn with CCHFV, which is the only virus with an identified molecular structure among orthonairoviruses, the presence of the conservative motifs CXCXHXC of cysteine and histidine (where X is any amino acid) was characteristic of the ββα-type zinc fingers of bunyaviruses (Figure 5 b). In some *in vitro* experiments, the possibility of RNA binding has been shown for this structure, which may indicate the participation of Gn-tail in interaction with viral RNA (Estrada et al., 2011). In general, the molecular structure of orthonairoviruses is poorly studied, even for the CCHFV proteins, and nor the spatial structure, nor the functional domain have been completely described.

The secondary structures of G1, Gn, and Gc, deduced from PSIPRED server and identified as the distribution of helices, strands, and coils, are shown in Supplementary Figures 1, 2, 3, respectively. It can be seen that the secondary protein structures are highly conservative for all compared viruses. A significant difference can be seen only at the N-terminus of G1 of the SAKV virus. The experimental high-resolution structural data have been determined only for Gc protein of several othounyaviruses: Rift Valley virus (RFV), Puumala virus (PUUV) and Heartland virus (HRTV) (Dessau and Modis, 2013, Willensky et al., 2016, Zhu et al., 2018). The structural studies have revealed that both hantaviral and phenuiviral Gc fold as a class II membrane fusion proteins, which is induced by a low pH during endocytosis and undergo a conformational change by exposing a hydrophobic fusion peptide or fusion loop. This is highly possible that the tertiary structure of Gc protein from strain 137 has the same post-fusion trimeric conformation and is also a member of the

class II fusion protein. The bioinformatical methods, considering the amino acid sequences of most closely related PUUV, have partially predicted the tertiary structure of the PARV strain 137 Gc protein (Figure 6a) (Biasini et al., 2014). The sequences of domains I and II match well with the sequence of the Puumala virus, but domain III differs to a large extent, and no spatial structure could be determined, same being true for the particularly sequenced M segment of PARV strain 133. However, a conserved block of the sequences forming putative fusion loops and the endosomal membrane anchors has been seen as a part of the domain II, which is distal to the domain I (Figure 6b). The fusion loop, which is responsible for the initial insertion into the target membrane, is the key element in the fusogenic process (White et al., 2009)

PARV strain 133 and PARV strain 137, as well as other bunyaviruses, have more than one fusion loop (Figure 6c) (Zhu et al., 2018). In comparison with the PUUV, on which the model was built, there is a significant difference in the first loop: for PARV strain 137 and PARV strain 133 they are outside of the bonding plane with the membrane (Figure 6d). However, this could be due to a flaw in the constructed model. The two remaining loops corresponding to the bc and ij loop of PUUV have all the conserved residues and two disulfide bonds fixing the fusion loop conformation. At the same position, PARV strain 137 Gc fusion loop contains a three-residue insertion 888P-T-W890, which differs from 777P-X-D779, mostly conserved among hantaviruses (Willensky et al., 2016). Instead of the conserved F907 of hantaviruses, a proline amino acid (P1057 in PARV strain 137 isolate) is present in the Sakhalin genogroup. As previously shown, Gc of Andes virus (ANDV) with single mutations in the conserved residues W773, N776, and D779, located at the fusion loop, eliminated the cell-cell fusion activity and infectivity of the ANDV pseudotyped particles (Cifuentes-Muñoz et al., 2011). The third amino acid (D) of PARV strain 137 and PARV strain 133 is replaced by W (W884, N887, and W890) and for PARV strain 133 N is changed to I. This suggests that, despite the fact that the structure of the glycoprotein C in the Sakhalin genogroup has a high homology with other representatives of bunyaviruses, the substitutions in the conserved residues most likely result in the differences in pathogenesis processes and pathways and pathogenicity to humans.

### 4. Discussion

In general, within the Sakhalin genogroup, the nucleotide identities ranged from 78% to 98% in the L segment, from 69% to 90% in the M segment, and from 66% to 99% in the S segment. Therefore, the S segment is known as the most variable region within this genogroup (see Table 3). However, based on our data, the sequences of the M segment among PARV strains share a lower identity in comparison to the average volume within the genogroup. This peculiarity likely explains why the

number of sequencing reads belonging to the segment M was the lowest and caused significant difficulties in the primer design and sequencing.

By using phylogenetic analysis, we observed close relations between the novel strains and PARV strain LEIV–1149K, also confirmed their association with the Sakhalin genogroup. Moreover, the phylogenetic analysis identified distinct and reliable grouping patterns among PARV LEIV–1149K and PARV strains 133, 137 in the different genome segments. This indicates that the M segment reassortment is common in PARV and also supports the assignment of all these strains to the same species, because their genome segments are genetically compatible and could be involved in the reassortment. Besides PARV, other orthonairoviruses that were analyzed also demonstrated signs of multiple reassortment, in which both M and S segments were involved. Such peculiarity is probably caused by genome structure and comparability and allows for the reassortment process.

In view of the change in tree topology at the M segment coding the region of PARV strain 133, recombination analysis was performed to detect the probable recombination events.

Using RDP 4 beta 80 software, two probable recombination events in a fragment from nucleotide positions 1,523–2,722 were revealed. The unknown strain that belongs to the Sakhalin genogroup, which is most closely related to the AVAV strain CanAr173, was the major parent in both events. The minor parent in the first event was the strain closely related to PARV LEIV–1149K, while in the second event it was the strain that is most closely related to PARV strain 137. Given the close location of the probable recombination events and a high nucleic acid identity between both minor parents, one expanded recombination event could be assumed in a fragment from nucleotide positions 1,523–2,722 bp (according to genome alignment of Sakhalin virus genogroup). In the M segment of PARV strain 133, this fragment is located from nucleotide positions 1,100 to 2,317.

A single probable recombination event was confirmed by SimPlot analysis. However, RDP 4 beta software indicated a narrower range of recombination site due to the use of several statistical models. The minor parent in this event was, probably, an unknown strain of PARV, sharing a number of features with PARV LEIV–1149K and PARV 137.

The pairwise analysis of the M segment's nucleic acid identity within the Sakhalin genogroup particularly confirmed this assumption. The M segment identity of PRMV strain 133 with other known PARV strains in the range 1523–2722 bp was significantly higher than that for the whole M segment. The inherited fragment of the major parent was closer to the strains of PARV, rather than to the strains of AVAV. However, the nucleic acid identity did not exceed 61%.

The recombination phenomenon as a source of genetic diversity of orthonairoviruses was previously described for the CCHFV (Lukashev, 2005). The recombination event along with the intersegment reassortation points to a high variability potential of the virus, which may lead to the formation of strains with new properties.

#### 5. Conclusions

In this study, we report the identification and nearly complete genome sequencing of two strains of Paramushir virus using a combination of high throughput sequencing and specific multiplex PCR following an earlier approach (Quick et al., 2017). The results of the sequence analysis confirmed the applicability of this approach for resequencing of known viral genomes, especially in the case of a lack of viral RNA.

Both strains of PARV were isolated in 2015 on the Tyuleniy Island in the Okhotsk Sea after a 43-years break. The phylogenetic analysis was used to determine their relation with the Sakhalin genogroup in the *Orthonairovirus* genus and with PARV strain LEIV–1149K. The signs of multiple reassortment among the orthonairoviruses, including PARV, and of probable recombination events within the Sakhalin genogroup were revealed. However, these phenomena need to be studied in more detail.

We hope our study will contribute to the knowledge of orthonairoviruses, which is still poor, especially regarding the Sakhalin genogroup of unknown pathogenicity. For a more detailed understanding of the complicated phylogeny within the Sakhalin genogroup, further ecological studies are needed, and additional isolates belonging to the genogroup should be analyzed.

More data regarding Paramushir virus could provide a better understanding of its diversity and relation within *Orthonairovirus* genus and could help define intragenic demarcation criteria which have not yet been established.

#### **Foundation**

This work was supported by RSF grant N. 17-74-20096

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Table 1. Primers used for broad-range PCR

Primer set no	Genera	Primer name	Sequence 5'-3'	Expected Product Length, bp	References
1	Flavivirus	PhIP2 PhIM2	GGCTACTTCAARAAYAARGANGA CTCTCTCAGICCICCRTGYTG	507	Klimentov et al., 2016
2	Orthonairo- virus	Nairo-F Nairo-R	TCTCAAAGAAACACGTGCCGC GTCCTTCCTCCACTTGWGRGCAGCCTGCTG GTA	400	Lambert and Lanciotti 2009
3	Phlebovirus	Phlebo F1 Phlebo R1	TTTGCTTATCAAGGATTTGACC TCAATCAGTCCAGCAAAGCTGGGATGCATCA T	370	Lambert and Lanciotti 2009
4	Ortho- bunyavirus	Bun F Cal/Bwa F Oropouche F Wyeomyia-F Bun R Cal/Bwa R Oropouche R Wyeomyia-R	CTGCTAACACCAGCAGTACTTTTGAC GCAAATGGATTTGATCCTGATGCAG GGCCCATGGTTGACCTTACTTT ATGTCTGAAATTGTATTTGATGATATTTGG TGGAGGGTAAGACCATCGTCAGGAACTG TTGTTCCTGTTTGCTGGAAAATGAT ACCAAAGGGAAGAAAGTGAAT TATTTCGATTCCCCGGAAAGT	222-286	Lambert and Lanciotti 2009 This study
5	Orbivirus	OrbiVP1 F2295a OrbiVP1 F2295b OrbiVP1-R2523a OrbiVP1 R2523b OrbiVP1 R2523c OrbiVP1 R2523d OrbiVP1 R2523e OrbiVP1 R2523f	CGGAGCAGTATGTGGGNGATGAYA CGGAGATGTATGTGGGNGATGAYA TCWGAWGAKATKATCATCATTCGRTCTTG TCAGACGAGATCAACATCATTCGRTCTTG TCAGAGGAGACCATCATCATTCGRTCTTG TCCGATGAAATGAACATCATTCGRTCTTG TCGGACGAAACTAGCATCATTCGRTCTTG TCAGAACTGACAATCATTCGRTCTTG	228	Dedkov et al. 2016
6	Orthoreovirus	Orthoreo -2090F Orthoreo -2334R	GGBTCMACNGCYACYTCBACYGAGCA CDATGTCRTAHWYCCANCCRAA	244	Landolfi et al. 2010

Table 2. Primers used for near complete genome sequencing of  $\ensuremath{\mathsf{PMRV}}$ 

Region No.	Pool	Left Primer Name	Left Primer Sequence	Right Primer Name	Right Primer Sequence	Produ ct Length
1	1	ParamL_1 _LEFT	TCCTTTAACTGGCATCCCTCTCT	ParamL_1_ RIGHT	CGTAACGCATGACATGGTCACA	386
2	2	ParamL_2 _LEFT	AACGAATGGGGCTCAACACTTG	ParamL_2_ RIGHT	TGGGGATCAGCTTTCGAACTCT	407
3	3	ParamL_3 _LEFT	GTTGGTGAACGAGGAAGAGCTG	ParamL_3_ RIGHT	GTCCTTAGCAACCCAGGTAGGT	373
4	1	ParamL_4 _LEFT	CTCTGACTCTAAGAACACTTGGTCAT	ParamL_4_ RIGHT	GGATGCTCGTACAGGTCCTTCT	365
5	2	ParamL_5 _LEFT	AGAAGAAGTTCATAACCAACTGCCT	ParamL_5_ RIGHT	GCTCTGTCAACTGCTCCTTGTC	382
6	3	ParamL_6 _LEFT	TGAAACAGAAGACTTTGGTCACCT	ParamL_6_ RIGHT	AGGAAGGATTGCCTGTGTTCCT	
7	1	ParamL_7 _LEFT	AGGAGAGTCTGAAGCTGAGCAC	ParamL_7_ RIGHT	TGAACTCTGGGTGAAGCTGAGT	400
8	2	ParamL_8 _LEFT	ATGGAGGTATGTGCCAGAGGAC	ParamL_8_ RIGHT	TTGATGAGGAGGCCGTCTTCAG	406
9	3	ParamL_9 _LEFT	CGGTGCCATTGAAGTGTTCGAA	ParamL_9_ RIGHT	TTAACCACTGACCCTGCCTTGA	427
10	1	ParamL_1 0_LEFT	TCCTCAAATCTTCAATCAGCTTTCTCT	ParamL_10_ RIGHT	GGTCTTCCTTTAGCCAGGCAAC	424
11	2	ParamL_1 1_LEFT	TCATTGACGCCTTTGAGAGAACA	ParamL_11_ RIGHT	GTGAGCCTTACCAGCCTATCCA	368
12	3	ParamL_1 2_LEFT	CCCACAGTGAGACGAAGAGTTG	ParamL_12_ RIGHT	CTGGCACTTTGAGTTGGCTTGA	403
13	1	ParamL_1 3_LEFT	AGTTGGCAGAGGAGAAGCTGAA	ParamL_13_ RIGHT	GCCTCAACATCAATTCCGCAGA	361
14	2	ParamL_1 4_LEFT	AGGAGACTACATCAGCAACAAGC	ParamL_14_ RIGHT	CAGGATGTAGCTGTAGGAGGCT	431
15	3	ParamL_1 5_LEFT	TGACCTGCTGTTTATACAATAGCAGT	ParamL_15_ RIGHT	GGAGCTGTTTGTTTTGCTGCTG	424
16	1	ParamL_1 6_LEFT	ACACTCATGACCACTTTATAAGGGTT	ParamL_16_ RIGHT	CGATGCAGCCCTCATCAAAGTT	430
17	2	ParamL_1 7_LEFT	TCTATGGAGCCTGGGTCAACAG	ParamL_17_ RIGHT	TCCCTTTCTACCTTGAACCCTGA	428
18	3	ParamL_1 8_LEFT	AGGTCGGTTGGTTCCAGTAACA	ParamL_18_ RIGHT	AGCATGTGTGTCTTTGGGACTG	387
19	1	ParamL_1 9_LEFT	TCCCACCTGATGCAATAGAAAAGT	ParamL_19_ RIGHT	GCTGCTAAGGTCTTAAGCCAGC	419

20	2	ParamL_2 0_LEFT	TTCTACCACTGGAAGGAGCTGG	ParamL_20_ RIGHT	AGTTAGAGTTGGGAGTGCAGCT	370
21	3	ParamL_2 1_LEFT	TTCAGAGTGCCTGGGAGAAGAC	ParamL_21_ RIGHT	CCAAGAGTTGGCAGTCAGGTTC	376
22	1	ParamL_2 2_LEFT	CCCTGGTGTGTCCACTACAAGT	ParamL_22_ RIGHT	CCATAGAAACTCATGCCTGCCG	370
23	2	ParamL_2 3_LEFT	GCGTGAGAAGGTTGTTCAAGGT	ParamL_23_ RIGHT	CCATCATCGTTCGTTGTGCTCA	427
24	3	ParamL_2 4_LEFT	AGGCTGTCGAGATCTACTGGTG	ParamL_24_ RIGHT	TGCAGAGGGTATTTCCACCTGT	389
25	1	ParamL_2 5_LEFT	GTGGAATGATGCAGCAGCTACT	ParamL_25_ RIGHT	TGAAAAGAAGTCTCGCGCTAGC	387
26	2	ParamL_2 6_LEFT	AGCTACAATCACATGGGTCAAGG	ParamL_26_ RIGHT	TGATCACTGCCGGTGTAACTCT	377
27	3	ParamL_2 7_LEFT	GCCCGATGCTGTCAGATGAAAG	ParamL_27_ RIGHT	AGCCGAGCAAGAGATAACCTCA	409
28	1	ParamL_2 8_LEFT	TGACGCTTGGGTCATTATCACC	ParamL_28_ RIGHT	TGGACTTTGGATCCAGAGGTCC	409
29	2	ParamL_2 9_LEFT	GCCAGAACCATCCAAGATTACTCC	ParamL_29_ RIGHT	GTTGCAAACCTGGTAGGAGCTC	407
30	3	ParamL_3 0_LEFT	GAAGACCCAATGATACAACTCATCCC	ParamL_30_ RIGHT	GTCTCTGTAGTTCGTGCTGTGC	361
31	1	ParamL_3 1_LEFT	ACTCTCAGTGCCTCTGTACACA	ParamL_31_ RIGHT	CAAGTTCTGCTTCTGGAGGCTG	414
32	2	ParamL_3 2_LEFT	CAATGACCTAGACATACTTCACAAAG AATT	ParamL_32_ RIGHT	ACTGCAGTCCAGATCCTGTTGT	371
33	3	ParamL_3 3_LEFT	CCACAAGGATAGATCTATCAAACCTG C	ParamL_33_ RIGHT	TGGCAGAGATGCAGTGGAGAAT	382
34	1	ParamL_3 4_LEFT	TCGTATGGGATCAAAGAAGGAAGGT	ParamL_34_ RIGHT	GGCCCTCTCCCCATGTCTTAAA	413
35	2	ParamL_3 5_LEFT	AAGGAAAAGTATAAGTGACCTTATCG ACT	ParamL_35_ RIGHT	AGAGAGACGGAGGCTATCTTGC	370
36	3	ParamL_3 6_LEFT	CGGACTTAGTCTCAAAGGAACAGC	ParamL_36_ RIGHT	AGTTTCGGAGAAAGTTGCATTT GC	384
37	1	ParamL_3 7_LEFT	AGAGACTCAGACCACATACCATGA	ParamL_37_ RIGHT	TCCTTGTCAGTGATGTTGCTGT	367
38	2	ParamL_3 8_LEFT	TCCTGTCAAGTGTTCTCTGGCA	ParamL_38_ RIGHT	ACTCATAAACGCATCAATGGTG CT	382
39	3	ParamL_3 9_LEFT	GCTTTCCGCTGTTTGGAACTTC	ParamL_39_ RIGHT	AGCATTAGGACGGACACAGTGA	361
40	1	ParamL_4 0_LEFT	TGTCGATTGTCAGAAGCAGAAGC	ParamL_40_ RIGHT	CTCTCGGTTGTTGCTGCTGATG	429

41	2	2 ParamL_4 CCACCATTGGCAACCTCTTCA		ParamL_41_ TCTGGCTCAGGAACCACT		368
42	3	ParamM_1 _LEFT	=		ACAAAATTAAGGCCGGTCCTGC	408
43	1	ParamM_2 _LEFT	AGTCTGTTTACCACTTTCATTGAAGG T	ParamM_2_ RIGHT	TTGTCACAGGAGGCCTTCTTGA	381
44	2	ParamM_3 _LEFT	CAAAGGAAGATAGAGCACCTTGGT	ParamM_3_ RIGHT	AATTGTGCACTCCTCGTTGTGG	375
45	3	ParamM_4 _LEFT	TGGTACCATGTCGAATTGGTGG	ParamM_4_ RIGHT	GTAAGCTTGTCACCCTTGTGCC	378
46	1	ParamM_5 _LEFT	GCATCCAACTCCACTTTGATTCCA	ParamM_5_ RIGHT	GCATTGGGCAGCTTATGTCGAT	431
47	2	ParamM_6 _LEFT	CTTCAGTGATGCAGTGGGTGAC	ParamM_6_ RIGHT	GAACTTGGTGCTGCAGTAAGGG	395
48	3	ParamM_7 _LEFT	TGCCCTGAATGCCTAGACAGAT	ParamM_7_ RIGHT	CTGGTTGTCACACTGGCTAACC	385
49	1	ParamM_8 _LEFT	TCATGCTTGTCTCTCCGGTGAA	ParamM_8_ RIGHT	CTTTGCTGCTGGAAAGTGACCA	424
50	2	ParamM_9 _LEFT	TGGGACTCAGAAGAAGAAGTTGGT	ParamM_9_ RIGHT	CATGAGAACTGCCAGGTGTTCC	406
51	3	ParamM_1 0_LEFT	GCAAGAAATTGGGGCTGCAATC	ParamM_10 _RIGHT	CGCTCTTGACAAGGTGATCCAC	431
52	1	ParamM_1 1_LEFT	ACATCCTGACATGGTCACGACT	ParamM_11 _RIGHT	GCTTTGCCTTGACATCCCAAGT	375
53	2	ParamM_1 2_LEFT	ACACAAAGTCGAAAGAGTGCTCT	ParamM_12 _RIGHT	ACTGACAGGGGAGAACAGCTTT	398
54	3	ParamM_1 3_LEFT	GCCTGACAGATTTGCAGTGCAT	ParamM_13 _RIGHT	ATGACAAGCCCAGCGCAAATAA	424
55	1	ParamM_1 4_LEFT	TGCTAGAAAACAGAGGGACCATCA	ParamM_14 _RIGHT	TCAGTGAGGTGGGAGAATGCTT	383
56	2	ParamS_1 _LEFT	GCAAGCTGGACTTTGAGGACAA	ParamS_1_R IGHT	CACTTGCAGTCTCGGTGAACTC	436
57	3	ParamS_2 _LEFT	TCAAGAAGCGTGTCCCTCTTCT	ParamS_2_R IGHT	TACAAGTTTGCACAGGCCAGTG	372
58	1	ParamS_3 _LEFT	TTAGGCGTAGGAACCTTGTGCT	ParamS_3_R IGHT	ATCATTGCACCGCTTGACTCTG	368
59	2	ParamS_4 _LEFT	AGATGGTTGAGGAGCCTGACAA	ParamS_4_R IGHT	CATGCAGCCATATCTGACAGCC	401
60	3	ParamS_5 _LEFT	CCTTCAAGTGGGGTAAGGGGAT	ParamS_5_R IGHT	GCCCTTCACTCTTGAGACGTTG	366

 $Table \ 3. \ Nucleotide \ Identities \ of \ the \ L, M \ and \ S \ segment \ sequences \ within \ the \ Sakhalin \ genogroup$ 

strain	PARV LEIV-1148K identity, nucleotide level, %			PARV nucleoti	133, identity, de level, %		PARV 137, iden		identity,
	L	M	S	L	M	S	L	M	S
PARV LEIV-1148K	100	100	100	98	64	97	98	90	98
PARV 133	98	64	97	100	100	100	98	64	99
PARV 137	98	90	98	98	64	99	100	100	100
Avalon CanAr173	78	75	82	78	57	83	78	74	83
Avalon Bres T/Ar/T-261	78	74	82	78	56	82	78	74	82
Sakhalin LEIV-71C	77	67	66	77	47	66	77	67	66
Sakhalin LEIV-71S	76	68	67	76	52	66	76	68	66
Tillamook RML 86	76	67	67	76	47	66	76	67	66
Taggert Ml 14850	73	68	67	73	48	67	73	67	67
Clo Mor SCOT Ar7	75	69	69	75	49	68	76	69	69

# **Figure Legends**

# Figure 1. Island territories of the Russian Far East (courtesy Google maps)

- a-Sakhalin Island and Kuril Ridge
- b- Tyuleniy Island the territory where PARV strain 133 and PARV strain 137 were isolated
- c-Paramushir Island the territory where the first isolation of PARV occurred

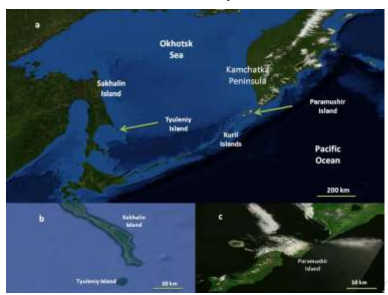


Figure 2. Library preparation for near complete genome sequencing using the Ion S5 system

- a PCR products after multiplex PCR
- b Mix of final RARMV libraries for Ion S5 system after size selection

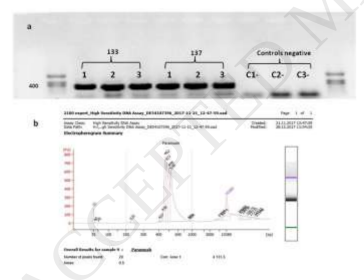


Figure 3. Dendrograms for near complete L, M and S sequences of PARV (at the nucleotide level)

Phylogenetic analysis was performed using the Jukes-Cantor substitution model and the trees were reconstructed using the Neighbor-Joining (NJ) tree algorithm in MEGA 6.0 software. The robustness of the trees was tested using 1000 bootstrap replicates. The locations on the trees of the Paramushir virus/Ixodes uriae/Russia/Tyuleniy Island/133/2015 and Paramushir virus/Ixodes uriae/Russia/Tyuleniy Island/137/2015 strains are marked with black dots.

A – Dendrogram for near complete L sequences of PARV. Other orthonairoviruses used for tree building

- B Dendrogram for near complete M sequences of PARV. Other orthonairoviruses used for tree building
- c Dendrogram for near complete S sequences of PARV. Other orthonairoviruses used for tree building



## Figure 4. Results of recombination analysis

- **a Bootscan analysis**: A Bootscan analysis of the M segment was performed using Simplot version 3.5.1 software. Probable recombination fragments are located in the region from 1366 to 2897 nt.
- **b- Analysis RDP 4 beta 80 software.** Two probable recombination fragments were revealed (first was located in the region from 1523 to 1910 nt, second was located in the region from 2210 to 2722 according to genome alignment of Sakhalin virus genogroup)

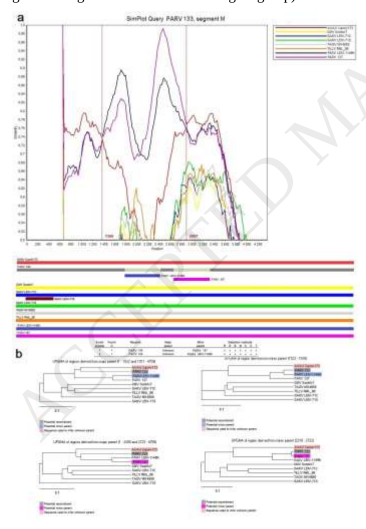


Figure 5. Putative proteins encoded in M segment of PARV

- a Schematic illustration of the structures of the polyglycoproteins encoded in the M segments of Sakhalin orthonairoviruses. Regions corresponding to the G1 (orange), Gn (green) and Gc (blue) are shaded. Dotted lines represent unsequenced regions. Predicted signal peptidase cleavage sites (SP) and potential SKI-I cleavage sites are shown. The predicted molecular weights (kDa) of GPCs are annotated.
- b. Sequence alignment of the Gn tails of the members from the Sakhalin genogroup with the known structure of zinc fingers of CCHF. The conserved zinc finger motifs (boxed) and conserved basic residues (in blue) are mapped.

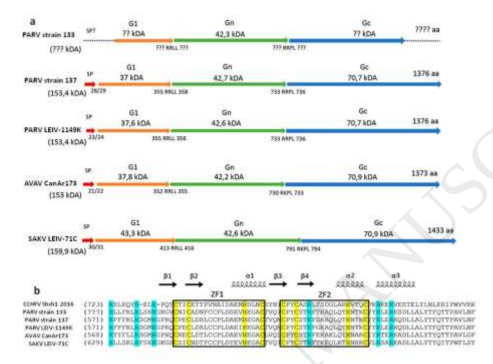


Figure 6. Schematic representation of PARV strain 137 Gc structure

- a Schematic 3D diagram of predicted PARV strain 137 Gc structure (domain I and II). A monomeric form with fusion loop residues is displayed in red.
- b Trimer form in different projections.
- C **Structure-based multiple sequence alignment of the fusion loop regions**. The PXX PUUV-like insertion is marked by a black box. The secondary structure elements are displayed below the sequences. Strictly conserved, identical, and highly similar residues are highlighted in yellow, blue and green, respectively. The residues correspond to polyprotein precursor numbering.
- **d** The putative fusion loop of Puumala GC and the fusion loops of the Sakhalin group viruses. The hydrophobic residues that anchor the protein to the cellular membrane are shown in red and the disulfide bonds in stick representation.
- \*All molecular graphics were produced using PyMOL (PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC).

