

1 **Revisiting rustrela virus – new cases of encephalitis and a solution to the**
2 **capsid enigma**

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25 **Running Title:** Revisiting rustrela virus

26 **Key Words:** rustrela virus; rubivirus; sequencing; capsid; intergenic region; encephalitis;
27 Eurasian otter; South American coati; yellow-necked field mouse.

28 Abstract

29 Rustrela virus (RusV, species *Rubivirus strelense*) is a recently discovered relative of
30 rubella virus (RuV) that has been detected in cases of encephalitis across a wide spectrum
31 of mammals, including placental and marsupial animals. Here we diagnosed two additional
32 cases of fatal RusV-associated meningoencephalitis in a South American coati (*Nasua*
33 *nasua*) and a Eurasian otter (*Lutra lutra*) that were detected in a zoological garden with
34 history of prior RusV infections. Both animals showed abnormal movement or unusual
35 behaviour and their brains tested positive for RusV using specific RT-qPCR and RNA *in situ*
36 hybridization. As previous sequencing of RusV proved to be very challenging, we employed
37 a sophisticated target-specific capture enrichment with specifically designed RNA baits to
38 generate complete RusV genome sequences from both detected encephalitic animals and
39 apparently healthy wild yellow-necked field mice (*Apodemus flavicollis*). Furthermore, the
40 technique was used to revise three previously published RusV genomes from two
41 encephalitic animals and a wild yellow-necked field mouse. Virus-to-host sequence ratio
42 and thereby sequence coverage improved markedly using the enrichment method as
43 compared to standard procedures. When comparing the newly generated RusV sequences
44 to the previously published RusV genomes, we identified a previously undetected stretch
45 of 309 nucleotides predicted to represent the intergenic region and the sequence encoding
46 the N-terminus of the capsid protein. This indicated that the original RusV sequence was
47 likely incomplete due to misassembly of the genome at a region with an exceptionally high
48 G+C content of >80 mol%, which could not be resolved even by enormous sequencing
49 efforts with standard methods. The updated capsid protein amino acid sequence now
50 resembles those of RuV and ruhugu virus in size and harbours a predicted RNA binding
51 domain that was not encoded in the original RusV genome version. The new sequence data
52 indicate that RusV has the largest overall genome (9,631 nucleotides), intergenic region
53 (290 nucleotides) and capsid protein-encoding sequence (331 codons) within the genus
54 *Rubivirus*.

55 **(316 words; max. 350 words)**

56 Introduction

57 Rubella virus (RuV; species *Rubivirus rubellae*) was the sole member of the family
58 *Matonaviridae* and the genus *Rubivirus* ¹, until recently its first relatives rustrela virus
59 (RusV; *Rubivirus strelense*) and ruhugu virus (RuhV; *Rubivirus ruteetense*) were
60 identified ². While RuhV was detected in apparently healthy cyclops leaf-nosed bats
61 (*Hipposideros cyclops*) in Uganda, RusV was associated with cases of fatal neurological
62 disease in placental and marsupial zoo animals in Germany. RusV was initially identified
63 using a metagenomic sequencing workflow from brain tissues of a donkey (*Equus asinus*),
64 a capybara (*Hydrochoeris hydrochaeris*), and a red-necked wallaby (*Macropus rufogriseus*)
65 between July 2018 and October 2019 ^{2,3}. All of these animals were housed in a zoological
66 garden located in northeast Germany, close to the Baltic Sea, and developed acute
67 neurological signs such as ataxia and lethargy, which ultimately resulted in death. RusV
68 was mainly detected in the central nervous system of these animals and only sporadically
69 and in very low concentrations in extraneural organs. RusV-infected wild yellow-necked field
70 mice (*Apodemus flavicollis*) were identified in close proximity to the encephalitic animals'
71 housings. These rodents were considered as a likely reservoir host, as they carried the virus
72 without obvious encephalitis whereas all tested individuals of other sympatrically occurring
73 rodent species at the same location were RusV-negative ². However, the mode of
74 transmission between potential reservoir and accidental spill-over hosts still remains to be
75 identified ². Currently, no isolates of either RusV or RuhV are available and therefore, most
76 data are limited to *in silico* predictions and analogies with RuV. Furthermore, sequencing
77 of RusV from organ samples proved to be extremely difficult and only three full-length
78 genome sequences and a few partial coding sequences are currently available (**Table 1**).

79 The genome of rubiviruses consists of single-stranded positive sense (+ss) RNA, that
80 contains two open reading frames (ORF) encoding the non-structural p200 and structural
81 p110 polyproteins, respectively ⁴. Both ORFs are separated by an untranslated intergenic
82 region (IGR). In RuV, co-translational cleavage of the p110 polyprotein results in three
83 structural proteins E1, E2, and the capsid protein ⁴. After cleavage by cellular signal
84 peptidase, the capsid protein remains in the cytoplasm while E1 and E2 enter the secretory
85 pathway using distinct translocation signals ⁵. Based on sequence comparison with RuV,
86 the genomes of RusV and RuhV are likewise predicted to encode the p110 polyprotein and
87 the mature capsid, E1 and E2 proteins ². In RuV, the capsid protein consists of a structurally
88 disordered N-terminal part that contains a RNA-binding domain (RBD) ⁶ and a structurally
89 ordered C-terminal domain (CTD) ^{7,8} containing the E2 signal sequence ⁹. While the

90 predicted capsid protein sequence and structure of RuhV is analogous to that of RuV, the
91 capsid protein of RusV was considered enigmatic as it appears truncated and lacking e.g.
92 the RBD ¹⁰.

93 Here we analysed the RusV genome sequences from two novel cases of RusV encephalitis
94 using a sophisticated target-specific capture enrichment with RNA baits prior to
95 sequencing. This resulted in markedly improved virus-to-background sequence ratios and
96 higher genome coverage particularly in regions of exceptionally high G+C ratios of
97 >80 mol%. The *de novo* assembled sequences suggested a 309 nucleotide (nt) longer
98 RusV genome sequence than initially reported. We also confirmed the sequence extension
99 by reanalysing samples from previously published diseased animals and potential reservoir
100 animals using the same methods, and finally solved the enigma of the unusual RusV capsid
101 protein sequence. So far, the clinical and pathological data are limited to the first
102 description of RusV fatal encephalitis ². We now present further clinical data and an in
103 depth pathological and histopathological evaluation of the two new cases.

104

105 **Material and Methods**

106 **Animals and samples included in this study**

107 Brain samples were collected from a South American coati that was housed in a zoological
108 garden in the Northern Germany, a wild Eurasian otter that was found nearby the zoo and
109 three yellow-necked field mice that had been trapped during pest control measures at the
110 zoo (**Table 1**). In addition, samples from previously published animals, including a donkey,
111 a capybara and seven yellow-necked field mice, were re-analysed during this study ².

112 **Histopathology, immunohistochemistry and RusV RNA *in situ* hybridization**

113 Routine staining, immunohistochemistry (IHC) as well as RNA *in situ* hybridization (RNA ISH)
114 was applied as described earlier with minimal adaptations summarized in Supplemental
115 Table S1 (see also ²). Briefly, formalin-fixed, paraffin-embedded (FFPE) brain tissues were
116 processed for haematoxylin and eosin (HE) staining and examination using light
117 microscopy. On consecutive slides, conventional Prussian Blue staining was used to
118 demonstrate haemosiderin, whereas Luxol Fast Blue Cresyl Violet was applied for detection
119 of myelin sheaths and Nissl substance. Immunohistochemistry was performed according
120 to standardized procedures using markers to detect T-cells (CD3), B-cells (CD79a),
121 microglial cells and macrophages (IBA1), astrocytes (glial fibrillary acidic protein, GFAP) and

122 apoptotic cells (active caspase 3). A bright red chromogen labelling was produced with 3-
123 amino-9-ethylcarbazole substrate (AEC, DAKO). Sections were counterstained with Mayer's
124 haematoxylin. RNA ISH was performed with the RNAScope 2-5 HD Reagent Kit-Red
125 (Advanced Cell Diagnostics, USA) according to the manufacturer's instructions using a
126 custom-designed probe against the RusV non-structural protein (p200, NSP) ORF, and a
127 negative control probe against the dihydrodipicolinate reductase (*DapB*) gene. Analysis and
128 interpretation were performed by a board-certified pathologist (AB).

129 **Total RNA extraction for sequencing**

130 Total RNA was extracted from frozen brain tissues as described previously ¹¹. Initially,
131 approximately 20 - 30 mg of tissue was snap-frozen in liquid nitrogen and disintegrated
132 using a cryoPREP impactor (Covaris, UK). The pulverized tissue was solubilized in pre-
133 heated lysis buffer AL and RNA was extracted using the RNAdvance Tissue Kit (Beckman
134 Coulter, Germany) in combination with a KingFisher Flex Purification System (Thermo
135 Fisher Scientific, Germany).

136 **RusV-specific RT-qPCR**

137 RusV-specific RNA was detected by TaqMan RT-qPCR using the AgPath-ID One-Step RT-PCR
138 reagents (Thermo Fisher Scientific, Germany) along with a modified primer/probe set
139 targeting the p200 ORF ². Briefly, 2.5 µl extracted RNA was reverse-transcribed and
140 amplified in a reaction mix of 12.5 µl total volume containing primers RusV_1072_A+
141 (5'-CGAGCGYGTCTACAAGTTYA-3'; final concentration 0.8 µM) and RusV_1237-
142 (5'-GACCATGATGTTGGCGAGG-3'; 0.8 µM) and probe RusV_1116_A_P
143 (5'-[FAM]CCGAGGARGACGCCCTGTGC[BHQ1]-3'; 0.4 µM). The reaction was performed with
144 the following cyclers setup: 45 °C for 10 min, 95 °C for 10 min, 45 cycles of 95 °C for 15
145 sec, 60 °C for 30 sec and 72 °C for 30 sec on a Bio-Rad CFX96 qPCR cycler (Bio-Rad,
146 Germany).

147 **Sequencing of total RNA**

148 Extracted total RNA was sequenced using a universal metagenomics sequencing
149 workflow ^{11, 12}. An amount of 350 ng total RNA per sample was reverse-transcribed into
150 cDNA using the SuperScript IV First-Strand cDNA Synthesis System (Invitrogen, Germany)
151 and the NEBNext Ultra II Non-Directional RNA Second Strand Synthesis Module (New
152 England Biolabs, Germany). Afterwards, cDNA was processed to generate Ion Torrent
153 compatible barcoded sequencing libraries as detailed described ^{2, 11}. Libraries were

154 quantified with the QIAseq Library Quant Assay Kit (Qiagen, Germany) and subsequently
155 sequenced on an Ion Torrent S5XL instrument using Ion 530 chips and chemistry for
156 400 base pair reads (Thermo Fisher Scientific, Germany).

157 **Sequencing of rRNA-depleted and poly(A)+ enriched RNA**

158 For rRNA depletion we used the NEBNext rRNA Depletion Kit for human, mouse, and rat
159 (New England Biolabs, USA) that specifically depletes cytoplasmic (5S, 5.8S, 18S and 28S
160 rRNA) and mitochondrial ribosomal RNA (12S and 16S rRNA). As the depletion is rRNA
161 sequence-specific, we first confirmed that the human-, mouse- and rat-specific panel would
162 be compatible with samples from yellow-necked field mice by comparing available
163 cytoplasmic and mitochondrial rRNA sequences of all species. Subsequently, 3 µg of the
164 total RNA from two selected yellow-necked field mice were treated with the NEBNext rRNA
165 Depletion Kit for human, mouse, and rat (New England Biolabs), following the
166 manufactures instructions.

167 Enrichment of poly(A)+ RNA from total RNA was considered appropriate, as the RusV
168 genome, like RuV¹³, comprises a poly(A) tail at the 3' terminus. For poly(A)+ enrichment,
169 3 µg of total RNA from the same yellow-necked field mice were treated with the Dynabeads
170 mRNA DIRECT Micro Purification Kit (Invitrogen, USA) following the manufacturer's
171 instructions.

172 Both, rRNA depleted and poly(A)+ enriched RNA, were used for strand-specific library
173 construction with the Colibri Stranded RNA Library Prep Kit (Thermo Fisher, USA). Libraries
174 were quality-checked using a 4150 TapeStation System (Agilent Technologies, USA) with
175 the High Sensitivity D1000 ScreenTape and reagents (Agilent Technologies) and were then
176 quantified using a Qubit Fluorometer (Thermo Fisher) along with the dsDNA HS Assay Kit
177 (Thermo Fisher). Libraries were pooled and sequenced on a NextSeq 500 (Illumina, USA)
178 using a NextSeq 500/550 Mid-output Kit v2.5 with 300 cycles (Illumina).

179 **Design of custom panRubi bait panels**

180 All available whole-genome sequences of the genus *Rubivirus* were received from NCBI
181 GenBank (86 RuV, one RuhV and three RusV sequences). The genome set was sent to
182 Daicel Arbor Biosciences (Ann Arbor, USA) and a tailored custom myBaits panel for target
183 enrichment via hybridization-based capture was designed. The resulting "panRubi" panel
184 consists of 19,178 RNA oligonucleotide baits with a length of 60 nt arranged every 20 nt
185 along the genomes (designated "panRubi bait set v1"). The set was later supplemented

186 with 22 additional baits covering the newly identified part of the capsid protein-encoding
187 sequence and IGR arranged every 16 nt. This set was mixed with the “panRubi bait set v1”
188 at a ratio of 1:10 to give the “panRubi bait set v2”. All bait sets were checked using Basic
189 Local Alignment Search Tool (BLAST) search against human, mouse, horse, and opossum
190 genomes and no BLAST hit was found.

191 **Application of RNA baits and sequencing**

192 The custom panRubi bait sets v1/v2 were applied to the sequencing libraries according to
193 the manufacturer’s instructions (myBaits manual v.5.00, Arbor Biosciences, Sep 2020).
194 Hybridization reactions were performed in 1.5 µl safe-lock tubes overlaid with one volume
195 of mineral oil (Carl Roth, Germany), to keep the volume constant during hybridization using
196 a ThermoMixer (Eppendorf, Germany) with 550 rotations per minute. We used the standard
197 protocol (according to ¹⁴) with a hybridization temperature of 65 °C and a hybridization time
198 of about 24 hours. The enriched and purified samples were amplified using the GeneRead
199 DNA Library L amplification Kit (Qiagen, Germany) according to manufacturer’s instructions
200 with 14 cycles and amplicons were purified using solid-phase paramagnetic bead
201 technology. Treated libraries were sequenced after quality check using a Bioanalyzer 2100
202 (Agilent Technologies) and quantification as described above.

203 **Read processing and *de novo* assembly**

204 Ion Torrent-derived reads from the myBaits capture enrichment approach were initially
205 quality-trimmed and specific adapters were removed using the 454 Sequencing Systems
206 Software (version 3.0). Instead of host/background removal using specific reference
207 sequences, a G+C content filter was applied to the trimmed reads, as the RusV genome
208 has a particularly high average G+C content of 70.6 mol% ². In detail, only reads with an
209 average G+C content of ≥60 mol% were filtered using PRINSEQ-lite (version 0.20.4; ¹⁵) and
210 subsequently used for *de novo* assembly with SPAdes genome assembler (version
211 3.15.2; ¹⁶) running in single cell mode (–sc) for Ion Torrent data (–iontorrent). The resulting
212 contigs were mapped to the RusV reference sequence MN552442 using Geneious generic
213 mapper (Geneious Prime 2021.0.1) with medium sensitivity allowing discovery of structural
214 variants, and short insertions/deletions (indels) of any size. A consensus sequence was
215 generated and reads were finally mapped back to the consensus sequence using Geneious
216 generic mapper in order to manually inspect genomic termini and possible frameshifts
217 caused by homopolymers.

218 Illumina-derived reads from rRNA-depleted and poly(A)+-enriched RNA were initially
219 trimmed using Trim Galore (version 0.6.6; ¹⁷) with automated adapter selection and reads
220 containing only poly(A) homopolymers were trimmed using BBDuk/BBDuk
221 (version 38.18, ¹⁸). For coverage analysis, the trimmed reads of each sample were mapped
222 to the respective assembled genome using Geneious Prime generic mapper in “Low
223 Sensitivity / Fastest” mode. The indexed BAM files were then processed with SAMtools
224 depth (version 1.11; ¹⁹).

225 **Phylogenetic analysis**

226 Complete RusV genome sequences were aligned using MAFFT (version 7.450; ²⁰) and then
227 used as input for approximately-maximum-likelihood reconstruction with Fast Tree
228 (version 2.1.11; ²¹) using the generalized time-reversible (GTR) model with 5 rate
229 categories and optimized Gamma20 likelihood. The resulting tree was inspected using
230 Geneious Prime (version 2021.0.1).

231

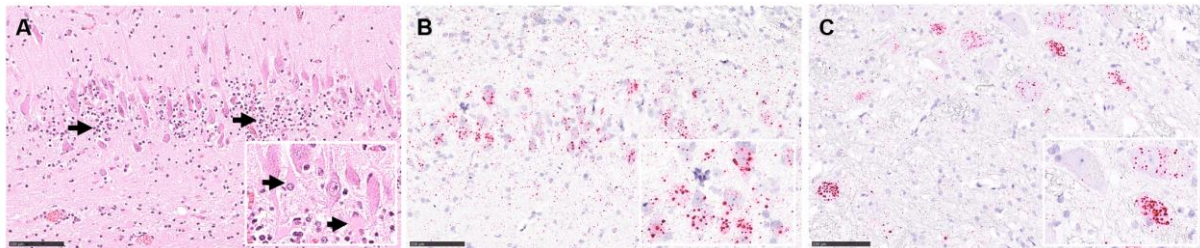
232 **Results**

233 **Two carnivoran mammals with neurological disorder**

234 In August 2020, a South American coati (*Nasua nasua*) kept in the zoo showed lethargy,
235 hind limb weakness, convulsion and tremor. Two days later and finally unmoving, the
236 animal was euthanized. Gross pathology revealed swelling of the liver and hyperkeratosis
237 of the footpads. Initial histopathology identified a non-suppurative meningoencephalitis.
238 Findings in the liver included scattered single cell necrosis of hepatocytes and minimal
239 microvesicular fatty change interpreted to be clinically irrelevant, while the footpad
240 hyperkeratosis was interpreted to be age-related. Standard diagnostic tests were negative
241 for mammalian bornaviruses, canine distemper virus and *Salmonella* spp.

242 In December 2020, a wild Eurasian otter (*Lutra lutra*) was found in the vicinity of the very
243 same zoological garden, without any reported link to the zoo areal, showing abnormal
244 movement. Prior to capturing, the animal was observed in the open waters of the nearby
245 Baltic Sea coast and then later found on the premises of a local school. The animal was
246 sent for clinical observation to the zoo, presenting in a state of malnourishment but with
247 increased food and water uptake, loss of natural shyness and an abrasion at the head
248 indicating blunt trauma. Abnormal movements were still present until the animal was found
249 dead three days later. Pathological examination confirmed hairless spots at the head with

250 a focal perforation of the skin but otherwise non-specific alterations interpreted to be
251 associated with acute, agonal cardiovascular failure. Initial routine histology identified a
252 non-suppurative meningoencephalitis but no further lesions in other organs. Standard
253 diagnostic tests were negative for mammalian bornaviruses, influenza A virus, canine
254 distemper virus, rabies virus, *Salmonella* spp. and *Toxoplasma gondii*.



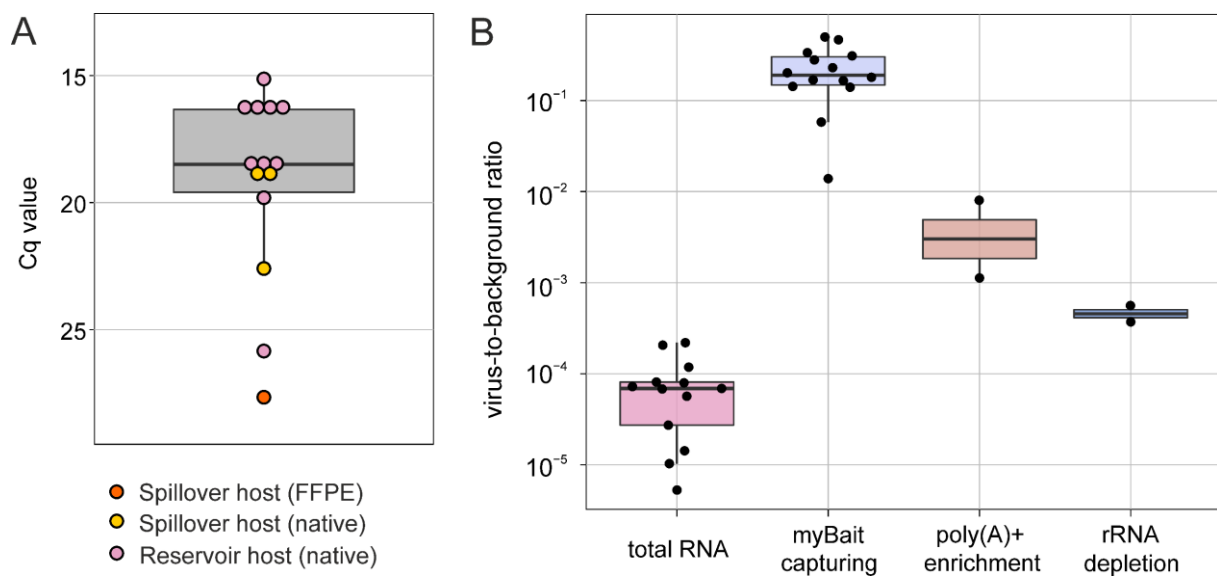
255
256 **Fig. 1:** Histopathology from cases of rustrela virus (RusV)-associated meningoencephalitis in a
257 Eurasian otter (*Lutra lutra*) and a South American coati (*Nasua nasua*). (A) Non-suppurative
258 meningoencephalitis in the hippocampus region of the otter, with mononuclear infiltrates
259 (arrows) and loss of Nissl substance indicating neuronal degeneration (inlay with arrows), HE
260 stain. Detection of RusV RNA in neurons of the hippocampus region of the otter (B and inlay) and
261 brain stem of the coati (C and inlay). RNA ISH, chromogenic labelling (fast red) with probes to
262 RusV non-structural polyprotein encoding region, Mayer's hematoxylin counter stain. Scale bar
263 100 μ m.

264 Histopathology confirms RusV-associated encephalitis

265 In general, follow-up histopathology of the RusV-infected South American coati and
266 Eurasian otter confirmed our results previously reported for the RusV-infected donkey,
267 capybara and wallaby from the same zoo. Associated with a non-suppurative
268 meningoencephalitis (Fig. 1A), RNA ISH confirmed the presence of RusV-specific RNA
269 within neuronal cell bodies and their processes in both animals (Fig. 1B and C). Routine HE
270 staining (Fig. 1A) as well as Luxol fast blue Cresyl violet staining (Supplemental Fig. S1A)
271 identified neuronal degeneration in the brain of the Eurasian otter but not the South
272 American coati. Scattered cells, in particular perivascularly, were active caspase 3-labelled,
273 indicating subtle apoptosis induction (Supplemental Fig. S1B). Multifocal perivascular cells
274 in brain samples from the otter were positive for iron in the Prussian Blue reaction,
275 confirming intravital hemorrhages (Supplemental Fig. S1C), potentially associated with a
276 suspected history of a blunt trauma. The non-suppurative meningoencephalitis was
277 characterized by perivascular and disseminated infiltrates and few microglial nodules.
278 Immunohistochemistry identified mainly infiltrating CD3-positive T-cells (Supplemental
279 Fig. S1D) but only single CD79-labelled B-cells (Supplemental Fig. S1D inlay). Numerous
280 IBA1-positive microglial cells and infiltrating macrophages were detected intralésionally
281 (Supplemental Fig. S1E). In addition, GFAP immunohistochemistry indicated activation of
282 astrocytes, exhibiting a plump cell shape (Supplemental Fig. S1F).

283 RT-qPCR confirms presence of RusV in encephalitic animals and reservoir hosts

284 A RusV-specific RT-qPCR confirmed the presence of viral RNA in the brain of the South
285 American coati (Cq 18.9) and the Eurasian otter (Cq 22.5). Furthermore, using the same
286 RT-qPCR setup, we also reanalysed samples from two previously investigated zoo animals
287 and from ten previously published or recently collected RusV-infected wild yellow-necked
288 field mice from within and around the zoo (**Table 1**). Cq values of frozen brain samples
289 ranged from 15.1 to 25.8, with a median of 18.4 (**Fig. 2A**), whereas FFPE brain tissue from
290 a capybara with encephalitis revealed the highest Cq of 27.6 corresponding to the lowest
291 amount of detectable RNA.



292

293 **Fig. 2: (A)** Comparison of RusV-specific RT-qPCR Cq values for frozen or FFPE brain samples from
294 potential reservoir (yellow-necked field mouse) and spill-over hosts (donkey, capybara, South
295 American coati, European otter). **(B)** Comparison of virus-to-background sequence ratio observed
296 in sequencing data using different RNA preparations and post library capturing methods.

297 Increasing RusV sequencing efficiency

298 Initially, we used total RNA from brain samples of the Eurasian otter and South American
299 coati for sequencing and *de novo* assembly. However, this resulted in incomplete and
300 highly fragmented genome sequences due to very low virus-to-background sequence ratios
301 of 0.021% and 0.001% for the South American coati and Eurasian otter, respectively. The
302 virus-to-background ratios observed during sequencing of total RNA from all samples
303 included in this study (**Table 1**) ranged from 0.00053% to 0.022%, with a median of
304 0.0069% (**Fig. 2B**). To increase the efficiency of RusV sequencing, we compared poly(A)+
305 enrichment, rRNA depletion as well as post library hybridization-based capturing (bait
306 capturing) for selected samples.

307 By reducing host-derived RNA, poly(A)+ enrichment and rRNA depletion increased virus-to-
308 background sequence ratios by factors of 67 and 6.8, respectively, as compared to total
309 RNA, resulting in median virus-to-background sequence ratios of 0.46% and 0.047%,
310 respectively (**Fig. 2B**). The application of bait capturing to libraries prepared from total RNA
311 achieved virus-to-background sequence ratios of 1.4% to 49.9% with a median of 19.1%
312 (**Fig. 2B**), corresponding to a median 2,772-fold increase.

313 The characteristic sequence coverage pattern observed for bait-captured libraries closely
314 resembled that of total RNA sequencing (Supplemental Fig. S2). In contrast, libraries from
315 poly(A)+ enriched RNA had a strong bias in coverage towards the 3' end of the RusV
316 genome. Depletion of rRNA resulted in a relatively uniform coverage across the genome
317 with a bias towards the 5' end of the genome (Supplemental Fig. S2). No coverage dropout
318 was noted for any of the applied methods.

319 **Generation and comparison of full length RusV genomes**

320 As bait capturing proved to be very efficient, we applied the technique to all 14 available
321 brain samples (**Table 1**), including South American coati and Eurasian otter, and used the
322 sequencing data for *de novo* assembly. The assembly of each library resulted in contigs
323 that were matched to the RusV genome MN552442. For all samples, a full-length RusV
324 genome without any gaps could be derived from the matched *de novo* assembled contigs.

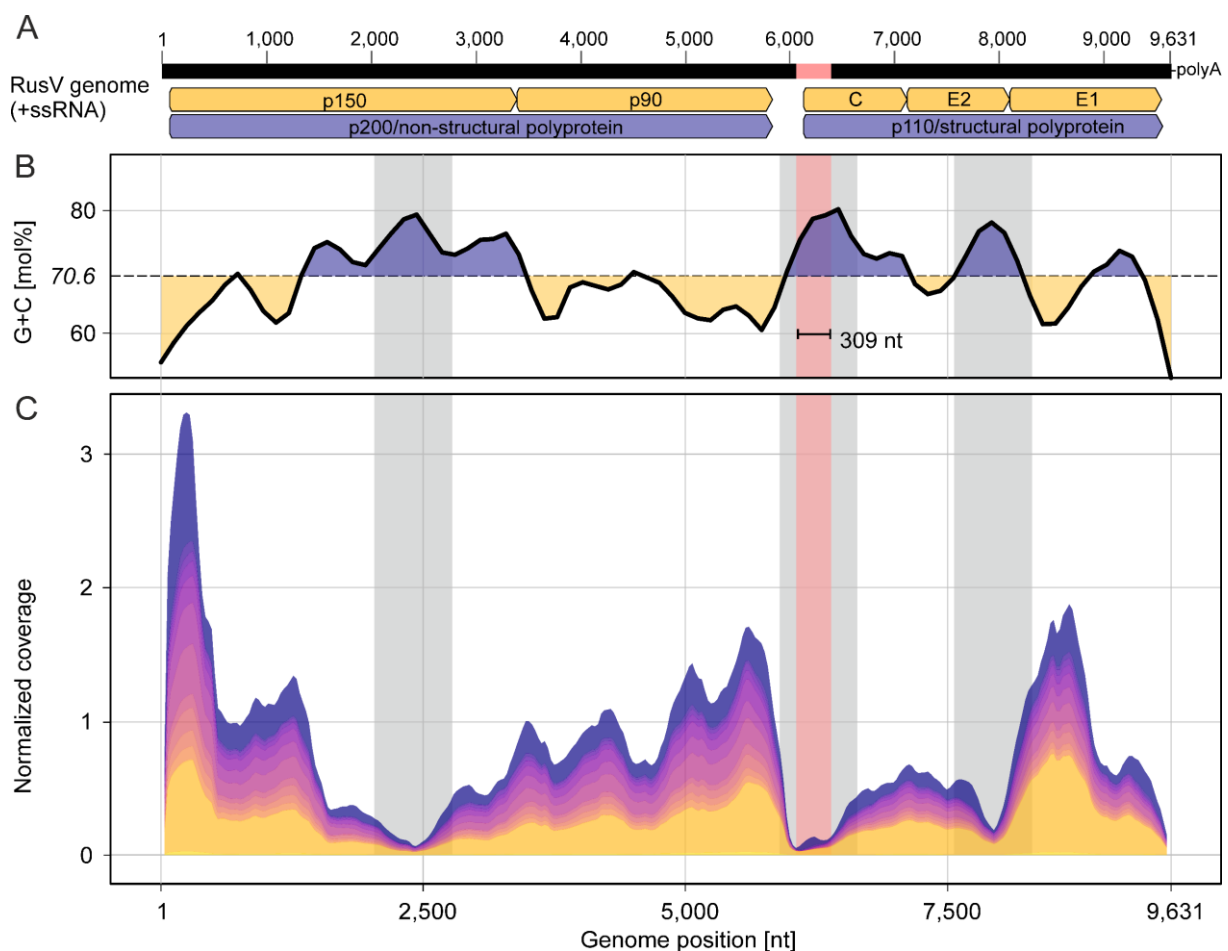
325 An alignment of all 14 RusV genome sequences showed only minor variation with an overall
326 pairwise nt identity of 98.8%. Phylogeny based on the aligned whole genome sequences
327 confirmed the high genetic identity of the RusV genomes originating from within or in close
328 proximity of the zoo (Supplemental Fig. S3). RusV sequences from apparently healthy
329 yellow-necked field mice and encephalitic mammals, including the South American coati
330 and Eurasian otter, clustered closely together. Notably, two RusV sequences from yellow-
331 necked field mice collected in a distance of about 10 km from the zoo grouped in a
332 separate genetic branch (Supplemental Fig. S3).

333 **Revised RusV genome and implications for the IGR and capsid protein-coding sequence**

334 All 14 RusV genomes assembled from bait-captured libraries showed a 309 nt stretch
335 ranging from pos. 6,062 to pos. 6,370 and covering part of the IGR and the N-terminal part
336 of the capsid protein-coding sequence. This stretch had not been present in the three
337 initially released RusV genomes generated by total RNA sequencing² but was now
338 identified when re-sequencing the very same sample materials using bait capturing

339 (Fig. 3A). As the panRubi v1 bait set did not comprise the extra 309 nt-long region, the bait
340 set was complemented with probes specifically targeting this region, leading to a further
341 improved coverage within the respective region (Supplemental Fig. S4).

342 In general, the observed sequencing coverage varied considerably across the genome,
343 showing pronounced maxima and minima in all samples (Fig. 3C). The three genomic
344 regions with the most prominent reduction in sequence coverage correlated with the
345 highest G+C content (Fig. 3B and C), while genome regions with very high coverage
346 correlated with lower G+C content. The newly identified 309 nt region correlated with a
347 G+C peak and possessed a particularly low sequence coverage (Fig. 3A-C).



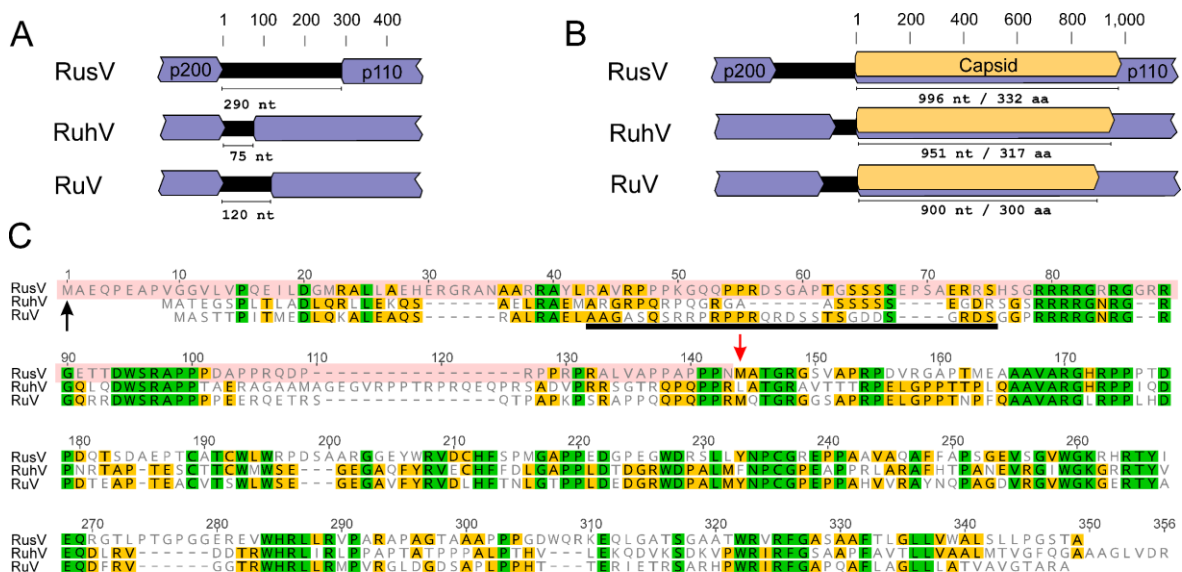
348

349 **Fig. 3:** Schematic rustrela virus (RusV) genome sequence (A) showing averaged G+C content (B)
350 and cumulated RusV sequence coverage of all 14 animals included in this study (C). The newly
351 identified 309 nt sequence stretch partly covering the intergenic region and p110 ORF is
352 highlighted in red. Note that the start of the p110 coding ORF is located within the newly identified
353 sequence stretch, leading to a longer capsid protein sequence as compared to the previously
354 published RusV genomes. Grey labelled areas in B and C indicate areas of particularly high G+C
355 content.

356

357 As a consequence, the IGR and the predicted p110 ORF are longer than initially reported
 358 (Fig. 4A and B). The IGR of all 14 full-length RusV sequences is spanning 290 nt between
 359 the stop codon of the predicted p200 ORF and start codon of the p110 ORF. In comparison,
 360 the IGRs of RuhV and RuV were reported to be 75 nt and 120 nt in length, respectively
 361 (Fig. 4A). Based on an ATG start codon in the newly identified 309 nt region and a prediction
 362 of the signal peptidase cleavage site (Supplemental Fig. S5), the predicted capsid protein-
 363 encoding sequence of RusV is 996 nt (332 amino acids, aa) in length. In comparison, the
 364 length of the capsid protein-encoding regions of RuhV and RuV was predicted to be 951 nt
 365 (317 aa) and 900 nt (300 aa), respectively (Fig. 4B).

366 Comparison of the revised capsid protein sequence of RusV to both RuhV and RuV revealed
 367 highly conserved stretches (Fig. 4C). The revised RusV capsid protein sequence comprised
 368 a part that has been predicted to be the RBD in RuV. This part had been absent in the
 369 initially published RusV genome. Directly downstream of the predicted RuV RBD region, a
 370 polybasic motif (RRRRG R/N RG) can be found that is highly conserved between RusV,
 371 RuhV and RuV. This polybasic motif is followed by a likewise highly conserved hydrophobic
 372 motif (DWSRAPP).



373

374 **Fig. 4:** Comparison of the rubivirus intergenic region and capsid protein-encoding sequences. (A)
 375 The size of the intergenic region between the non-structural (p200) and structural polyprotein
 376 (p110) ORFs of rustrela virus (RusV; MN552442.2), ruhugu virus (RuhV; MN547623) and rubella
 377 virus (RuV; NC_001545) is shown. (B) The predicted length of the capsid protein-coding sequence
 378 (highlighted in orange) is shown for RusV, RuhV and RuV. (C) The sequences of the capsid protein
 379 from RusV, RuhV and RuV are compared using an amino acid sequence alignment. Amino acid
 380 residues highlighted in green or yellow are conserved in all three or at least in two of the viruses,
 381 respectively. The N-terminal part of the RusV mature capsid protein (highlighted in red; start
 382 marked by black arrow) has been determined in this study. The red arrow indicates the predicted
 383 start of the capsid protein in the previously published RusV sequence. The RNA binding site of the
 384 RuV capsid protein is indicated by the black bar.

385 Discussion

386 We recently discovered RusV in the central nervous system of encephalitic zoo animals and
387 wild yellow-necked field mice on the basis of metagenomic sequencing of total RNA, RT-
388 qPCR and RNA ISH ². More than 300 million reads from different sequencing platforms and
389 numerous samples and subsamples were used in order to generate the first RusV genomes
390 originating from three individuals (MN552442.1, MT274724.1 and MT274725.1). A
391 combination of *de novo* assembly, mapping, BLASTx, and manual inspection was used to
392 generate these RusV genomes ². The assembly was exceptionally difficult, as most parts of
393 the RusV genome have a G+C content of >70 mol% with low complexity G+C stretches and
394 coverage dropping drastically at several positions (**Fig. 3B** and **C**). In the IGR, the G+C
395 content even exceeds 85 mol%. Nevertheless, despite very large efforts in sequence
396 determination and characterisation of the RusV genome, questions remained regarding its
397 unusual IGR and capsid protein, which appeared to be rather short in comparison to RuV
398 and RusV and lacked a potential RBD ¹⁰.

399 We now investigated two new cases of RusV-associated fatal meningoencephalitis in a
400 European otter and a South American coati that clinically and histologically closely
401 resembled previous RusV-associated cases and that further broadened the spectrum of
402 infected mammals, which now includes placental mammals of the orders Rodentia
403 (families Caviidae and Muridae), Carnivora (families Procynoidae and Mustelidae), and
404 Perissodactyla (family Equidae) as well as marsupials of the order Diprotodontia (family
405 Macropodidae). This broad host spectrum is in clear contrast to rubella virus, for which
406 humans are the only host ⁸. So far, we assume that the wild yellow-necked field mouse may
407 act as reservoir host. However, the transmission route to the other hosts remains unclear.

408 Despite the relatively low RusV-specific RT-qPCR C_q values in native organ samples,
409 detected using a RusV-specific RT-qPCR (**Fig. 2A**), the virus-to-host sequence ratio observed
410 when sequencing total RNA was unsatisfying. It has been shown, that genome length, virus
411 species, virus- and host-derived RNA concentrations as well as the overall composition of
412 the sample matrix impact the virus-background ratio ²². RT-qPCR results often do not reflect
413 this complex interplay and may lead to false expectations for sequencing.

414 As sequencing of RusV genomes from total RNA proved to be very difficult, we attempted
415 to increase the sequencing efficiency by poly(A)⁺ enrichment, rRNA depletion and post-
416 library bait capturing. Poly(A)⁺ enrichment was more efficient than rRNA depletion,
417 resulting in higher virus-to-background ratios (**Fig. 2B**). This observation is in accordance

418 with other studies, in which rRNA-depleted RNA preparations contained many host-derived
419 small or long non-coding RNAs that are absent in poly(A)+ enriched RNA preparations ²³.
420 However, poly(A)+ selection introduced a 3' sequence coverage bias resulting in poor
421 5' coverage. This bias has been reported previously for poly(A)+ enrichment methods and
422 is most likely caused by partially degraded transcripts particularly in samples with highly
423 degraded RNA ²⁴. RNA quality and integrity plays a major role in sequencing experiments
424 and is directly connected to sampling conditions, transportation and storage ²⁵⁻²⁷. While
425 we used qualified and robust methods for RNA extraction and preservation ¹¹, the RNA
426 preparations used for comparison of the different methods originated from brain tissues
427 of two wild-trapped rodents that were sampled under suboptimal conditions.

428 Hybridization-based capturing has been shown to markedly increase efficiency of RNA virus
429 sequencing previously ^{14, 28-30} and was also found to be most efficient in this study,
430 increasing the median virus-to-background sequence ratio 2,772-fold. Using this
431 technique, we sequenced or re-sequenced 14 full-length RusV genomes from cases of
432 encephalitis and from wild yellow-necked field mice. Using the hybridization-based bait
433 capturing method, the overall sequence coverage and especially the coverage in
434 challenging regions was markedly improved. However, we found a correlation between
435 sharp drops in sequencing coverage within regions of very high G+C content exceeding
436 ~75 mol%. This may indicate a technical limit of the used sequencing platforms and has
437 been described for different technologies ³¹⁻³⁴. It has also been suggested, that extreme
438 G+C contents may negatively affect *de novo* assemblies ³⁵.

439 Within a region of high G+C content, spanning IGR and the 5'-end of the capsid protein-
440 encoding sequence, we now found a notable sequence difference, namely a previously
441 unidentified stretch of 309 nt, in comparison to the initially reported RusV genome.
442 Thereby, the predicted capsid protein of RusV is longer than described earlier and now
443 includes the typical rubivirus capsid protein features, such as the RBD that might be crucial
444 for virion formation ^{6, 7}. The RBD was unexpectedly missing in the initially annotated RusV
445 capsid protein as pointed out recently in detail by Das and Kilian ¹⁰. An alignment of the
446 capsid proteins of RusV, RuhV and RuV showed highly conserved motifs that were initially
447 absent in the predicted RusV protein. However, the region identified as RBD in RuV ⁶
448 appears to be only poorly conserved on aa sequence level in the RuhV or RusV when
449 compared in an alignment. Whether conserved motifs are involved in RNA binding or other
450 structural features remains unclear, as no structural model is currently available for the N-
451 terminal part of the RuV capsid protein ⁷.

452 The revised version of the RusV genome reveals RusV to have the longest capsid protein-
453 encoding sequence and IGR among all three currently known matonaviruses (**Fig. 2A** and
454 **B**). It has been shown for RuV, that the p110 polyprotein is translated from a subgenomic
455 RNA by using a separate promoter within the IGR^{36, 37}. However, based on the coverage
456 along the genome, we could not find any indication for the presence of subgenomic RNA in
457 the analysed samples. This may indicate that either RusV does not translate the p110
458 polyprotein from a subgenomic RNA or the RusV replication cycle includes stages without
459 presence of subgenomic RNA. Future studies should address these open questions.

460 In conclusion, we were able to markedly increase RusV sequencing efficiency leading to an
461 improved genome coverage by employing a bait capturing-based enrichment strategy.
462 Overall, 14 high-quality whole-genomes from RusV-related encephalitis cases and reservoir
463 hosts could be generated applying this strategy. By *de novo* assembly, we identified an
464 extra 309 nt sequence spanning the partial RusV IGR and 5'-end of the capsid protein-
465 encoding region. The RusV example impressively demonstrates the difficulties in correctly
466 determining sequences with an extreme G+C content, but also suggests possible solutions
467 that are now available, such as targeted enrichment via RNA baits. The updated RusV
468 sequence now allows further studies about the function of conserved regions of RusV, but
469 also about viral replication using reverse genetics.

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474 **Data availability**

475 Revised versions of previously published RusV genome sequences are available under
476 DDBJ/ENA/GenBank accession numbers MN552442.2, MT274724.2, and MT274725.2.
477 Novel RusV genomes from this study are available under DDBJ/ENA/GenBank accession
478 numbers: OL960716 - OL960726

479 **Ethics statement**

480 This study involved no animal experiments. All animal materials were from routine
481 diagnostics or pest rodent control measures.

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486 Competing interests

487 The authors declare no competing interests.

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489 **Conceptualization:** DR, MB, RGU

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497

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606 **Table 1:** Rustrela virus-infected zoo animals, Eurasian otter and yellow-necked field mice from Northern Germany included in this study

Strain	Organism	Sampling date	Location	Study	Sequence data#	Previous accessions	Updated and new full genome accession
Yellow-necked field mouse/Mu09-1341/2009/Germany	<i>Apodemus flavicollis</i>	Jul 2009	~2 km distance to zoo	Bennet <i>et al.</i> 2020	T, B	MT274737.1, MT274731.1	OL960721
Donkey/19_041-1/2019/Germany	<i>Equus asinus</i>	Mar 2019	housed in zoo	Bennet <i>et al.</i> 2020	T, B	MN552442.1	MN552442.2
Capybara/P19-643/2019/Germany	<i>Hydrochoerus hydrochaeris</i>	Oct 2019	housed in zoo	Bennet <i>et al.</i> 2020	T, B, B+	MT274724.1	MT274724.2
Yellow-necked field mouse/KS19-928/2019/Germany	<i>Apodemus flavicollis</i>	Sep 2019	on zoo grounds	Bennet <i>et al.</i> 2020	T, B, B+	MT274725.1	MT274725.2
Yellow-necked field mouse/KS20-1296/2020/Germany	<i>Apodemus flavicollis</i>	Oct 2020	~10 km distance to zoo	Bennet <i>et al.</i> 2020	T, B	MT274732.1, MT274726.1	OL960722
Yellow-necked field mouse/KS20-1340/2020/Germany	<i>Apodemus flavicollis</i>	2020	on zoo grounds	Bennet <i>et al.</i> 2020	T, B	MT274733.1, MT274727.1	OL960726
Yellow-necked field mouse/KS20-1341/2020/Germany	<i>Apodemus flavicollis</i>	2020	on zoo grounds	Bennet <i>et al.</i> 2020	T, B	MT274734.1, MT274728.1	OL960725
Yellow-necked field mouse/KS20-1342/2020/Germany	<i>Apodemus flavicollis</i>	2020	on zoo grounds	Bennet <i>et al.</i> 2020	T, B, B+	MT274735.1, MT274729.1	OL960724
Yellow-necked field mouse/KS20-1343/2020/Germany	<i>Apodemus flavicollis</i>	2020	on zoo grounds	Bennet <i>et al.</i> 2020	T, B, R, P	MT274736.1, MT274730.1	OL960723
Yellow-necked field mouse/KS20-1512/2020/Germany	<i>Apodemus flavicollis</i>	2020	on zoo grounds	this study	T, B, R, P	n.a.	OL960720
Yellow-necked field mouse/KS20-1513/2020/Germany	<i>Apodemus flavicollis</i>	2020	on zoo grounds	this study	T, B	n.a.	OL960719
Yellow-necked field mouse/KS20-1535/2020/Germany	<i>Apodemus flavicollis</i>	Jun 2020	~10 km distance to zoo	this study	T, B	n.a.	OL960718
South American coati/20_131/2020/Germany	<i>Nasua nasua</i>	Aug 2020	housed in zoo	this study	T, B, B+	n.a.	OL960717
Eurasian otter/21_002/2020/Germany	<i>Lutra lutra</i>	Dec 2020	~3 km distance to zoo	this study	T, B, B+	n.a.	OL960716

607 #T: total RNA, B: initial panRubi myBait set v1, B+ modified pabRubi myBait set v2, P: polyA selected RNA, R: rRNA depleted RNA; n.a., not applicable