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Version 2

# Rabies Virus Bat-Clade Sequencing V.2

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We use this protocol and it's working

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## Disclaimer

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#### Abstract

Join us in advancing global genomic surveillance of the rabies virus.

We are actively engaged in pioneering a pan-clade whole genome amplicon sequencing protocol for the rabies virus. If you're interested in collaborating on this crucial endeavor and wish to obtain a primer aliquot from our established protocol or the one currently in development, please do not hesitate to contact us at richardssalvato@gmail.com.

#### **Background**

Rabies virus (RABV) causes fatal encephalitis in domestic animals and humans. This high-impact zoonotic virus is poorly studied from the genomic perspective so establishing genomic surveillance protocols for RABV is crucial to track their evolution and monitor spillover events and wildlife reservoirs.

Despite the importance of RABV zoonotic cycle, there is limited complete genome sequence data of RABVs from some hosts as bats. We developed a novel rapidly deployable, cost-effective, and flexible ampliconbased sequencing approach usable with protocols widely established during the COVID-19 pandemic and suitable to different hosts based on a one-health context. We used PrimalScheme to generate a primers panel and then aligned them with a RABV sequences dataset from different species and manually degenerated the primers to cover a wider diversity of hosts.

The set of 47 primers is compatible and ready to use with COVIDSeq sequencing protocol and Illumina DNA Prep, and allows to sequence up to 384 samples per run on the Illumina MiSeq system or to accommodate in libraries with other sample types. Additionally, we included primers for amplification of a fragment of the mitochondrial gene COI to host species identification.

## COI

#### Initial Validation

In the initial validation, we sequenced 160 complete RABV genomes from different species from five distinct families (Bovine, Equine, Caprine, Felines, Microchiroptera) with an average coverage of 98%, most of them recovering the whole genome (88/160).

#### **Conclusions**

Here, we introduced a cost-effective and easy-to-use sequencing protocol for RABV bat-clade in order to support genomic surveillance of a re-emerging zoonotic disease allowing targeting viral control programs and adequate public health policies.

## **Acknowledgements**



We would like to thank *Instituto de Pesquisas Veterinárias Desidério Finamor* for sample processing and initial rabies virus diagnosis.



## **Materials**

A	В	С	D
primer	sequence	pool	
raiva_1_LEFT	ATGTGGAAGGRARTTGGGCTCT	1	
raiva_1_RIGHT	TTGACKGTTCCGTCATCTGCC	1	
raiva2_LEFT	CATGAGATGTCWGTTCTTGGRGG	2	
raiva2_RIGHT	GTGACATAGGATATGATCTCCTCRAC	2	
raiva_3_LEFT	AGATTTTTGTCARYCCAAGTGCG	1	
raiva_3_RIGHT	CATCTCAGAGGGAGYTTGGATCC	1	
raiva4_LEFT	TGAARATGAACCTTGAYGACATAGTC	2	
raiva4_RIGHT	CATGTTRATACACCAAATYCTKCC	2	
raiva5_LEFT	GAACTGGGTATAYAARTTGAGGAGAAC	1	
raiva5_RIGHT	CTTGAATGTGGTRGTGACATAACC	1	
raiva6_LEFT	AAGGTGGGRTACATMTCYGCCA	2	
raiva6_RIGHT	TATGCCTTYCCAAAYCCMGG	2	
raiva7_LEFT	ACYGTAAARACCACYAARGAGT	1	
raiva7_RIGHT	ACAGARGACTCYARCAGCTCCA	1	
raiva8_LEFT	TRATGAKTGCAGGTGSTCTGG	2	
raiva8_RIGHT	CTTAAGATRTTGGGRAYGGYGGG	2	
raiva9_LEFT	RACAGACAAYTGYTCKAGGTCWTACA	1	
raiva9_RIGHT	CCTCMAGYTGACYCACCTTRTCYC	1	
raiva10_LEFT	CCTTGGAYTGGGATGARGAGAA	2	
raiva10_RIGHT	TGTTTGGGAGGCCAYGTYTG	2	
raiva11_LEFT	CYAAATGGTATCTTGATYCGCGAC	1	
raiva11_RIGHT	YTGACATGTCCAGACARTATATTTGATC	1	
raiva_12_LEFT	TCTGTACTYGATCAAGTGTTYGGA	2	



А	В	С	D
raiva_12_RIGHT	GARAACTGACGTATRTGGAACCTT	2	
raiva_13_LEFT	GCTGTMTTCCATTACYTGCTST	1	
raiva_13_RIGHT	CCCAACATYTCAGAGGGRTGRG	1	
raiva_14_LEFT	SATGACMCAGACTCCCCAAAGG	2	
raiva_14_RIGHT	GCACDGCTCCMGAAACCATTCTA	2	
raiva_15_LEFT	TTGGCATCTTMGATGTAACAAGTG	1	
raiva_15_RIGHT	GARCTCATTTGTCKYAAGTTGG	1	
raiva_16_LEFT	TCWGACTTTAGAAGYTCYAAGATGAC	2	
raiva_16_RIGHT	GTGACCTCHGCATCACAAATGA	2	
raiva_17_LEFT	TGATGGCRTCAGGRACACAYC	1	
raiva_17_RIGHT	CTGCAGCATATGTTGAAGTGTCTC	1	
raiva_18_LEFT	GYTDATGTCTGATTTTGCAYTRTC	2	
raiva_18_RIGHT	TCARCCTGATCCAGTGAGAWGA	2	
raiva_extra_1_LEFT	ACGCTTAACRACAAAATCAG	2	
raiva_extra_1_RIGHT	ATGTTTGTCTTGTAATTGCC	2	
raiva_extra_2_LEFT	ATATTCAACAAGACYTTRAT	2	
raiva_extra_2_RIGHT	GTACAACTCCCATGARGATA	2	
raiva_extra_3_LEFT	CGAYTTGCCTCCTATGAAGG	1	
raiva_extra_3_RIGHT	AGCCAAAGGGAGATCATMGA	1	
raiva_extra_4_LEFT	TACAACAGACCCATAACYTA	1	
raiva_extra_4_RIGHT	ACGCTTAACAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	1	
raiva_extra_5_RIGHT	ACTTGGAACGAGATCATCCC	2	
raiva_extra_5_LEFTA	CCTATGAAGGACACAAGCAA	2	
raiva_extra_5_LEFTB	CCTATGAAGGACCCTAGCAA	2	
Mod_RepCOI_F	TNTTYTCMACYAACCACAAAGA	1	
Mod_RepCOI_R	TTCDGGRTGNCCRAARAATCA	2	
VertCOI_7194_F	CGMATRAAYAAYATRAGCTTCTGAY	2	



A	В	С	D
VertCOI_7216_R	CARAAGCTYATGTTRTTYATDCG	1	

Primers sequences and respective pools

# Troubleshooting



## **Primer Preparation**

- 1 Reconstitute each primer shown in Table 1 (See Materials section), using nuclease-free water to get a 100  $\mu$ M stock solution.
- 2 Prepare RABV-BAT primer pools A and B as described here.
- 2.1 Separate all primers at 100  $\mu$ M into two separate boxes labeled as Pool A and Pool B, according to Table 1.
- 2.2 Label a 2.0 ml microtube as Pool A and another as Pool B.
- 2.3 Vortex and spin down all the primers tubes.

2.4 Add  $\perp$  1080  $\mu$ L of nuclease-free water into the tube with Pool A. Add  $\perp$  1215  $\mu$ L of nuclease-free water into the tube with Pool B.

Now you have Pool A and B at a concentration of 10  $\mu M$  and ready to use.

2.5 Pooled Primers Can Be Stored at 4 -20 °C

## Sample Extraction and Cycle threshold (Ct) determination

28m 45s

- 3 Samples were extracted as described below and we used a previously published Realtime RT-PCR on all samples to determine viral load with Cycle threshold (Ct) value. Samples with a Ct value <28 are recommended for optimal results.
- 3.1 A total of  $\[ \underline{A} \]$  200  $\[ \mu L \]$  of homogenate samples, containing  $\[ \underline{A} \]$  50 mg of brain tissue fragments macerated in Basal Medium Eagle (BME), were added to  $\[ \underline{A} \]$  800  $\[ \mu L \]$  of



TRIzoITM Reagent (Invitrogen, Grand Island, NY, USA) in 2.0 ml microtubes.

- 3.2 The tissue was disrupted using a vortex at maximum speed for  $\bigcirc$  00:00:15 , followed by Room temperature incubation for  $\bigcirc$  00:05:00 and then  $\bot$  180  $\mu$ L of chloroform was added.
- 3.3 The tissue was disrupted using a vortex at maximum speed for 00:00:15, followed 5m 15s by Room temperature incubation for 00:05:00 and then Δ 180 μL of chloroform was added.
- 3.4 The mixture was mixed for  $\bigcirc$  00:00:15 , incubated at  $\blacksquare$  Room temperature for  $\bigcirc$  00:03:00 , and centrifuged at  $\bigcirc$  12000 rcf at  $\blacksquare$  4 °C for  $\bigcirc$  00:15:00 .
- 3.5 Viral RNA extraction was carried out using Δ 200 μL of supernatant using a commercially available Extracta Kit Fast DNA e RNA Viral (MVXA-PV96-B FAST) and the Loccus Extracta 96 equipment (Loccus, Sao Paulo, Brazil) following the manufacturer's instructions. Extracted RNA samples were stored at –80°C until tested by RT-qPCR.
- 3.6 For Cycle threshold (Ct) determination TaqMan single-step RT-PCR assay was conducted according previously described by Crystal Gigante (See protocol below).

# NAME LN34 pan-lyssavirus real-time RT-PCR for post-mortem diagnosis of rabies in animals CREATED BY Crystal Gigante Preview

4 cDNA synthesis and amplification of amplicons can be performed using the Illumina COVIDSeq Test (RUO version) or Illumina DNA Prep. Choose one of these approaches below.

STEP CASE

Illumina COVIDSeq Test (RUO version)

27 steps

5m 15s



- 5 This step reverse transcribes the RNA fragments primed with random hexamers into first-strand cDNA using reverse transcriptase.
  - \* Include a negative PCR control (NTC; nuclease-free water) for each pool.
- 5.1 Label new PCR plate CDNA1.
- 5.2 Add  $\triangle$  8.5  $\mu$ L EPH3 to each well.
- 5.3 Add  $\triangle$  8.5 µL eluted sample to each well.
- 5.4 Seal and shake at (5) 1600 rpm for (5) 00:01:00 .

1m

- 5.5 Centrifuge at  $1000 \times g$  for 1 minute.
- 5.6 Place on the preprogrammed thermal cycler and run the COVIDSeq Anneal program.

COVIDSeq Anneal program:

- Choose the preheat lid option
- Set the reaction volume to 17  $\mu$ l
- 65°C for 3 minutes
- Hold at 4°C
- 5.7 In a 1.7 ml tube, combine the following volumes to prepare First Strand cDNA Master Mix. \*Multiply each volume by the number of samples.

Reagent overage is included to account for small pipetting errors.

5.8 Add 4 8 µL master mix to each well of the CDNA1 plate.



5.9 Seal and shake at (5 1600 rpm for (5) 00:01:00 .

1m

5.10 Centrifuge at (1000 x g) for (500:01:00).

1m

5.11 Place on the preprogrammed thermal cycler and run the COVIDSeq FSS program.

COVIDSeq FSS program:

- Choose the preheat lid option
- Set the reaction volume to 25 µl
- 25°C for 5 minutes
- 50°C for 10 minutes
- 80°C for 5 minutes
- Hold at 4°C

## **Amplicon Generation**

- 6 This step uses two separate PCR reactions to amplify cDNA using the previously prepared primers pool A and B.
  - \* Include a negative PCR control (NTC; nuclease-free water) for each pool.
- 7 Label two new PCR plates POOL A and POOL B. The plates represent two separate PCR reactions.
- 8 In a 15 ml tube, combine the following volumes to prepare PCR 1 Master Mix and PCR 2 Master Mix. Multiply each volume by the number of samples.
  - \*Reagent overage is included to account for small pipetting errors.

А	В	С
Reagent	PCR 1 Master Mix (μl)	PCR 2 Master Mix (μΙ)
IPM	15	15
RABV-BAT Pool A	4.3	N/A
RABV-BAT Pool B	N/A	4.3



A	В	С
Nuclease-free water	4.7	4.7

- 9 Add Δ 20 μL PCR 1 Master Mix to each well of the POOL A plate corresponding to each well of the CDNA1 plate.
- 9.1 Add  $\sqsubseteq$  5  $\mu$ L first strand cDNA synthesis from each well of the CDNA1 plate to the corresponding well of the POOL A plate.
- 9.2 Add  $\underline{\mathbb{Z}}$  20  $\mu L$  COVIDSeq PCR 2 Master Mix to each well of the POOL B plate corresponding to each well of the CDNA1 plate.
- 9.3 Add  $\sqsubseteq$  5  $\mu$ L first strand cDNA synthesis from each well of the CDNA1 plate to the corresponding well of the POOL B plate.
- 9.4 Seal and shake at (5 1600 rpm for (5) 00:01:00 .
- 9.5 Centrifuge at  $\bigcirc$  1000 x g for  $\bigcirc$  00:01:00 .
- 9.6 Place in the preprogrammed thermal cycler and run the COVIDSeq PCR program.

COVIDSeq PCR program:

- Choose the preheat lid option
- Set the reaction volume to 25 µl
- 98°C for 3 minutes
- 35 cycles of:
- ---- 98°C for 15 seconds
- ---- 63°C for 5 minutes
- Hold at 4°C
- 10 \*Safe Stopping Point:

1m

1m



Amplicons can be stored at 🖁 -20 °C until ready to use

## **Amplicon checking**



We recommend using agarose gel electrophoresis or automated electrophoresis to check the amplicons before proceeding to library construction.

## Library preparation

The following steps were conducted according to the standard protocol, starting from "Tagment PCR Amplicons" step on page 9.



Tagment PCR Amplicons
Post Tagmentation Clean Up
Amplify Tagmented Amplicons
Pool and Clean Up Libraries
Quantify and Normalize Libraries
Pool and Dilute Libraries
Prepare for Sequencing

## Sequencing

The sequencing can be performed in any Illumina platform or flowcell using 2×150 nt reads.

#### Note

For sequencing, we recommend generating at least 50,000 reads per sample or 100,000 reads for optimal sequencing coverage.

# **Bioinformatics Analysis**

14 Sequencing reads should be analyzed by an amplicon-based sequencing pipeline. We recommend using ViralFlow Workflow which performs several genomic analyses based



on reference genome assembly. See details at <a href="https://viralflow.github.io/">https://viralflow.github.io/</a>

From our experience, better coverage and depth are obtained when using subcladespecific references. So, we usually assembly the sequences using the bat clade reference (JQ685956), then we run the consensus resulting sequence at http://rabvglue.cvr.gla.ac.uk/ typing tool, and getting the nearest reference we re-assembly using this reference genome.

## **Reference Sequence**



JQ685956.fasta 11KB

### **BED file**



JQ685956.bed 1KB

## **COI Analysis**

COI amplicon analysis can be performed using **ampliseq** pipeline, an amplicon sequencing analysis workflow using DADA2 and QIIME2.