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# Viral Sequencing, from Gunk to Graph (Two-step, strand-switching) v.2

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Coronavirus Method Development Community



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## ABSTRACT

This is a fast "gunk to graph" protocol for analysing viral RNA from nasopharyngeal swabs. The approach involves swab lysis and inactivation at the point of sampling, uses a cellulose binding / wash protocol to reduce extraction cost, incorporates sample-specific barcodes during first-strand synthesis, nanopore rapid-attachment primers during PCR amplification, and nanopore sequencing with parallel RAMPART analysis for fast assembly and phylogenetics.

## MATERIALS

NAME	CATALOG #	VENDOR
Q5 Hot Start High-Fidelity 2X Master Mix - 500 rxns	M0494L	New England Biolabs
MinION sequencer	View	Oxford Nanopore Technologies
ONT MinION Flow Cell R9.4.1	FLO-MIN106D	Oxford Nanopore Technologies

## MATERIALS TEXT

Additional materials TBA.

## SAFETY WARNINGS

This protocol is UNTESTED, and is in the early stages of development. Do not trust the protocol; question everything.

Assume samples are potentially infectious during extraction, and make sure to use proper sterile technique to avoid cross-contamination.

## Swab Lysis

- 1 Prepare a **1.5 ml** centrifuge tube with heated lysis buffer and a cellulose disc

1.1 Add **500 µl** lysis / RNase inactivation buffer ([Twitter reference](#)) to 1.5ml centrifuge tube:

- **10 Milimolar (mM) Tris**
- **10 Milimolar (mM) EDTA**
- **0.5 % volume SDS**
- **150 Milimolar (mM) NaCl**

OR **500 µl** extraction buffer #2 (see [paper](#)):

- **800 Milimolar (mM) guanidine hydrochloride**
- **50 Milimolar (mM) Tris [pH 8]**
- **0.5 % volume Triton X100**
- **1 % volume Tween-20**

1.2 Add a **3 mm diameter** punched disc from [Whatman #1 filter paper](#) (see [paper](#))

1.3 Preheat **1.5 ml** tube to **60 °C**

2 Collect sample using a sterile polystyrene swab with a 30mm breakpoint (e.g. [Puritan 25-3606-U; PurFlock Ultra 6" Sterile Elongated Flock Swab w/Polystyrene Handle, 30mm Breakpoint](#)).

#### RNA Wash

3 Transfer disc to a new **1.5 ml** tube containing **200 µl** wash buffer using a pipette tip to remove contaminants:

- **10 Milimolar (mM) Tris [pH 8.0]**
- **0.1 % volume Tween-20**

4 Incubate tube at **Room temperature** for **00:01:00**

#### cDNA Synthesis

5 Transfer disc to a new **200 µl** PCR tube using a pipette tip

6 Add the following additional components into the **200 µl** PCR tube (see the [Nanopore protocol for Sequence-specific cDNA-PCR Sequencing \(SQK-PCS109\)](#)) in a **11 µl** reaction:

- **1 µl** x **2 Micromolar (µM)** reverse primers
- **1 µl** x **10 Millimolar (mM)** dNTPs
- **9 µl** RNase-free water

Reverse primers should be prefixed with sample-specific barcode sequences (if used) and the ONT reverse anchor sequence, i.e. [5' - ACTGCGCTGCTCGCTCTATCTTC - [barcode] - [sequence-specific] - 3']

A potential primer pool are the reverse/right ARCTIC primers with barcodes and ONT anchor sequences from [here](#).

An alternative protocol using both forward and reverse primers can be found [here](#).

7 Mix gently **by flicking the tube** and spin down ⌚ **00:00:05**

8 Denature RNA and anneal primers at **65 °C** for ⌚ **00:05:00** and then snap cool on a pre-chilled freezer block for ⌚ **00:01:00**

9 In a separate tube, mix together the following in an **8 µl** reaction:

- **4 µl** 5X RT Buffer
- **1 µl** RNaseOUT
- **1 µl** Nuclease-free water
- **2 µl** x **10 Micromolar (µM)** ONT Strand-switching primer (SSP)

10 Mix gently **by flicking the tube** and spin down ⌚ **00:00:05**

11 Add the strand-switching buffer to the snap-cooled, annealed RNA, mix by **flicking the tube** and spin down

12 Incubate at **42 °C** for ⌚ **00:02:00**

13 Add **1 µl** of Maxima H Minus Reverse Transcriptase, to a total volume of **20 µl**

14 Mix gently by **flicking the tube** and spin down ⌚ **00:00:05**

- 15 Incubate using the following protocol:

Cycle step	Temperature	Time	No. of cycles
Reverse transcription and strand-switching	42° C	90 mins	1
Heat inactivation	85° C	5 mins	1
Hold	4° C	∞	

Thermal cycler settings for reverse transcription and strand switching

#### PCR amplification

- 16 In four new **200 µl** PCR tubes, prepare the following reaction at **Room temperature** in a **50 µl** reaction:

- **25 µl** 2X Q5 Hot Start High-Fidelity Master Mix
- **1.5 µl** cDNA primer (cPRM)
- **18.5 µl** Nuclease-free water
- **5 µl** Reverse-transcribed cDNA from the previous step (pool, or single sample)

- 17 Amplify using the following cycling conditions:

Cycle step	Temperature	Time	No. of cycles
Initial denaturation	95 °C	30 secs	1
Denaturation	95 °C	15 secs	10-40*
Annealing	62 °C	15 secs	10-40*
Extension	65 °C	50 secs per kb	10-40*
Final extension	65 °C	6 mins	1
Hold	4 °C	∞	

Thermal cycler settings for PCR amplification

\* Starting from viral RNA, the recommended starting point is 20 cycles - adjust this depending on experimental needs.

- 18 Add **1 µl** of NEB Exonuclease 1 (20 units) directly to each PCR tube to remove unextended primers. Mix by **pipetting**.

- 19 Incubate the reaction at **37 °C** for **00:15:00**, followed by **80 °C** for **00:15:00**

- 20 Run 1 µl of amplified product on a gel (or similar length-based QC device) to verify that amplified products exist at the expected length. Because this is a strand-switch protocol, there may be a smear of template DNA rather than specific bands.

#### Bead Cleanup

- 21 Add 160 µl of resuspended AMPure XP beads to the **1.5 ml** tube and mix by **pipetting**

- 22 Incubate on a gentle agitator (e.g. hula mixer or rotator mixer) for **00:05:00** at **Room temperature**

- 23 Spin down ⌚00:00:05 the sample and pellet on a magnet. Keep the tube on the magnet, and pipette off the supernatant.
- 24 Keep the tube on the magnet and wash the beads with 📏200 µl of freshly-prepared 📏70 % volume ethanol without disturbing the pellet. Remove the ethanol using a pipette and discard.
- 25 Repeat the previous step: wash with 📏200 µl 📏70 % volume ethanol , and discard the ethanol / wash liquid.
- 26 Spin down ⌚00:00:05 and place the tube back on the magnet. Pipette off any residual ethanol. Allow to dry for ⌚00:00:30 [at most] but do not dry the pellet to the point of cracking (the magnetic beads should just start to lose their shiny sheen).
- 27 Remove the tube from the magnetic rack and resuspend pellet in 📏12 µl of Elution Buffer (EB).
- 28 Incubate at 🌡 Room temperature for ⌚00:10:00
- 29 Pellet beads on magnet ⌚00:05:00 until the eluate is clear and colourless
- 30 While still on the magnet, carefully remove and retain 📏12 µl of eluate into a clean 📏1.5 ml Eppendorf DNA LoBind tube
- 31 Quantify 1 µl of the amplified cDNA library using the Quantus Fluorometer using the ONE dsDNA assay (see [ncov 2019 sequencing protocol, step 16](#))

#### Adapter Addition

- 32 Add 📏1 µl of Rapid Adapter (RAP) to the amplified cDNA library
- 33 Mix by *pipetting* and spin down ⌚00:00:05
- 34 Incubate the reaction for ⌚00:05:00 at 🌡 Room temperature
- 35 Store the prepared library 🌡 On ice until ready to load onto a flow cell.


#### Nanopore Sequencing

- 36 Load 📏20 ng sequencing library onto a MinION flow cell (see [ncov 2019 sequencing protocol, step 21](#))

- 37 Start the sequencing run using MinKNOW, using SQK-PCS109 as the sample preparation protocol (see [ncov 2019 sequencing protocol, step 22](#))

#### RAMPART Analysis

- 38 Analyse the run results using RAMPART (see <https://artic.network/ncov-2019/ncov2019-using-rampart.html>)

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