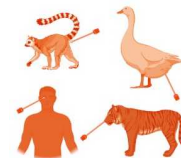


Jun 29, 2023

## TELEVir Field Protocol

DOI

[dx.doi.org/10.17504/protocols.io.n2bvj694xIk5/v1](https://dx.doi.org/10.17504/protocols.io.n2bvj694xIk5/v1)



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DOI: <https://dx.doi.org/10.17504/protocols.io.n2bvj694xIk5/v1>

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**Protocol status:** Working

**We use this protocol and it's working**

**Created:** April 06, 2022

**Last Modified:** June 29, 2023

**Protocol Integer ID:** 60387

**Keywords:** metagenomic virus detection, random amplification for metagenomic virus detection, televir field protocol, emerging virus threat, virus threats for human, minion device, vir project, oxford nanopore technology, vir project key aim, tele, wildlife animal

**Funders Acknowledgements:**

Horizon 2020

Grant ID: 773830

### Disclaimer

This protocol was prepared for a workshop held the 23-24 June 2022 at Statens Serum Institut, Denmark. Here 25 TELEVIR partners from 10 European countries participated to train in the usage of a field-deployable point-of-incidence toolbox to identify emerging virus threats. The protocol was used to detect SARS-CoV-2 and Human Papillomavirus a RNA and DNA virus detection test-of-concept.



## Abstract

The TELE-Vir project key aim is to develop a fast point-of-incidence (poi) toolbox for identification and characterisation of emerging virus threats for humans and/or domestic and wildlife animals.

Presented here is a protocol including sample pretreatment, NA extraction and random amplification for metagenomic virus detection using the MinION device (Oxford Nanopore Technologies)

## Protocol materials

☒ DNase Set I **Zymo Research** **Catalog #E1010**

☒ MagNA Pure LC Total Nucleic Acid Isolation Kit **Roche** **Catalog #03038505001**

☒ MagNA Pure LC Total Nucleic Acid Isolation Kit **Roche** **Catalog #03038505001**

☒ REPLI-g Cell WGA & WTA **Qiagen** **Catalog #150052**

☒ Rapid Sequencing Kit **Oxford Nanopore Technologies** **Catalog #SQK-RAD004**

## Troubleshooting



## Pretreatment

15m

1

15m


**DNase Set | Zymo Research Catalog #E1010**

### Equipment

**BD Disposable Syringe with Luer-Lok Tip (5ml)** NAME

Syringe TYPE

Becton Dickinson BRAND

309649 SKU

### Equipment

**Syringe Filter pore size 0.2 µm** NAME


Filter TYPE


Minisart BRAND

16532 SKU


<https://www.sartorius.com/shop/ww/en/usd/applications-laboratory-filtration-sterile-filtration/minisart-syringe-filter%2C-polyethersulfone-%28pes%29%2C-pore-size-0-22-%2C%2B5m%2C-ethylene-oxide%2C-female-luer-lock%2C-male-luer-lock%2C-pack-size-50/p/16532-> LINK

Polyethersulfone (PES), Pore Size 0.22 µm, Ethylene Oxide, Female Luer Lock, Male Luer Lock, Pack Size 50 SPECIFICATIONS


 400 µL Sample


 50 µL DNase I


 50 µL DNase Digestion buffer


 00:15:00 at room temperature

### Note

If 400 µL sample is not available, downscale the reagents to fit the lower sample volume.

2 Add  500 µL PBS (or equivalent to 1 ml sample material in total)

3 Poor the diluted sample into the lid of the Syringe Filter (Minsart). The filter can be placed on a sterile surface meanwhile e.g. the Syringe Filter paper lid (inner side).

4 Suck in air corresponding to app 1 ml in a 5 ml syringe



- 5 Extract the 1 ml sample material from the petri dish to the 5 ml syringe
- 6 Attach the 0,22 µM Minisart syringe filter to the 5 ml syringe with Luer-Lok
- 7 Filter the sample and air directly into a 4.5 ml cryotube containing 1 ml MPLB-buffer. Put lid on, mix by turning tube.

☒ MagNA Pure LC Total Nucleic Acid Isolation Kit **Roche Catalog #03038505001**

## Inactivation

20m

- 8 Incubate 🕒 00:20:00 at room temperature for viral inactivation

### Citation

Vinner L, Fomsgaard A (2007). Inactivation of orthopoxvirus for diagnostic PCR analysis.. Journal of virological methods.

### Citation

Rosenstjerne MW, Jensen CE, Fomsgaard A (2018)  
. Rapid, Safe, and Simple Manual Bedside Nucleic Acid Extraction for the Detection of Virus in Whole Blood Samples..  
Journal of visualized experiments : JoVE.

<https://doi.org/10.3791/58001>

LINK

## Extraction

10m

- 9 Kit used for hand held NA extraction:  
☒ MagNA Pure LC Total Nucleic Acid Isolation Kit **Roche Catalog #03038505001**



## Equipment

4.5 ml cryotube

NAME

tube

TYPE

Thermo Fisher Scientific

BRAND

363452

SKU

<https://www.thermofisher.com/order/catalog/product/363452><sup>LINK</sup>

## Equipment

3.6 ml cryotube

NAME

Tube

TYPE

Thermo Fisher Scientific

BRAND

379189

SKU

<https://www.thermofisher.com/order/catalog/product/363452><sup>LINK</sup>

Cylinder magnet

A rubber band to attach the magnet to a tube.

A 50 ml Nunc tube or similar to pour excess solutions into for disposing.

Following the protocol by:

## Citation

Rosenstjerne MW, Jensen CE, Fomsgaard A (2018)

. Rapid, Safe, and Simple Manual Bedside Nucleic Acid Extraction for the Detection of Virus in Whole Blood Samples..





Journal of visualized experiments : JoVE.

<https://doi.org/10.3791/58001>

LINK

10 Per *X* number of samples, aliquot in cryotubes or Eppendorf tubes

10m


 960 µL Magnetic Glass Particles (MGPs) in an Eppendorf tube 4 mL Wash buffer I in a 4.5 ml cryotube 1.5 mL Wash buffer II in an Eppendorf tube 3 mL Wash buffer III in a 3.6 ml cryotube 100 µL Elution buffer in an Eppendorf tube

Prepare one sample at the time.

Approximate time  00:10:00 per sample

11 Add 960 µl MGPs to 2 ml solution of sample and MPLB-buffer



- 12 Put lid on and turn tube in hand gently until well mixed
- 13 Attach magnet and trap beads.
- 14 Pour solution into trash tube. The MGPs should stay behind trapped on the side of the tube.
- 15 Pour Wash buffer I in the sample tube
- 16 Put a lid on, remove magnet and turn the tube in hand gently until well mixed
- 17 Attach magnet and trap beads.
- 18 Pour solution into trash tube. The MGPs should stay behind trapped on the side of the tube.
- 19 Pour Wash buffer II in the sample tube
- 20 Put a lid on, remove magnet and turn the tube in hand gently until well mixed
- 21 Attach magnet and trap beads.
- 22 Pour solution into trash tube. The MGPs should stay behind trapped on the side of the tube.
- 23 Pour Wash buffer III in the sample tube
- 24 Put a lid on, remove magnet and turn the tube in hand gently until well mixed
- 25 Attach magnet and trap beads.
- 26 Pour solution into trash tube. The MGPs should stay behind trapped on the side of the tube.
- 27 Pour Elution buffer in the sample tube
- 28 Put a lid on, remove magnet and turn the gently until beads are eluted.
- 29 Transfer  10 µL Elution Buffer with MGPs to **2 x** 0.2 ml PCR tube

## Whole Transcriptome and Genome Amplification

4h

- 30
- Equipment to enable portability
- A MiniPCR to run isothermal incubations
  - A powerbank that can provide 20V. Use one to power the miniPCR and another to power the MinION for approximately 5 hours. Use brand of choice.
  - A salad swing as a hand driven centrifuge. Use brand of choice.
  - A coolbox for reagents and cooling samples. Use brand of choice.

Equipment	
All-in1 Laptop Powerbank 24000	NAME
Power Bank	TYPE
Sanberg	BRAND
420-57	SKU
<a href="https://sandberg.world/da-dk/product/all-in-1-laptop-powerbank-2400">https://sandberg.world/da-dk/product/all-in-1-laptop-powerbank-2400</a>	LINK
Any powerbank that can produce 20V is acceptable. Have 2-3 to run the MiniPCR and the MinION	SPECIFICATIONS

Equipment	
miniPCR® mini8 thermal cycler	NAME
Thermal cycler	TYPE
miniPCR®	BRAND
mini8	SKU
<a href="https://www.minipcr.com/">https://www.minipcr.com/</a>	LINK
	

## Equipment

### A Salad Spinner Centrifuge

Centrifuge

undefined

undefined

[https://www.oxo.com/salad-spinner.html?](https://www.oxo.com/salad-spinner.html?queryID=975936c98686e2b0325b684dd613daff&objectID=2251&indexName=magentc)

[queryID=975936c98686e2b0325b684dd613daff&objectID=2251&indexName=magentc](https://www.oxo.com/salad-spinner.html?queryID=975936c98686e2b0325b684dd613daff&objectID=2251&indexName=magentc)



Kit used for WTA and WGA

REPLI-g Cell WGA & WTA Qiagen Catalog #150052

Quantiscript RT Enzyme Mix, Ligase Mix, and REPLI-g SensiPhi DNA Polymerase should be thawed on ice.

All other components can be thawed at room temperature (15–25°C)

- 31 The two aliquots of same sample enables uniform whole genome amplification (WGA) and whole transcriptome amplification (WTA) in parallel reactions.

### 32 Cleanup for WTA

Add 2 µL gDNA Wipeout Buffer to PCR tube with 10 µl eluted sample material

00:10:00 42 °C on MiniPCR

10m

### 33 Repair (WGA) and Reverse Transcription (WTA)

- 33.1 For **WGA**, mix following and add 10 µL Repair reagent mix to sample

Reagents	Volume (µL) per sample
RT/Polymerase Buffer	4
gDNA Wipeout buffer	2
H2O sc	1
Random primer	1
Random Hex-P primer (20µM)	1
WGA Ready Enzym	1
<b>Total</b>	<b>10</b>

This modified protocol uses 5'-phosphorylated random hexamers (P-N6) instead of kit provided oligo-dT primers according to Rosenstierne et al (2014).



## Citation

Rosenstierne MW, McLoughlin KS, Olesen ML, Papa A, Gardner SN, Engler O, Plumet S, Mirazimi A, Weidmann M, Niedrig M, Fomsgaard A, Erlandsson L (2014)

. The microbial detection array for detection of emerging viruses in clinical samples--a useful panmicrobial diagnostic tool..

PLoS one.

<https://doi.org/10.1371/journal.pone.0100813>

LINK

33.2 For **WTA**, mix following and add 8 µL RT reagent mix to sample

Reagents	Volume (µL) per sample
RT/Polymerase Buffer	4
H2O sc	1
Random primer	1
Random Hex-P primer (20µM)	1
Quantiscript RT Enzym Mix	1
<b>Total</b>	<b>8</b>

33.3 Incubate

01:00:00

42 °C

On MiniPCR

1h 3m

Afterwards immediatly

00:03:00

95 °C

on MiniPCR

34 **Ligation**

Mix following and add 10 µL Ligation mix to each tube

35m

Reagents	Volume (µL) per sample
Ligation buffer	8
Ligase Mix	2
<b>Total</b>	<b>10</b>

Incubate

00:30:00

24 °C This is the Repli-G recommended temperature, however, the miniPCR only goes to 25C. Instead of incubation in the MiniPCR, try table-top incubation.

00:05:00

95 °C

Hold at 4 °C

35 **Amplification**

Mix following and add 30 µL amplification mix to each tube

2h 5m

Reagents	Volume (μL) per sample
Repli-g reaction buffer	29
Repli-g SensiPhi DNA polymerase	1
<b>Total</b>	<b>30</b>

Incubate

🕒 02:00:00 🌡️ 30 °C

🕒 00:05:00 🌡️ 65 °C

Hold at 🌡️ 4 °C

36 For library preparation use amplified material **without** MGPs.

## Library Preparation

37

Multiple rapid library preparation kits are available through **Oxford Nanopore Technology**.

For Workshop we use.

🔗 Rapid Sequencing Kit **Oxford Nanopore Technologies Catalog #SQK-RAD004**

### Equipment

MinION Mk1C	NAME
Sequencer	TYPE
Oxford Nanopore Technology	BRAND
MIN-101C	SKU
<a href="https://store.nanoporetech.com/minion-mk1c.html">https://store.nanoporetech.com/minion-mk1c.html</a>	LINK





## Citations

### Step 33.1

Rosenstierne MW, McLoughlin KS, Olesen ML, Papa A, Gardner SN, Engler O, Plumet S, Mirazimi A, Weidmann M, Niedrig M, Fomsgaard A, Erlandsson L. The microbial detection array for detection of emerging viruses in clinical samples--a useful panmicrobial diagnostic tool.

<https://doi.org/10.1371/journal.pone.0100813>

### Step 8

Vinner L, Fomsgaard A. Inactivation of orthopoxvirus for diagnostic PCR analysis.

### Step 8

Rosenstierne MW, Jensen CE, Fomsgaard A. Rapid, Safe, and Simple Manual Bedside Nucleic Acid Extraction for the Detection of Virus in Whole Blood Samples.

<https://doi.org/10.3791/58001>

### Step 9

Rosenstierne MW, Jensen CE, Fomsgaard A. Rapid, Safe, and Simple Manual Bedside Nucleic Acid Extraction for the Detection of Virus in Whole Blood Samples.

<https://doi.org/10.3791/58001>