

COVID19 RTLAMP Assay

DOI

dx.doi.org/10.17504/protocols.io.bkx6kxre

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Protocol Citation: matt, Eugene Joseph <eugene@primediscoveries.com>, Arun Manoharan Arunprimediscoveriescom . COVID19 RTLAMP Assay. **protocols.io** <https://dx.doi.org/10.17504/protocols.io.bkx6kxre>

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Created: September 06, 2020

Last Modified: September 08, 2020

Protocol Integer ID: 41694

Keywords: rtlamp assay, assay

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Materials

MATERIALS

✕ RNAse and DNase-free 1.5ml tubes

✕ Tubes, strips of 8 **Thermo Fisher Catalog #AB0452**

STEP MATERIALS

✕ Binding Solution

✕ Binding Solution

✕ Wash Solution

✕ Resuspension Buffer

✕ Reaction Mix

✕ Primer Mix

✕ Positive Control

✕ Lysis Buffer

✕ Lysis Buffer



Protocol materials

⊗ Tubes, strips of 8 **Thermo Fisher Catalog #AB0452**

⊗ Binding Solution

⊗ Primer Mix

⊗ Lysis Buffer

⊗ RNase and DNase-free 1.5ml tubes

⊗ Binding Solution

⊗ Wash Solution

⊗ Resuspension Buffer

⊗ Reaction Mix

⊗ Positive Control

⊗ Lysis Buffer

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⊗ Binding Solution

⊗ Binding Solution

⊗ Wash Solution

⊗ Resuspension Buffer

⊗ Lysis Buffer

Troubleshooting

Safety warnings

! Biohazardous Materials


Before start

Heat incubator



Sample Lysis / Inactivation

- 1 Centrifuge the Sample in a centrifuge that can fit 15ml tubes, or let the solid debris settle for 30 min on Ice.


 00:30:00

Thaw the Lysis Buffer on ice

- 2 **If using a Heat Block and Following the Full Method (Includes Purification / Concentration)**

Transfer 100ul of the sample Input Material (or clarified supernatant if Fresh Saliva) to a new 1ml tube containing an equal volume of 100ul of Lysis Buffer and pipette mix. Label this tube to indicate the sample name.


 Lysis Buffer

 100 μ L Lysis Buffer

 100 μ L Clarified Sample

- 3 Use a tube cap lock to prevent tubes from popping open.

- 4 Place the tubes in a Heat block or water bath set at 95°C for 5 min

 00:05:00

Equipment

ThermoMixer

NAME

Benchtop Incubator

TYPE

Eppendorf

BRAND

5382000023

SKU

<https://online-shop.eppendorf.us/US-en/Temperature-Control-and-Mixing-44518/Instruments-44519/Eppendorf-ThermoMixerC-PF-19703.html>


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Any heat block will suffice

SPECIFICATIONS





- 5 Carefully transfer the tubes to ice or a 4°C Cold Block for 30 sec, or let sit at Room Temperature for 5 min.

 00:00:30


 4 °C

Or

 00:05:00

 Room temperature

- 6 Centrifuge for 3 sec at 2000xg to spin down any condensation.

 00:00:03



Equipment

Mini-centrifuge

NAME

Centrifuge

TYPE

Fisher

BRAND

S67601B

SKU

<https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-mini-centrifuge/s67601b>

LINK

Any standard mini centrifuge with adapters for different tube sizes will suffice

SPECIFICATIONS




Samples are ready go to Step #11

7 If using a 96 well Thermal Cycler and following the Quick Method (Without Purification / Concentration)


Transfer 20ul of the sample Input Material (or clarified supernatant if Fresh Saliva) to a new 200ul tube containing an equal volume of 20ul of Lysis Buffer and pipette mix. Label this tube to indicate the sample name.

 Lysis Buffer

 20 µL Lysis Buffer

 20 µL Clarified Sample

8 Create a Method on the Thermal Cycler for 95°C for 5 min followed by a Hold at 4°C, with the lid set to 105°C.

 00:05:00

 95 °C

Hold at  4 °C

Equipment

SimpliAmp Thermal Cycler

NAME

PCR

TYPE

Applied Biosystems

BRAND

A24811

SKU

<https://www.thermofisher.com/order/catalog/product/A24811>^{LINK}


Any standard PCR thermocycler will suffice

SPECIFICATIONS



9 Place the tubes in the Thermal Cycler and run the method

10 Centrifuge for 3 sec at 2000xg to spin down any condensation.

 00:00:03

Equipment

Mini-centrifuge

NAME

Centrifuge

TYPE

Fisher

BRAND

S67601B

SKU

<https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-mini-centrifuge/s67601b>

LINK

Any standard mini centrifuge with adapters for different tube sizes will suffice

SPECIFICATIONS



Samples are ready go to step #25


Sample Purification / Concentration

11

Preheat the Heat Block or Water Bath


12


Vortex to mix the Binding Solution so there is no sediment.

 Binding Solution

13

To each sample tube, add 800ul of Binding Solution

 Binding Solution

 800 μ L Binding Solution



- 14 Let the Tubes sit at Room Temperature for 10 min, every 2 min invert the tube to suspend the Binding Solution.

 00:10:00 at  Room temperature

- 15 Centrifuge for 3 sec at 2000xg to pellet the Binding Reagent in the Binding Solution Note: it should appear as a white smear at the bottom of the tube. The Binding Reagent should adhere to the bottom of the tube with a short spin on a simple benchtop centrifuge. Depending on tube type and input sample viscosity, a longer spin may be required until it is sufficiently pelleted.

Equipment

Mini-centrifuge

NAME

Centrifuge

TYPE

Fisher

BRAND

S67601B

SKU

<https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-mini-centrifuge/s67601b>

LINK

Any standard mini centrifuge with adapters for different tube sizes will suffice


SPECIFICATIONS




- 16 Remove the supernatant with a 1ml Pipette taking care not to disturb the pellet.


 1000 μ L Supernatant

- 17 Add 1ml of Wash Solution, wash the pellet by closing and inverting the tube several times.

 Wash Solution

 1000 µL Wash Solution

18 Centrifuge for 3 sec at 2000xg to pellet the Binding Reagent in the Binding Solution

 00:00:03

Equipment

Mini-centrifuge

NAME

Centrifuge

TYPE

Fisher

BRAND

S67601B

SKU

<https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-mini-centrifuge/s67601b>


LINK

Any standard mini centrifuge with adapters for different tube sizes will suffice


SPECIFICATIONS



19 Remove the supernatant with a 1ml Pipette taking care not to disturb the pellet.

 1000 µL Wash Supernatant

20 Centrifuge for 3 sec at 2000xg to collect any residual Wash Solution.

 00:00:03



Equipment

Mini-centrifuge

NAME

Centrifuge

TYPE

Fisher

BRAND

S67601B

SKU

<https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-mini-centrifuge/s67601b>

LINK

Any standard mini centrifuge with adapters for different tube sizes will suffice


SPECIFICATIONS



- 21 Remove the residual supernatant with a 200ul Pipette taking care not to disturb the pellet.

 200 μ L Wash Supernatant

- 22 Air dry in a clean ventilated area until the Binding Reagent is dry, it should appear opaque and not reflective, air drying should take 10-15 min, some sample types may take longer to dry, do not let drying take longer than 30 min. Note: If the lab is equipped, air drying should take place in a ventilated air cabinet to prevent cross contamination. To speed up the air drying, if available it is possible to use a Centrifuge with open tube lids, or a heated vacuum concentrator. If using these it would take less than 15 minutes as it should be evaporating less than 5-10ul of Wash Solution.

 00:15:00

- 23 To each tube add 10ul of Resuspension Buffer, pipette mix to resuspend the pellet. Depending on the Sample Input, the pellet may resuspend easily or stay as clumps that will take more pipetting to break apart.

⚗ Resuspension Buffer

🧪 5 µL Resuspension Buffer

- 24 Vortex until the Binding Reagent is resuspended, you can briefly centrifuge for 3 sec at 2000xg to collect droplets from the sides of the tube. Note: As the sample contains extracted resuspended RNA, the RT-LAMP reaction should proceed the same day. The extracted samples should not be stored or frozen as it would impact the RNA integrity.

🕒 00:00:03

Equipment

Mini-centrifuge

NAME

Centrifuge

TYPE

Fisher

BRAND

S67601B

SKU

<https://www.fishersci.com/shop/products/fisherbrand-standard-mini-centrifuge-standard-mini-centrifuge/s67601b>

LINK

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SPECIFICATIONS





Samples are ready go to Step #25

RT-LAMP Reaction


- 25 Prepare the qPCR instrument, have Ice on hand to prepare the master Assay mix. Calculate the amount of samples being run and make sure to include the Positive Control and the H2O.
- 26 Prepare the Assay mix by adding Reaction Mix and Primer mix according to the amount of samples being run.



 Reaction Mix

 Primer Mix

 12 μ L Reaction Mix per Sample

 3 μ L Primer Mix per Sample

27 Transfer 15ul of Assay mix into the Assay wells.

 15 μ L Assay Mix

28 Transfer 5ul of each sample into the appropriate Assay well, pipette mix.

 5 μ L Sample

29 Add the Positive Control to the Plate or Strip Tube

 Positive Control

 5 μ L Positive Control

30 Transfer 5ul of H2O into the appropriate Negative Control well.

 5 μ L H2O

31 Seal the plate and run it in the qPCR instrument with the method:

Capture 2 Channels:

FAM - No Quencer = Assay Target

Cy3 - No Quencher = Control Target



00:00:30



68.5 °C

and Capture Image

Repeat 80x