



Oct 04, 2021

SARS-CoV-2 Illumina MiSeq protocol v.2 V.2

Public Health Ontario¹

¹Public Health Ontario Laboratory

Works for me



dx.doi.org/10.17504/protocols.io.bs98nh9w

wgscov

DISCLAIMER

DISCLAIMER - FOR INFORMATIONAL PURPOSES ONLY: USE AT YOUR OWN RISK

The protocol content here is for informational purposes only and does not constitute legal, medical, clinical, or safety advice, or otherwise; content added to protocols.io is not peer reviewed and may not have undergone a formal approval of any kind. Information presented in this protocol should not substitute for independent professional judgment, advice, diagnosis, or treatment. Any action you take or refrain from taking using or relying upon the information presented here is strictly at your own risk. You agree that neither the Company nor any of the authors, contributors, administrators, or anyone else associated with protocols.io, can be held responsible for your use of the information contained in or linked to this protocol or any of our Sites/Apps and Services.

ABSTRACT

ARTIC amplicon sequencing protocol adapted from Josh Quick's https://www.protocols.io/view/ncov-2019sequencing-protocol-v2-bdp7i5rn for illumina sequencing of SARS-CoV-2

dx.doi.org/10.17504/protocols.io.bs98nh9w

PROTOCOL CITATION

Public Health Ontario 2021. SARS-CoV-2 Illumina MiSeg protocol v.2. protocols.io https://dx.doi.org/10.17504/protocols.io.bs98nh9w

Version created by wgscov

LICENSE

 $_{\!\scriptscriptstyle \Box}$ This is an open access protocol distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

CREATED

Mar 12, 2021

LAST MODIFIED

Oct 04, 2021

PROTOCOL INTEGER ID

48160

DISCLAIMER:

DISCLAIMER - FOR INFORMATIONAL PURPOSES ONLY; USE AT YOUR OWN RISK

The protocol content here is for informational purposes only and does not constitute legal, medical, clinical, or safety

mprotocols.io

10/04/2021

Citation: Public Health Ontario (10/04/2021). SARS-CoV-2 Illumina MiSeq protocol v.2. https://dx.doi.org/10.17504/protocols.io.bs98nh9w

advice, or otherwise; content added to <u>protocols.io</u> is not peer reviewed and may not have undergone a formal approval of any kind. Information presented in this protocol should not substitute for independent professional judgment, advice, diagnosis, or treatment. Any action you take or refrain from taking using or relying upon the information presented here is strictly at your own risk. You agree that neither the Company nor any of the authors, contributors, administrators, or anyone else associated with <u>protocols.io</u>, can be held responsible for your use of the information contained in or linked to this protocol or any of our Sites/Apps and Services.

Mix the following components:	15r	m						
Component	Volume							
50 μM random hexamers	⊒1 μl							
10mM dNTPs mix (10mM each)	□ 1 μl							
Total Mastermix volume	⊒2 μl							
(template RNA)	⊒11 μl							
Total Reaction volume	⊒13 μl							
Prepare Mastermix (1:1) of random hexamers and dNTP.								
Mix gently and pulse centrifuge to collect liquid at the bottom of the Mastermix tube.								
The Mastermix should be prepared in a clean room and the nucleic acids added in a BSC or workbench exclusive for RNA work.								
Aliquot 🗆 2 ul of this mix into ea	5r ch well of a 96 well plate. Keen the plate in a cold block	m						
, inquest == processing time time time.								
Use multichannel pipette to aliquo	5r t □11 μl of RNA to the plate from step 2. Seal plate, mix gently on plate mixer, an	m 1d						
briefly centrifuge the plate to collect the liquid at bottom of the wells.								
Incubate the reaction mix in thern	ocycler as follows:	m						
8 65 °C © 00:05:00								
8 4 °C © 00:01:00								
F Prepare the following mastermix: 15m								
5 Prepare the following mastermix:								
Component	Volume							
SSIV Buffer	34 μ Ι							
100mM DTT	⊒1 μl							
	Total Mastermix volume (template RNA) Total Reaction volume Prepare Mastermix (1:1) of randor Mix gently and pulse centrifuge to The Mastermix should be preparent for RNA work. Aliquot 2 µl of this mix into ease of the plate to colled the plate to colled the plate the plate to colled the pla	50 μM random hexamers						

⊒1 μl RNaseOUT RNase Inhibitor SSIV Reverse Transcriptase **□**1 μl **Total Mastermix volume □**7 μl (denatured RNA) **⊒13** μl **Total Reaction volume ⊒20 μl** Add 🖵 7 µl of mastermix to the denatured RNA from the previous step. Cover the plate with seal, mix gently on a plate mixer, and pulse spin the plate to collect liquid at the bottom of the tube. The Mastermix should be prepared in in a clean room and added to the denatured RNA in a BSC or workbench exclusive for RNA work. 1h Incubate in a thermocycler as follows: 6 **© 00:50:00** 8 42 °C 8 70 °C **© 00:10:00** 84°C Hold Multiplex PCR 4h 10m Prepare the multiplex PCR reactions as follows and aliquot in each well of a 96-well plate x2 (1 for each pool): Pool 2 Component Pool 1 5X Q5 Reaction Buffer **□**5 μl **⊒**5 μl 10 mM dNTPs **□**0.5 μl **□**0.5 μl Q5 Hot Start DNA Polymerase **■**0.25 µl **■**0.25 µl Primer Pool 1 or 2 (10µM) **■3.6** μl **■**3.6 μl Nuclease-free water **■**13.15 μl **■**13.15 μl **Total Mastermix volume ■22.5** µl **■22.5** μl (cDNA) **⊒2.5** μl **□2.5** ul Total reaction volume **⊒**25 μl **⊒25** μl Prealiquot 22.5 µl of each mastermix(pool1 and pool2) to each plate (pool1 and pool2) accordingly. In a BSC or workbench exclusive for RNA work, add 2.5 µl of cDNA from step 6 to each plate. Cover the plate with seal, mix gently on a plate mixer, and pulse spin the plate to collect liquid at the bottom of the tube. 3h 30m Run the 3.5 hours PCR program for each pool: **Temperature** Step Time Cycles mprotocols.io 3

Citation: Public Health Ontario (10/04/2021). SARS-CoV-2 Illumina MiSeq protocol v.2. https://dx.doi.org/10.17504/protocols.io.bs98nh9w

10/04/2021

```
      Heat Activation
      § 98 °C
      © 00:00:30
      1

      Denaturation
      § 98 °C
      © 00:00:15
      35

      Annealing
      § 65 °C
      © 00:05:00
      35

      Hold
      § 4 °C
      1
```

Amplicon Clean-up 1h

5m

- 10 Combine the two pools of amplicons:

 Add 12.5 μl of each Pool 1 and Pool 2 (total 25μl) in an 0.2 ml PCR plate (low binding plate).
- 11 Perform AMPure XP bead cleanup according to directions, as follows.

45m

- 11.1 Add 25 μl of AMPure XP beads(well-vortexed and at & Room temperature) to the combined amplicons plate. Cover the plate with seal, mix gently on a plate mixer, and pulse spin the plate to collect liquid at the bottom of the tube. Incubate at & Room temperature for © 00:05:00.
- 11.2 Place the plate on a magnetic rack for **© 00:05:00**, or until the beads have pelleted and the supernatant is completely clear.
- 11.3 Remove and discard the liquid from each well with a multichannel pippette, being careful not to touch the bead pellet.
- 11.4 Add 200 μl of freshly prepared, & Room temperature 80% ethanol to each well, incubate for © 00:00:30, remove the ethanol carefully with a multichannel pipette.
- 11.5 Repeat ethanol wash (step 11.3 and 11.4).
- 11.6 Discard all ethanol and carefully remove as much residual ethanol as possible using a multichannel pipette. With the plate uncovered, incubate for 3-5 min or until the pellet loses its shine (if the pellet dries completely it will crack and become difficult to resuspend).
- 11.7 Remove from magnetic rack, add 28 μl of EB buffer to wells and mix gently on a plate mixer, ensuring beads are well re-suspended. Briefly centrifuge the plate to collect the liquid at the bottom of the wells. Incubate at δ Room temperature for © 00:05:00.
- 11.8 Place the plate on magnetic rack and incubate for © 00:02:00 to © 00:05:00 or until the beads have pelleted and the supernatant is completely clear.

	11.9	Transfer $\ \ \ \ \ \ \ \ \ \ \ \ \ $					
Gel elec	trophoresis	1h					
12	20m						
13	Load 2 μl of th	e 100 bp ladder into gel on either side of each row of wells.					
14	Dispense 2 μl c	of 6X loading dye into each sample with a multichannel pipette, mix and load 2 μ l of this mix into the gel.					
15	Run at 240V fo	or © 00:20:00 . Visualize PCR products, confirm bands of approximately 300bp size.					
Amplico	on quantification	and normalization 2h					
16	Quantify amplicons using Qubit dsDNA High Sensitivity kit and plate reader according to directions, as follows.						
	16.1 Create Qubit dsDNA HS working solution by mixing 99.5 μl X buffer and 0.5 μl X dye (X is the total number of samples, including 6 standards). Using a reservoir and multichannel pipette, dispense 98 μl into required number of wells of a Costar 3590 flat-bottom plate (or as appropriate for plate reader).						
	16.2	Dilute the clean, pooled amplicons (from step 11.9) 1:10 by mixing $\Box 3 \mu I$ of the amplicons in $\Box 27 \mu I$ of nuclease free water.					
	Make up serial standards using 1:2 dilutions of 10 ng/ul stock (Standard 2) from the Qubit HS. This creates 5 standards in the following concentrations: [M]10 ng/ul [M]5 ng/ul [M]2.5 ng/ul [M]1.25 ng/ul [M]0.625 ng/ul plus Standard 1 [M]0 ng/ul standard 1.						
	16.4	Mix $\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \$					
17	Based on the a	mplicon concentration, normalize of all the samples amplicon concentration to [M] 0.2 ng/ul .					
	This can be done by adding 2.5 µl of diluted amplicon to a plate with prealiquoted, appropriate amount of nuclea free water.						
Library	preparation	2h					
18	Prepare seque	ncing libraries with Nextera XT DNA Library Prep kit at half volume, as follows.					

i protocols.io 5 10/04/2021

30m Tagment DNA.

19

Thaw the following Nextera XT reagents on ice:

Amplicon tagment mix (ATM)

Tagment DNA buffer (TD)

Nextera PCR master mix (NPM)

Thaw the index primers, mix by vortex each vial and spin down the liquid at the bottom of the vials.

Neutralization buffer (NT) at § Room temperature

- 19.1 Add the following reagents in order:
 - 1. $\square 5 \mu I$ of TD buffer
 - 2. $\square 2.5 \mu l$ of [M] 0.2 ng/ul amplicon (from step 17)
 - 3. **2.5** μ**l** of ATM

Cover plate with plate seal, mix gently on plate mixer and centrifuge for 1 min.

19.2 Incubate in thermocycler with the following steps:

```
8 55 °C © 00:05:00
§ 10 °C hold
```

19.3 Add 22.5 µl of NT buffer to each well and mix by pipetting up and down for 3 times, briefly spin down the plate and incubate at $\$ **Room temperature** for \bigcirc **00:05:00**.

1h

20 PCR Amplification.

Thaw the following reagents on ice:

NMP

Index primers

Invert all reagents 3 - 5 times, followed by pulse spin.

- 20.1 Add $7.5 \,\mu l$ of Nextera PCR mastermix to each well.
 - 20.2 From the pre-aliquoted index plate, add $\Box 5 \mu I$ ($\Box 2.5 \mu I$ of each i5 and i7 index of the corresponding index combination to each well. Cover plate with plate seal, gently mix on plate mixer, and centrifuge for 1 min.
 - 20.3 Run the PCR program to amplify the libraries:

Step	Tempe	rature	Time	9	Cycles
1	8 72 °C	© 00:03: 0	00	1	
2	8 95 °C	© 00:00:3	30	1	

mprotocols.io 10/04/2021

```
3
                                          8 95 °C
                                                      © 00:00:10
                                                                         12
                        3
                                                      © 00:00:30
                                                                        12
                                          8 55 °C
                        3
                                         A 72 °C
                                                      © 00:00:30
                                                                        12
                                         8 72 °C
                                                      © 00:05:00
                                                                        1
                        4
                        5
                                          8 4 °C
                                                      Hold
                                                                       1
Library Clean-up
                               2h
                                                                                                                      45m
        Clean Up Libraries
  21
        Repeat the same clean up process as step 11 using 20 µl of AMPure XP beads and 28 µl of resuspension
        buffer.
Library Quantification
                                    2h
                                                                                                                      30m
  22
        Repeat the same quantification process as Step 16 but do NOT dilute libraries.
Normalization and loading on Miseq
                                                  2h
                                                                                                                      30m
  23
        Normalize each library to [M]4 Nanomolar (nM) by dilution with nuclease free water.
                                                                                                                      15m
        Pool equal volume (e.g. □5 µl ) from each of the normalized libraries into a single □1.5 mL microtube.
                                                                                                                      20m
        Verify fragment size and concentration using Agilent D5000 Assay on TapeStation 4200 as follows.
               25.1
                        Add 2 µl of Sample Buffer and 2 µl of your pooled libraries in triplicate in a strip tube.
               25.2 Vortex using the adapter at 2000 rpm for 1 min.
               25.3
                        Load tubes, tapes, and tips. Start run. Using library concentration and fragment size, calculate the
                        molarity of the libraries using the following formula:
                        Molarity = concentration ng/uL * (1515.1515/fragment size(bp))
                                                                                                                      10m
        Denature and load pooled libraries as follows.
  26
               26.1
                        Denature the pooled libraries by mixing \Box 5 \mu I of pooled libraries and \Box 5 \mu I of freshly made 0.2N
                        NaOH solution.
                        Incubate for © 00:05:00.
```

ு protocols.io 7 10/04/2021

- 26.2 Add **3990 μI** of HT1 buffer and mix well with denatured pooled library by pipetting up and down 10 times with P1000.
- 26.3 Load **600** μI of the denatured, diluted pooled library into the loading position of the Illumina reagent cartridge (V2, 300 cycle kit). Load reagent cartridge, flow cell, and PR2 buffer into Miseq instrument, confirm the metrics and start the run.