

Aug 25, 2020

Version 3

nCoV-2019 sequencing protocol v3 (LoCost) V.3

 Version 1 is forked from [Ebola virus sequencing protocol](#)

 In 1 collection

DOI

<https://dx.doi.org/10.17504/protocols.io.bp2l6n26rgqe/v3>

Josh Quick¹

¹University of Birmingham

ARTIC

Coronavirus Method De...

1 more workspace



Josh Quick

University of Birmingham



Create & collaborate more with a free account

Edit and publish protocols, collaborate in communities, share insights through comments, and track progress with run records.

Create free account

OPEN  ACCESS



DOI: <https://dx.doi.org/10.17504/protocols.io.bp2l6n26rgqe/v3>

Protocol Citation: Josh Quick 2020. nCoV-2019 sequencing protocol v3 (LoCost). [protocols.io](#)
<https://dx.doi.org/10.17504/protocols.io.bp2l6n26rgqe/v3> Version created by [Josh Quick](#)

**Manuscript citation:**

Improvements to the ARTIC multiplex PCR method for SARS-CoV-2 genome sequencing using nanopore. John R Tyson, et. al. bioRxiv 2020.09.04.283077; doi:<https://doi.org/10.1101/2020.09.04.283077>

License: 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

Protocol status: Working

We use this protocol and it's working

Created: July 01, 2020

Last Modified: August 25, 2020

Protocol Integer ID: 38778

Keywords: sequencing protocol v3, ultra ii ligation module, sequencing protocol, protocol for sar, ta ligase master mix, lunascript rt supermix, superscript iv

Abstract

Amplicon sequencing protocol for SARS-CoV-2 v3 (LoCost)

We thank the ARTIC network, Oxford Nanopore Technologies, New England Biolabs, BCCDC, COG-UK, CanCOGen and protocols.io commenters for their assistance developing this protocol.

Changes in this version:

- Up to 95 samples per run with EXP-NBD196 native barcode kit
- Substitution of SuperScript IV for LunaScript RT SuperMix and reaction volume reduced to 10 uL.
- Substitution of Ultra II Ligation Module for Blunt/TA Ligase Master Mix and reaction volume reduced to 10 µL.
- Native barcode ligation reaction volume reduced to 10 uL.
- SFB wash volume reduced.

Materials

	Component	Supplier	Part number
	ARTIC nCoV-2019 V3 panel (100uM)	IDT	See links below
	LunaScript RT SuperMix Kit	NEB	E3010
	Q5 Hot Start HF Polymerase or	NEB	M0493
	Q5 Hot Start High-Fidelity 2X Master Mix	NEB	M0494
	dNTP Solution Mix (10 mM ea.)	NEB	N0447
	Nuclease-free water (100 mL)	NEB	B1500
	NEBNext Ultra II End Repair/dA-tailing module	NEB	E7546
	Blunt/TA Ligase Master Mix	NEB	M0367
	Native Barcoding Expansion Kit 1-12 and/or	ONT	EXP-NBD104
	Native Barcoding Expansion Kit 13-24 or	ONT	EXP-NBD114
	Native Barcoding Expansion Kit 96	ONT	EXP-NBD196
	AMPure XP beads	Beckman	A63881
	NEBNext Quick Ligation Module	NEB	E6056S
	Sequencing Auxiliary Vials	ONT	EXP-AUX001
	Short Fragment Buffer Expansion Kit	ONT	EXP-SFB001



	Qubit dsDNA HS Assay Kit	Thermo	Q32 854
	Flow Cell Priming Kit	ONT	EXP- FLP 002
	Flow Cell Wash Kit (optional)	ONT	EXP- WS H00 3
	R9.4.1 flow cells	ONT	FLO- MIN1 06

IDT premixed **ARTIC nCoV-2019 V3 panel** or order oligos **individually**.

Before start

Prepare between 11 and 95 RNA samples plus 1 negative control using this protocol.



cDNA preparation

30m

- 1 Prepare between 11 and 95 RNA samples plus 1 negative control of nuclease-free water per library. If previously frozen, mix by briefly vortexing and pulse spin to collect liquid. Keep samples on ice at all times.

Note

At least 11 samples are required to have sufficient material to load on the sequencer at the end. If you process >23 you will need the EXP-NBD196 expansion kit.

Note

A positive control can also be included which may be a synthetic RNA constructs or high-titre clinical sample which can be diluted. This can help monitor run performance.

- 2 Mix the following components in a PCR strip-tubes/plate. Gently mix by pipetting and pulse spin the tube to collect liquid.

Component	Volume
LunaScript RT SuperMix (5X)	2 μ L
Template RNA	8 μ L
Total	10 μL

Note

Viral RNA input from a clinical sample should be between Ct 18-35. If Ct is between 12-15, then dilute the sample 100-fold in water, if between 15-18 then dilute 10-fold in water. This will reduce the likelihood of PCR-inhibition.

Note

To prevent pre-PCR contamination the mastermix should be added to the PCR strip-tubes/plate in the **mastermix** cabinet which should be cleaned with decontamination wipes and UV sterilised before and after use.

RNA samples should be added in the **extraction/sample addition** cabinet which should be cleaned with decontamination wipes and UV sterilised before and after use.

- 3 Incubate the reaction as follows:

25 °C for 00:02:00

55 °C for 00:10:00

95 °C for 00:01:00

Hold at 4 °C

Primer pool preparation (optional)

2h

- 4 If making up primer pools from individual oligos fully resuspend lyophilised oligos in 1xTE to a concentration of 100 micromolar (μM) , vortex thoroughly and spin down.

Note

If using IDT ARTIC nCoV-2019 V3 Panel (100 micromolar (μM)) skip to step 6.

- 5 Sort all odd regions primers into one or more tube racks. Add 5 μL of each odd region primer to a 1.5 mL Eppendorf tube labelled "Pool 1 (100 micromolar (μM))". Repeat the process for all even region primers for Pool 2. These are your 100 micromolar (μM) stocks of each primer pool.

Note

Primers should be diluted and pooled in the **mastermix** cabinet which should be cleaned with decontamination wipes and UV sterilised before and after use.

**Note**

For more information see Figure 2 in;

Quick, J. et al. Multiplex PCR method for MinION and Illumina sequencing of Zika and other virus genomes directly from clinical samples. Nat Protoc 12, 1261–1276 (2017).
<https://doi.org/10.1038/nprot.2017.066>

- 6 Dilute 100 micromolar (μM) pools 1:10 in molecular grade water, to generate 10 micromolar (μM) primer stocks.

Note

Primers are used at a final concentration of 15 nanomolar (nM) per primer. In this case V3 pools have 110 primers in pool 1 and 108 primers in pool 2. so the requirement is ~ 4 μL primer pool (10 micromolar (μM)) per 25 μL reaction.

Note

Make up multiple 100 μL aliquots of 10 micromolar (μM) primer dilutions and freeze them in case of degradation or contamination.

Multiplex PCR

4h

- 7 Set up the two PCR reactions per sample as follows in strip-tubes or plates. Gently mix by pipetting and pulse spin the tube to collect liquid at the bottom of the tube.

Component	Reaction 1	Reaction 2
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
	V3 Pool 1 (10µM)	4 µL	0 µL
	V3 Pool 2 (10µM)	0 µL	4 µL
	Nuclease-free water	12.75 µL	12.75 µL
	Total	22.5 µL	22.5 µL

For M0493

or

Component	Reaction 1	Reaction 2
Q5 Hot Start High-Fidelity 2X Master Mix	12.5 µL	12.5 µL
V3 Pool 1 (10µM)	4 µL	0 µL
V3 Pool 2 (10µM)	0 µL	4 µL
Nuclease-free water	6 µL	6 µL
Total	22.5 µL	22.5 µL

For M0494

Note

Q5 Hot Start High-Fidelity 2X Master Mix can also be used instead of the component kit. Half-scale PCR reactions can also be used to save costs as you will only require

 2.5 µL for downstream steps.

Note

To prevent pre-PCR contamination the mastermix for each pool should be made up in the **mastermix** cabinet which should be cleaned with decontamination wipes and UV sterilised before and after use and aliquoted into PCR strip-tubes/plate



- 8 Add  2.5 μL cDNA to each of the PCR reactions, gently mix by pipetting and pulse spin the tube to collect liquid at the bottom of the tube.

Note

Up to  5 μL cDNA can be added to each PCR reaction (in place of nuclease-free water) to improve amplification of low titre samples. Using  5 μL cDNA will require a  20 μL cDNA reaction and may be more likely to cause inhibition so use cautiously.

Note

cDNA should be added in the **extraction and sample addition** cabinet which should be cleaned with decontamination wipes and UV sterilised before and after use.

- 9  0 μL Set-up the following program on the thermal cycler:

Step	Temperature	Time	Cycles
Heat Activation	 98 °C	 00:00:30	1
Denaturation	 98 °C	 00:00:15	25-35
Annealing	 65 °C	 00:05:00	25-35
Hold	 4 °C	Indefinite	1

Note

Cycle number should be 25 for Ct 18-21 up to a maximum of 35 cycles for Ct 35.



Note

Thermocycler calibration can vary instrument to instrument. If you see amplicon 64 dropout then decrease the annealing/extension temperature to 63°C . Denaturation temperature of 95°C can also be used and may slightly increase PCR yields.

PCR dilution

10m

- 10 Label strip-tubes/plate and combine the following volumes of each PCR reaction for $10\ \mu\text{L}$ each sample:

Component	Volume
Pool 1 PCR reaction	2.5 μL
Pool 2 PCR reaction	2.5 μL
Nuclease-free water	45 μL
Total	50 μL

Note

The PCR post-clean up concentration is typically around $100\ \text{Mass Percent}$. This means we can pool them without quantification/normalisation to make a significant time saving. If you require very even barcode representation perform clean-up and normalise to $10\ \text{Mass Percent}$ then continue.

Note

Amplicons should be added in the **post-PCR** cabinet which should be cleaned with decontamination wipes and UV sterilised before and after use.



Native barcoding

2h

- 11 Barcode the amplicon pools using the one-pot native barcoding approach.

Protocol

NAME

One-pot native barcoding of amplicons v3 (LoCost)

CREATED BY

Josh Quick

[Preview](#)

- 11.1 In a new PCR strip-tube/plate set up the following reaction for each sample:

Component	Volume
PCR dilution from previous step	3.3 μL
Ultra II End Prep Reaction Buffer	1.2 μL
Ultra II End Prep Enzyme Mix	0.5 μL
Nuclease-free water	5 μL
Total	10 μL

Note

Make a master mix of end-preparation reagents and nuclease-free water and aliquot into strip-tube/plate to improve reproducibility.

- 11.2 Incubate at room temperature for 00:15:00

Incubate at 65 °C for 00:15:00



Incubate on ice for  00:01:00

11.3 In a new PCR strip-tube/plate set up the following reaction for each sample:

Component	Volume
End-preparation reaction mixture	0.75 μL
NBXX barcode	1.25 μL
Blunt/TA Ligase Master Mix	5 μL
Nuclease-free water	3 μL
Total	10 μL

Note

Use one native barcode from the EXP-NBD104 (1-12), EXP-NBD114 (13-24) or EXP-NBD196 per sample. Use 12 or more barcodes per library or there will be insufficient total material to achieve good yields.

11.4 Incubate at room temperature for  00:20:00

Incubate at  65 °C for  00:10:00

Incubate on ice for  00:01:00

Note

The 65°C incubation is to inactivate the DNA ligase to prevent barcode cross-ligation when reactions are pooled in the next step.

11.5 In a new  1.5 mL Eppendorf tube pool all one-pot barcoding reactions together.

**Note**

If processing 12-24 samples pool all  10 μL from each native barcoding reaction.
if processing 48 samples pool  5 μL from each native barcoding reaction.
If processing 96 samples pool  2.5 μL from each native barcoding reaction so as not to exceed a pool volume of  240 μL which would make the clean-up volume too large.

- 11.6 Add 0.4x volume of SPRI beads to the sample tube and mix gently by either flicking or pipetting. For example add  96 μL SPRI beads to  240 μL pooled one-pot barcoding reactions.

Note

0.4x volume of SPRI is sufficient to bind 400 bp amplicons in the presence of ligation buffer, do not use 1x as this will result in an excessive large bead pellet.

- 11.7 Mix by vortexing and pulse centrifuge to collect all liquid at the bottom of the tube. Incubate for  00:05:00 at room temperature.
- 11.8 Place on magnetic rack and incubate for  00:02:00 or until the beads have pelleted and the supernatant is completely clear. Carefully remove and discard the supernatant, being careful not to touch the bead pellet.
- 11.9 Add  250 μL SFB and resuspend beads completely by pipette mixing. Pulse centrifuge to collect all liquid at the bottom of the tube and place on the magnet. Remove supernatant and discard.

Note

SFB will remove excess adapter without damaging the adapter-protein complexes. Do not use 70% ethanol as in early clean-ups.



- 11.10 Repeat steps 11.9 to perform a second SFB wash. Pulse centrifuge and remove any residual SFB.

Note

You do not need to allow to air dry with SFB washes.

- 11.11 Add  200 μL of room-temperature  70 % volume ethanol to bathe the pellet. Carefully remove and discard ethanol, being careful not to touch the bead pellet.

Note

Only perform 1x 70% ethanol wash

- 11.12 Pulse centrifuge to collect all liquid at the bottom of the tube and carefully remove as much residual ethanol as possible using a P10 pipette.

- 11.13 With the tube lid open incubate for  00:01:00 or until the pellet loses it's shine (if the pellet dries completely it will crack and become difficult to resuspend).

- 11.14 Resuspend pellet in  30 μL  10 millimolar (mM) Tris pH 8.0, mix gently by either flicking or pipetting and incubate for  00:02:00 .

- 11.15 Place on magnet and transfer sample to a clean  1.5 mL Eppendorf tube ensuring no beads are transferred into this tube.

- 12 Quantify  1 μL of the barcoded amplicons using the Quantus Fluorometer using the ONE dsDNA assay. Concentration will vary depending on number and Ct of samples and but you need about  30 ng total at this stage to acheive maximum run yield.



Protocol

NAME

DNA quantification using the Quantus fluorometer

CREATED BY

Josh Quick

[Preview](#)

- 12.1 Remove Lambda DNA 400 ng/ μ L standard from the freezer and leave on ice to thaw. Remove ONE dsDNA dye solution from the fridge and allow to come to room temperature.

 QuantiFluor(R) ONE dsDNA System, 500rxn **Promega Catalog #E4870**

- 12.2 Set up two  0.5 mL tubes for the calibration and label them 'Blank' and 'Standard'

- 12.3 Add  200 μ L ONE dsDNA Dye solution to each tube.

- 12.4 Mix the Lambda DNA standard 400 ng/ μ L standard by pipetting then add  1 μ L to one of the standard tube.

- 12.5 Mix each sample vigorously by vortexing for  00:00:05 and pulse centrifuge to collect the liquid.

- 12.6 Allow both tubes to incubate at room temperature for  00:02:00 before proceeding.

- 12.7 Selection 'Calibrate' then 'ONE DNA' then place the blank sample in the reader then select 'Read Blank'. Now place the standard in the reader and select 'Read Std'.



- 12.8 Set up the required number of  0.5 mL tubes for the number of DNA samples to be quantified.

Note

Use only thin-wall, clear, 0.5mL PCR tubes such as Axygen #PCR-05-C

- 12.9 Label the tubes on the lids, avoid marking the sides of the tube as this could interfere with the sample reading.

- 12.10 Add  199 μ L ONE dsDNA dye solution to each tube.

- 12.11 Add  1 μ L of each user sample to the appropriate tube.

Note

Use a P2 pipette for highest accuracy.

- 12.12 Mix each sample vigorously by vortexing for  00:00:05 and pulse centrifuge to collect the liquid.

- 12.13 Allow all tubes to incubate at room temperature for  00:02:00 before proceeding.

- 12.14 On the Home screen of the Quantus Fluorometer, select 'Protocol', then select 'ONE DNA' as the assay type.

Note

If you have already performed a calibration for the selected assay you can continue, there is no need to perform repeat calibrations when using ONE DNA pre diluted dye solution. If you want to use the previous calibration, skip to step 11. Otherwise, continue with step 9.

- 12.15 On the home screen navigate to 'Sample Volume' and set it to  1 µL then 'Units' and set it to ng/µL.
- 12.16 Load the first sample into the reader and close the lid. The sample concentration is automatically read when you close the lid.
- 12.17 Repeat step 16 until all samples have been read.
- 12.18 The value displayed on the screen is the dsDNA concentration in ng/µL, carefully record all results in a spreadsheet or laboratory notebook.
- 13 Set up the following AMII adapter ligation and clean-up with SFB.

Protocol

NAME

Adapter ligation with AMII v2

CREATED BY

Josh Quick

Preview

- 13.1 In a new  1.5 µL Eppendorf tube set up the following AMII adapter ligation reaction.

Component	Volume
Barcoded amplicon pool	30 µL
NEBNext Quick Ligation Reaction Buffer (5X)	10 µL
Adapter Mix (AMII)	5 µL
Quick T4 DNA Ligase	5 µL
Total	50 µL



13.2 Incubate at room temperature for  00:20:00

13.3 Add  50 μL (1:1) of SPRI beads to the sample tube and mix gently by either flicking or pipetting. Pulse centrifuge to collect all liquid at the bottom of the tube.

Note

Vortex SPRI beads thoroughly before use to ensure they are well resuspended, the solution should be a homogenous brown colour.

Note

There will be some variation in clean-up efficiencies but expect to carry around 50% through this clean-up

13.4 Incubate for  00:05:00 at room temperature.

13.5 Place on magnetic rack and incubate for  00:02:00 or until the beads have pelleted and the supernatant is completely clear. Carefully remove and discard the supernatant, being careful not to touch the bead pellet.

13.6 Add  250 μL SFB and resuspend beads completely by pipette mixing. Pulse centrifuge to collect all liquid at the bottom of the tube. Remove supernatant and discard.

Note

SFB will remove excess adapter without damaging the adapter-protein complexes. Do not use 70% ethanol as in early clean-ups.

13.7 Repeat steps 13.6 to perform a second SFB wash.



- 13.8 Pulse centrifuge and remove any residual SFB. Add  15 μL EB (ONT) and resuspend beads by pipette mixing.

Note

You do not need to allow to air dry with SFB washes.

- 13.9 Incubate at room temperature for  00:02:00 .

- 13.10 Place on magnetic rack until clear. Transfer final library to a new 1.5mL Eppendorf tube.

- 14 Quantify  1 μL of the final library using the Quantus Fluorometer using the ONE dsDNA assay. Concentration will vary depending on number and Ct of samples but  15 ng final library is usually required to achieve maximum run yield.

Protocol

NAME

DNA quantification using the Quantus fluorometer

CREATED BY

Josh Quick

Preview

Note

Final library can be now be stored in  10 millimolar (mM) Tris  8 at  4 $^{\circ}\text{C}$ for up to a week if needed otherwise proceed directly to MinION sequencing.

- 14.1 Remove Lambda DNA 400 ng/ μL standard from the freezer and leave on ice to thaw. Remove ONE dsDNA dye solution from the fridge and allow to come to room

temperature.

 QuantiFluor(R) ONE dsDNA System, 500rxn **Promega Catalog #E4870**

- 14.2 Set up two  0.5 mL tubes for the calibration and label them 'Blank' and 'Standard'
- 14.3 Add  200 μ L ONE dsDNA Dye solution to each tube.
- 14.4 Mix the Lambda DNA standard 400 ng/ μ L standard by pipetting then add  1 μ L to one of the standard tube.
- 14.5 Mix each sample vigorously by vortexing for  00:00:05 and pulse centrifuge to collect the liquid.
- 14.6 Allow both tubes to incubate at room temperature for  00:02:00 before proceeding.
- 14.7 Selection 'Calibrate' then 'ONE DNA' then place the blank sample in the reader then select 'Read Blank'. Now place the standard in the reader and select 'Read Std'.
- 14.8 Set up the required number of  0.5 mL tubes for the number of DNA samples to be quantified.
- #### Note
- Use only thin-wall, clear, 0.5mL PCR tubes such as Axygen #PCR-05-C
- 14.9 Label the tubes on the lids, avoid marking the sides of the tube as this could interfere with the sample reading.
- 14.10 Add  199 μ L ONE dsDNA dye solution to each tube.



14.11 Add  1 μL of each user sample to the appropriate tube.

Note

Use a P2 pipette for highest accuracy.

14.12 Mix each sample vigorously by vortexing for  00:00:05 and pulse centrifuge to collect the liquid.

14.13 Allow all tubes to incubate at room temperature for  00:02:00 before proceeding.

14.14 On the Home screen of the Quantus Fluorometer, select 'Protocol', then select 'ONE DNA' as the assay type.

Note

If you have already performed a calibration for the selected assay you can continue, there is no need to perform repeat calibrations when using ONE DNA pre diluted dye solution. If you want to use the previous calibration, skip to step 11. Otherwise, continue with step 9.

14.15 On the home screen navigate to 'Sample Volume' and set it to  1 μL then 'Units' and set it to ng/ μL .

14.16 Load the first sample into the reader and close the lid. The sample concentration is automatically read when you close the lid.

14.17 Repeat step 16 until all samples have been read.

14.18 The value displayed on the screen is the dsDNA concentration in ng/ μL , carefully record all results in a spreadsheet or laboratory notebook.

MinION sequencing

1d

15 Prime the flowcell and load  15 ng sequencing library onto the flowcell.



Protocol

NAME

Priming and loading a MinION flowcell v2

CREATED BY

Josh Quick

Preview

Note

From experience we know  15 ng is optimum loading input for short amplicons. Speed drop during the run indicates excessive library was loaded. Low run yield <20M reads indicates insufficient library.

15.1 Thaw the following reagents at room temperature before placing on ice:

Sequencing buffer (SQB)

Loading beads (LB)

Flush buffer (FLB)

Flush tether (FLT)

15.2 Add  30 µL FLT to the FLB tube and mix well by vortexing.

15.3 If required place a new MinION flowcell onto the MinION by flipping open the lip and pushing one end of the flowcell under the clip and pushing down gently.

15.4 Rotate the inlet port cover clockwise by 90° so that the priming port is visible.

15.5 Take a P1000 pipette and tip and set the volume to  800 µL . Place the tip in the inlet port and holding perpendicularly to the plane of the flowcell remove any air from the inlet port by turning the volume dial anti-clockwise.

Note

Be careful not to remove so much volume that air is introduced onto the rectangular array via the outlet.

15.6 Load  800 μL of FLB (plus FLT) into the flow cell via the inlet port, dispense slowly and smoothly trying to avoid the introduction of any air bubbles.

15.7 Wait for  00:05:00 .

15.8 Gently lift the SpotON cover to open the SpotON port.

15.9 Load another  200 μL of FLB (plus FLT) into the flow cell via the inlet port, this will initiate a siphon at the SpotON port to allow you to load the library dilution.

15.10 In a new tube prepare the library dilution for sequencing:

	Component	Volume
	SQB	37.5 μL
	LB	25.5 μL
	Library	12 μL
	Total	75 μL

Note

Mix LB immediately before use as they settle quickly.

Make up with EB if less than 12 μL library is required.



- 15.11 Mix the prepared library gently by pipetting up and down just prior to loading.
- 15.12 Add the  75 µL library dilution to the flow cell via the SpotON sample port in a dropwise fashion. Ensure each drop siphons into the port before adding the next.
- 15.13 Gently replace the SpotON sample port cover, making sure the bung enters the SpotON port, close the inlet port and close the MinION lid.
- 16 Start the sequencing run using MinKNOW.

Protocol

NAME

Starting a MinION sequencing run using MinKNOW

CREATED BY

Josh Quick

Preview

Note

If using Live basecalling ensure to turn on double-ended barcoding in the basecalling settings.

- 16.1 If required plug the MinION into the computer and wait for the MinION and flowcell to be detected.
- 16.2 Choose flow cell 'FLO-MIN106' from the drop-down menu.
- 16.3 Then select the flowcell so a tick appears.



16.4 Click the 'New Experiment' button in the bottom left of the screen.

16.5 On the New experiment popup screen, select the running parameters for your experiment from the individual tabs:

Experiment: Name the run in the experiment field, leave the sample field blank.

Kit: Selection: Select LSK109 as there is no option for native barcoding (NBD104).

Run Options: Set the run length to 6 hours (you can stop the run once sufficient data has been collected as determined using RAMPART).

Basecalling: Leave basecalling turned but select 'fast basecalling'.

Output: The number of files that MinKNOW will write to a single folder. By default this is set to 4000 but can be reduced to make RAMPART update more frequently.

Click 'Start run'.

16.6 Monitor the progress of the run using the MinKNOW interface.