"Midnight" SARS-CoV2 genome sequencing protocol using 1200bp amplicon primer set v2 and the Nanopore Rapid library kit

Forked from SARS-CoV2 genome sequencing protocol (1200bp amplicon "midnight" primer set, using Nanopore Rapid kit)

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ABSTRACT

This protocols is part of the ANU Biosecurity mini-research project #2 "An SARS-COV2 incursion scenario: Genomics, phylogenetics, and incursions." This mini-research project is modeled on the yearly Quality Assurance Program of The Royal College of Pathologists of Australia (RCPAQAP), we take part in together with ACT Pathology.

This research project is split into two major parts, identical to how the official RCPAQAP is run every year.

Part #1 is focusing on the 'wet-lab' by sequencing SARS-COV2 from real world RNA samples provided by ACT Pathology especially for our ANU biosecurity course (Thank YOU!). Here you will amplify and sequence five (5) RNA samples per research group. You will assess the SARS-COV2 genome sequences for their lineage assignments using online programs, put sequences into a global context, estimate the collection date based on genetic information, and describe mutations in the spike protein.

Part #2 is focusing on the 'dry-lab' by investigating a hypothetical incursion scenario in the so-called city Fantastica. You will combine genomic surveillance of SARS-COV2 with case interview data to trace the spread into of SARS-COV2 in the community and into high risk settings. We will provide you with real publicly available SARS-COV2 genome and fantasized case interviews. You will put these two together to trace the spread and suggest potential improvements in containment strategies with a focus on high risk settings.

This protocol describes the wet-lab component of Part #1. It is an adaption (fork) of
a previous protocol published by Nikki Freed and Olin Silander during the early days of the pandemic here and here. They designed the protocol to enable faster, easier sequencing of SARS-COV2 genomes with fewer steps than previous methods, we use multiplexed 1200 base pair PCR amplicons with the Oxford Nanopore RAPID barcoding kit.

The original publication that describes this protocol is here along with important considerations is here:
https://academic.oup.com/biometrika/article/5/1/bpaa014/5873518?login=true

Primers were all designed using Primal Scheme: http://primal.zibraproject.org/, described here https://www.nature.com/articles/nprot.2017.066.


The final goal is to achieve the following:

Week 7:
- cDNA reverse transcription of RNA into DNA.
- Pooled amplification of the SARS-COV2 genomes chunked into ~1200bp amplicons with non-overlapping pools A and B.

Week 8:
- Library preparation using the Nanopore Rapid barcoding kit.
- Sequencing your own library on a Nanopore flongel.

This protocol is applicable for week 7 and 8.
GUIDELINES

You must have read, understood, and follow the health and safety instructions provided in the "Overview Mini-Research Project #2 BIOL3106/6106" provided on Wattle (ANU learning portal).

You must have signed and returned one copy of the "Student Safety Declaration Form For Practical Class Work" before starting any laboratory work.

You must have read and understood the Hazard Sheets (Risk assessment) of all chemicals listed below in the "Safety Warnings" section. These Hazard Sheets are provided on Wattle as part of the "Overview Mini-Research Project #2 BIOL3106/6106" document.

MATERIALS

PRIMERS


REVERSE TRANSCRIPTION

- A PCR strip of 8 with six tubes containing 2uL each.

PCR

- Q5 High-Fidelity 2X Master Mix; Week 7: Two tubes with 81.25 ul (6.5*12.1u) each for PCR Pool A and Pool B.

CLEAN UP

- One 1.5ml tube with 500ul for each library prep.

SEQUENCING PREPARATION FOR OXFORD NANOPORE

- Rapid barcoding kit SQK-RBK110.96 https://store.nanoporetech.com/rapid-barcoding-kit-1.html
- Flongle https://nanoporetech.com/products/flongle

Consumables

- 1.5 ml tubes. Week 7: One for each PCR pool master mix. Week 8: One for each PCR reaction pooled (A+B) and one for the dilution.
■ PCR strip tubes of 8. Week 7: Three strips, one for cDNA and two for the PCR pools A and B. Week 8: One stripe for the barcoding reaction.
■ Primers @ 10uM. PoolA and PoolB.
■ Nuclease Free water
■ Six tubes with 198ul broad range qubit.
■ Ice bucket with ice.

**Equipment needed**
■ Benchtop centrifuge for 1.5ml/2ml tubes.
■ Mini benchtop spinner for PCR strip tubes of 8.
■ Vortex.
■ PCR machines.
■ Laptops 9x.
■ MinIONs 9x.
■ Flongle adapters 9x.

**SAFETY WARNINGS**

This protocol does not require any hazardous substances or infectious agents. However, it includes RNA extracted from human swap samples.

You need to wear safety equipment at all times including lab coats, gloves, and safety goggles when handling chemicals and biological agents. While the extracted RNA does not include any infectious agents treat them with care. Do not remove them from the laboratory. Use a dedicated notebook and pen to make notes during the mini-research project. Do not put anything into your mouth while in the laboratory. Wash your hands each time you leave the laboratory.

**BEFORE START INSTRUCTIONS**

You must study the protocol carefully before you start. If anything is unclear post questions directly here on protocols.io.

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**Week 7: cDNA synthesis and two pooled PCR reactions**

1 In week 7 you will perform the cDNA synthesis and PCR reactions on five samples and one negative control. You will be provided with five samples of 10 ul each and one tube of negative control.
You will receive a PCR strip of eight with 6 tubes containing 2uL Lunascript RT supermix. Label the tubes with 1-6 and your RG name. Note down which sample corresponds to which number.

Mix the following components in an 0.2mL 8-strip tube on ice:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>LunaScript® RT SuperMix Kit (5x)</td>
<td>2 µL</td>
</tr>
<tr>
<td>Template RNA</td>
<td>8 µL</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>10 µL</strong></td>
</tr>
</tbody>
</table>

This means add 8uL of sample to each tube separately. Gently mix by pipetting up and down during the adding of each sample. Change the tip between each sample addition. Once you added all samples close the lid.

Pulse spin the tube to collect liquid at the bottom of the tube.

Incubate the reaction in a PCR machine:

- **25 °C** 00:02:00
- **55 °C** 00:10:00
- **95 °C** 00:01:00
- Hold at **20 °C**

You will have to prepare two PCR master mixes one for each primer pool A and B. For this, you will have two tubes of 81.25uL Q5 hot start master mix provided. Label each tube with pool A or B and your RG name.

<table>
<thead>
<tr>
<th>Component and volume per sample</th>
<th>Pool A</th>
<th>Pool B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q5® Hot Start HF 2x Master Mix</td>
<td>12.5 µL</td>
<td>12.5 µL</td>
</tr>
</tbody>
</table>
Primer Pool A (10µM)  
1.1 µL  
Primer Pool B (10µM)  
0 µL  
Nuclease-free water  
8.9 µL  

**Total**  
22.5 µL

Considering you have five samples and one negative control, generate the following two master mixes.

<table>
<thead>
<tr>
<th></th>
<th>Pool A</th>
<th>Pool B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q5® Hot Start HF 2x Master Mix (two tubes provided)</td>
<td>81.25ul</td>
<td>81.25ul</td>
</tr>
<tr>
<td>Primer Pool A (10µM)</td>
<td>7.15ul</td>
<td>0</td>
</tr>
<tr>
<td>Primer Pool B (10µM)</td>
<td>0</td>
<td>7.15ul</td>
</tr>
<tr>
<td>Nuclease-free water</td>
<td>57.85ul</td>
<td>57.85ul</td>
</tr>
</tbody>
</table>

Once you added all ingredients, mix by vortexing and spin down to collect the liquid. Place on ice.

6. Label two PCR strips of eight with the PCR pool A or B, number 1-6, and your RG name. Aliquot out 22.5 ul of each master mix (Pool A or B) into the corresponding PCR strips.

7. Collect your cDNA containing strip tubes (the samples will have a blue color). Spin them down briefly.

   add 2.5 µL each cDNA sample [1-6] to a tube containing 22.5 µL Pool A that is labelled accordingly. Mix well by pipetting up and down. Change tip after each addition.

   Add 2.5 µL each cDNA sample [1-6] to a tube containing 22.5 µL Pool B that is labelled accordingly. Mix well by pipetting up and down. Change tip after each addition.

Leave the strip tubes on ice. Close the tubes with a lid when done adding all samples.
8. Pulse centrifuge the tubes to collect the contents at the bottom of the tube.

9. Set-up the following program on the thermal cycler:

<table>
<thead>
<tr>
<th>Step</th>
<th>Temperature</th>
<th>Time</th>
<th>Cycles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat Activation</td>
<td>98 °C</td>
<td>00:00:30</td>
<td>1</td>
</tr>
<tr>
<td>Denaturation</td>
<td>98 °C</td>
<td>00:00:15</td>
<td>35</td>
</tr>
<tr>
<td>Annealing and Extension</td>
<td>65 °C</td>
<td>00:05:00</td>
<td>35</td>
</tr>
<tr>
<td>Hold</td>
<td>20 °C</td>
<td>Indefinite</td>
<td>1</td>
</tr>
</tbody>
</table>

Expected result

Final concentrations of PCR products typically range from ~5 - 150ng/ul as measured by Qubit.

Hand over your PCR reactions to demonstrators as they will run them till next week.

Week 8: Library preparation and sequencing

10. In week 8 you will perform your own Nanopore sequencing library preparation and sequencing. You will do this by using your PCR reactions from week 7.

Pooling and PCR quantification

11. Label a 1.5 mL Eppendorf tube with sample name and RG name. In total this will be six tubes.

For each sample combine the PCR reactions (pool A and B) by pipetting all 25ul of each pool into one tube. This means each 1.5 ml eppendorf tube will contain 50ul in total.

Mix by vortexing and spin down samples briefly.
11.1 Quantify DNA using a Qubit.

You will be provided with 6 tubes of 198ul Qubit HS solution. Label the tubes with the numbers 1-6. Add 2uL of each combined PCR reactions into one Qubit tube. Vortex briefly, hand centrifuge, incubate for 2 mins, and measure with the Qubit machine. Note down your concentration in ng/ul for each sample.

Normalisation

12 Label a 1.5 eppendorf tube for each sample so you can generate a stock PCR solution of 13.33 ng/ul the final volume being 20uL.

Each tube must be labelled as stock 13.33 ng/ul, sample name, and RG name.

The important formula here is:

\[ n = c \times V \]

with \( n \) being the amount in ng, \( c \) the concentration in ng/ul, and \( V \) the volume.

You want \( V = 20\text{ul} \) and \( c = 13.33 \text{ ng/ul} \). This means you will have 267ng in 20 ul.

The question is how much volume of the original PCR do you need to obtain 267ngs?

Here now \( n = 267\text{g} \) and \( c \) equals your measured concentration. Let’s assume your measured \( c = 56 \text{ ng/ul} \) for an assumed sample. Than we can resolve the equation as follows.

\[ n = c \times V \rightarrow V = \frac{n}{c} \rightarrow 267\text{ng}/56\text{ng/ul} = 4.7 \text{ ul} \]

So you need to use 4.7ul in 20ul total volume to obtain 267ng. This means you can combine 13.7ul H20 with 4.7ul of your combined PCR sample. Vortex it and spin down.

In case you concentration is < 13.33 ng/ul use 20ul of sample as stock without dilution.
Multiple samples can be run on the same flow cell by barcoding. Up to 96 samples at a time can be sequenced using the RBK110.96 kit. Amplicons from each sample will be individually barcoded in the following steps.

You will receive a PCR strip of eight with six tubes containing 2.5ul of the fragmentation mix each. Label the strip with sample number [1-6] and RG name.

13.1 Add 7.5 µL of each diluted PCR stock [13.33 ng/ul] reaction from step 12 to the labeled PCR tube.

Set up the following reaction for each sample:

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA amplicons from step 12 (100ng total)</td>
<td>7.5 µL</td>
</tr>
<tr>
<td>Fragmentation Mix (one for each sample, already aliquoted for you)</td>
<td>2.5 µL</td>
</tr>
<tr>
<td>Total</td>
<td>10 µL</td>
</tr>
</tbody>
</table>

Important keep samples on ice!

Mix gently by pipetting up and down briefly, flick the tubes, and spin down.

13.2 Incubate the reaction in a PCR machine:

- 30 °C for 00:01:00
- 80 °C for 00:01:00
- 4 °C for 00:00:30

13.3 From here on out you will combine your barcoded samples with one other RG on your bench. Each final library will contain 12 samples to be sequenced together. [With the exception of RG0 who have a single flongel to themselves]. RG1+2, RG3+4 and so on will go together.

For each pair of RGs, label one 1.5 ml Eppendorf tube with BC and RG names. Combine all your barcoded samples in this tube. This tube will contain 120ul total. [RG0 will have 60ul]

Mix by vortexing and spin down.

14 SPRI Bead Cleanup. Use a 1:1 ratio of sample to beads. This means in case of 120ul combined
barcoded samples add 120ul of beads. **Ask for help of a demonstrator as needed.**

14.1 Add 120uL of SPRI beads to your combined barcoded samples. Mix by flicking the tube. Hand centrifuge to collect all liquid in the bottom of the tube while the brown beads stays suspended.

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

14.3 Place tube on magnetic rack and incubate for 00:03:00 until the beads have pelleted towards the magnet and the supernatant is completely clear.

14.4 Leave tube on the magnet and pipette off the supernatant, being careful not to touch the bead pellet. The DNA is now bound to the beads.

14.5 Add 1ml of freshly prepared room-temperature 80 % volume ethanol to wash the pellet.

14.6 **go to step #14.4** and repeat one more ethanol wash.

14.7 Leave tube on the magnet and pipette off the supernatant, being careful not to touch the bead pellet. The DNA is now bound to the beads.

14.8 Pulse centrifuge to collect all liquid at the bottom of the tube, place tube back to the magnet, and carefully remove as much residual ethanol as possible using a P10 pipette.
14.9 With the tube lid open incubate for 00:03:00 to let the pellet try.

14.10 Remove the tube from the magnetic rack. Resuspend pellet in 15 µL Elution Buffer (EB). Incubate for 00:03:00 at RT. with flicking the tube.

Label a new 1.5ml tube with BCD and your RG names.

14.11 Place on magnet, incubate for 00:03:00 so that all beads are on the side of the tube towards the magnet. Transfer 10 µL to a clean labelled 1.5mL Eppendorf tube ensuring no beads are transferred into this tube.

15 Label a new 1.5ml Eppendorf tube with L and your RG name.

Transfer 5 µL of your cleaned combined barcoded samples into this tube.

Ask a demonstrator to add 0.5 µL of Rapid Adapter F (RAP-F) to 5 µl of barcoded DNA.

Mix gently by flicking the tube, and spin down.

Incubate the reaction for 00:05:00 at RT.

16 Now prepare the final library to be loaded by combining the following a 1.5ml tube labelled FL and your RG name.

- 15 µL Sequencing Buffer II (SBII)
- 10 µL Loading Beads II (LBII) mixed immediately before use
- 2.5 µL EB buffer
- 2.5 µL of RAP-F adapted barcoded DNA from step 15.

Place library to the side till you are ready to load in step 23.

Loading the library onto a Flongel
Please familiarize yourself with how to load a flongel in this video [https://nanoporetech.com/resource-centre/video/ncm22/how-to-load-a-flongle-flow-cell](https://nanoporetech.com/resource-centre/video/ncm22/how-to-load-a-flongle-flow-cell) [0-12 minutes only]. **Ask a demonstrator to help with the loading!**

Place the Flongle adapter into the MinION.

**IMPORTANT**
Do NOT touch the reverse side of the Flongle flow cell array or the contact pads on the Flongle adapter. ALWAYS wear gloves when handling Flongle flow cells and adapters to avoid damage to the flow cell or adapter.

The diagram below shows the components of the Flongle flow cell:

The seal tab, air vent, waste channel, drain port and sample port are visible here. The sample port, drain port and air vent only become accessible once the seal tab is peeled back.

The adapter should sit evenly and flat on the MinION. This ensures the flow cell assembly is flat during the next stage.
19 Plug in the MinION contain the flongle adapter in the computer.

**IMPORTANT**
The adapter needs to be plugged into your device, and the device should be plugged in and powered on before inserting the Flongle flow cell.

20 Place the flow cell into the Flongle adapter, and press the flow cell down until you hear a click. The flow cell should sit evenly and flat inside the adapter, to avoid any bubbles forming inside the fluidic compartments.

21 Take the tube labelled with FL that contains 117 μl of Flush Buffer (FB) with 3 μl of Flush Tether (FLT). Mix it by flicking and collect it at the bottom of the tube.

Peel back the seal tab from the Flongle flow cell, up to a point where the sample port is exposed, as follows:

1. Lift up the seal tab:
2. Pull the seal tab to open access to the sample port:

3. Hold the seal tab open by using adhesive on the tab to stick to the MinION Mk 1B lid:
22 Take the 120ul of FL. To prime your flow cell with the mix of Flush Buffer (FB) and Flush Tether (FLT) that was prepared earlier, ensure that there is no air gap in the sample port or the pipette tip. Place the P200 pipette tip inside the sample port and slowly dispense the priming fluid into the Flongle flow cell by slowly pipetting down. We also recommend twisting the pipette plunger down to avoid flushing the flow cell too vigorously.

23 Take the final library from step 16 and mix by flicking so it is milky again. Collect by a quick hand centrifuge. Take the 30 ul and load.

To add the final library to the flow cell, ensure that there is no air gap in the sample port or the pipette tip. Place the P200 tip inside the sample port and slowly dispense the Sequencing Mix
into the flow cell by slowly pipetting down. We also recommend twisting the pipette plunger
down to avoid flushing the flow cell too vigorously.

24 Seal the Flongle flow cell using the adhesive on the seal tab, as follows:
1. Stick the transparent adhesive tape to the sample port.

2. Replace the top (Wheel icon section) of the seal tab to its original position.
Close the MinION lid.

**Data acquisition and basecalling**

25 Open the MinKNOW software on the laptop and get a demonstrator to help you get the sequencing run started.