

# Rapid, Sensitive, Full-Genome Sequencing of Severe Acute Respiratory Syndrome Coronavirus 2

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We describe validated protocols for generating high-quality, full-length severe acute respiratory syndrome coronavirus 2 genomes from primary samples. One protocol uses multiplex reverse transcription PCR, followed by MinION or MiSeq sequencing; the other uses singleplex, nested reverse transcription PCR and Sanger sequencing. These protocols enable sensitive virus sequencing in different laboratory environments.

In December 2019, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the etiologic agent of coronavirus disease 2019 (COVID-19), emerged in Wuhan, China. Since then, it has rapidly spread worldwide (1–3), causing 7,039,918 confirmed cases, including 404,396 deaths, in 188 countries or regions as of June 9, 2020 (4). Because SARS-CoV-2 has shown the capacity to spread rapidly and lead to a range of manifestations in infected persons, from asymptomatic infection to mild, severe, or fatal disease, it is essential to identify genetic variants to track spread and understand any changes in transmissibility, tropism, and pathogenesis.

We describe the design and use of 2 PCR-based methods for sequencing SARS-CoV-2 clinical specimens. The first is a multiplex PCR panel, followed by sequencing on either the Oxford Nanopore MinION apparatus (<https://nanoporetech.com>) or an Illumina MiSeq apparatus (<https://www.illumina.com>). When coupled with MinION sequencing, our protocol can be implemented outside a traditional laboratory and can be completed in a single workday, similar to previous mobile genomic surveillance of Ebola and Zika virus outbreaks (5,6). In

addition, we provide a complementary singleplex, nested PCR strategy, which improves sensitivity for samples with lower viral load and is compatible with Sanger sequencing.

## The Study

On January 10, 2020, the first SARS-CoV-2 genome sequence was released online (7). That day, we designed 2 complementary panels of primers to amplify the virus genome for sequencing.

For the first panel, we used the PRIMAL primer design tool (5) to design multiplex PCRs to amplify the genome by using only a few PCRs (Appendix, <https://wwwnc.cdc.gov/EID/article/26/10/20-1800-App1.pdf>). The final design consists of 6 pools of primers optimized for sensitivity and assay flexibility. The amplicons average 550 bp with 100-bp overlaps to enable sequencing on either the Oxford MinION or Illumina MiSeq.

For the second panel, we designed sets of primers to generate nested, tiling amplicons across the SARS-CoV-2 genome (Appendix) for enhanced sensitivity in samples with lower viral loads. Each amplicon is 322–1,030 bp with an average overlap of 80 bp. These amplicons are designed to be amplified and sequenced individually on Sanger instruments but might also be pooled for sequencing on next-generation sequencing platforms.

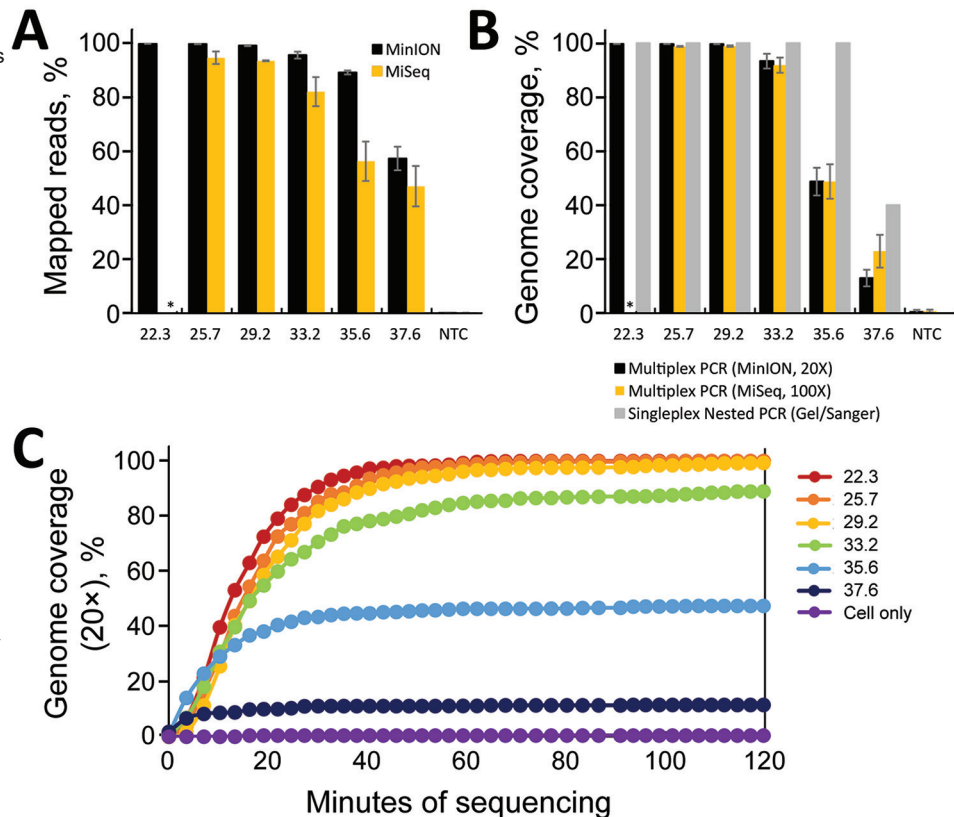
To determine the sensitivity of each sequencing strategy, we generated a set of 6 ten-fold serial dilutions of a SARS-CoV-2 isolate (J. Harcourt, unpub. data, <https://doi.org/10.1101/2020.03.02.972935>). Virus RNA was diluted into a constant background of A549 human cell line total nucleic acid (RNaseP cycle threshold [ $C_t$ ] 29). We quantitated each dilution by using the Centers for Disease Control and Prevention SARS-CoV-2 real-time reverse transcription PCR for the nucleocapsid 2 gene (8). The 6

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**Figure 1.** Limits of detection for sequencing severe acute respiratory syndrome coronavirus 2. Triplicate serial dilutions of virus isolate A12 (J. Harcourt, unpub. data, <https://doi.org/10.1101/2020.03.02.972935>) were amplified by using the singleplex or multiplex primer set. Multiplex amplicons were barcoded, library-prepped, and sequenced on an Oxford MinION apparatus (<https://nanoporetech.com>) or an Illumina MiSeq apparatus (<https://www.illumina.com>). A) Percentage of reads that map to the virus genome for each sample. B) Percentage of virus genome that is covered at >20× depth by the multiplex amplicons on the MinION (black) or >100× depth on the MiSeq (orange), or covered by the nested, singleplex amplicons (gray) (measured by presence or absence on a gel). C) Real-time analysis of MinION sequencing data. Each data point represents the average 20× genome coverage of three replicates. NTC, nontemplate controls (human cell nucleic acid carried through the PCR and library preparation). Asterisk (\*) indicates that samples were not analyzed at that dilution.



dilutions spanned  $C_t$  values from 22 to 37, corresponding to  $\approx 2 \times 10^0$  to  $1.8 \times 10^5$  copies. We amplified triplicate samples at each dilution by using the multiplex PCR pools. Next, we pooled, barcoded, and made libraries from amplicons of each sample by

using the ligation-based kit and PCR barcode expansion kit (Appendix). MinION sequencing was performed on an R9.4.1 or R10.3 flow cell (Oxford) until we obtained >1–2 million raw reads. From those reads, 50%–60% of them could be demultiplexed. In

**Table 1.** Genome consensus accuracy for sequencing severe acute respiratory syndrome coronavirus 2\*

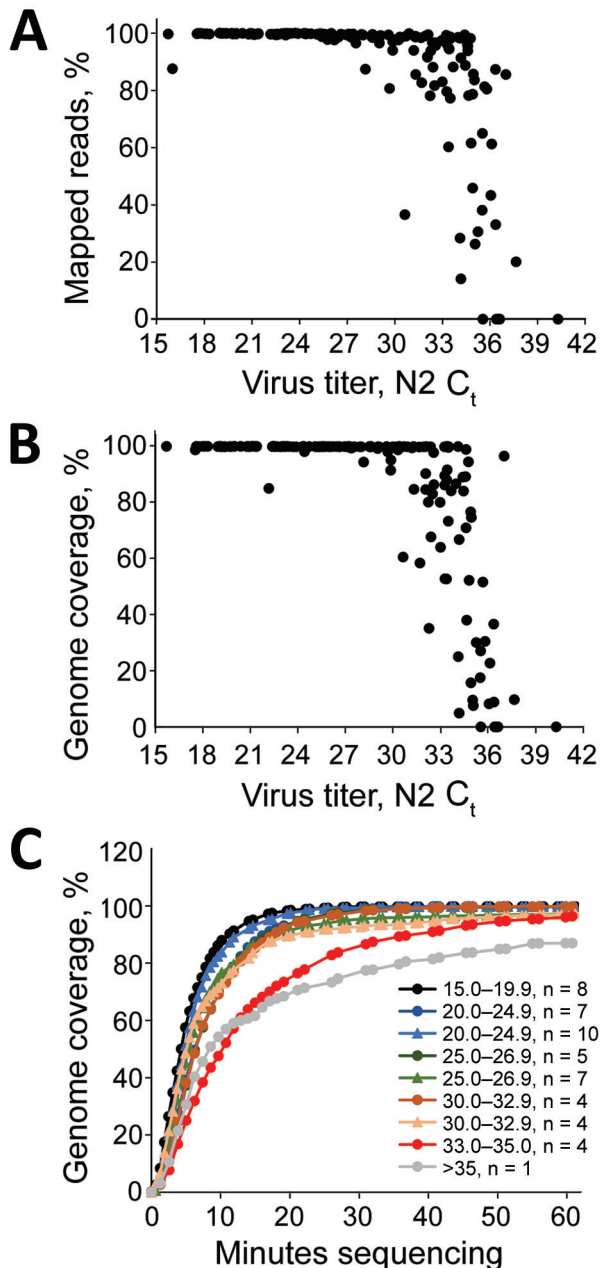
Virus titer (cycle threshold)	% Coverage, 20×†	Indels	Indel bases	Single-nucleotide polymorphisms	% Identity†
22.3	99.659	0	0	0	100
	99.722	0	0	0	100
	99.635	0	0	0	100
25.7	99.635	0	0	0	100
	99.615	0	0	0	100
	99.642	0	0	0	100
29.2	99.508	0	0	0	100
	99.635	0	0	0	100
	99.615	0	0	0	100
33.2	93.024	1	1	0	100
	93.603	2	35	0	100
	87.894	0	0	0	100
35.6	41.653	1	1	0	100
	51.266	0	0	1	99.993
	50.821	1	15	2	99.987
37.6	14.634	0	0	1	99.977
	9.317	0	0	0	100
	12.363	0	0	0	100

\*Because the 5' and 3' ends are primer sequences, 100% coverage is not possible.

†Percentage of covered bases identical to reference sequence, excludes indels and low-coverage bases.

addition, we sequenced these amplicons by using the Illumina MiSeq for comparison (Appendix).

For MinION sequencing, the reads were basecalled and analyzed by using an in-house read mapping



**Figure 2.** Sequencing of severe acute respiratory syndrome coronavirus 2 clinical samples. A, B) Percentage mapped (A) and percentage genome coverage (B) for 167 clinical severe acute respiratory syndrome coronavirus 2 samples amplified by using a multiplex PCR strategy and sequenced on the MinION apparatus (<https://nanoporetech.com>). C) Time-lapse of 20x genome coverage obtained for clinical specimens at the indicated cycle threshold values. Data points indicate average coverage over time for various samples and grouped by run and the indicated  $C_t$  values.  $C_t$ , cycle threshold; N2, nucleoprotein 2.

pipeline (Appendix). For samples with  $C_t \leq 29$ , we obtained >99% SARS-CoV-2 reads and >99% genome coverage at 20x depth, decreasing to an average of 93% genome coverage at  $C_t$  33.2 and 48% at  $C_t$  35 (Figure 1, panels A, B). Furthermore, we were able to obtain full genomes at >20x reading depth within the first 40–60 min of sequencing (Figure 1, panel C).

Consensus accuracy, including single-nucleotide polymorphisms and indels, is critical for determining coronavirus lineage and transmission networks. For high-consensus-level accuracy, we filtered reads based on length, mapped them to the reference sequence (GenBank accession no. RefSeq NC\_045512), trimmed primers based on position, and called variants with Medaka (<https://github.com>) (Appendix). Each Medaka variant was filtered by coverage depth (>20x) and by the Medaka model-derived variant quality (>30). We used the variant quality score as a heuristic to filter remaining noise from the Medaka variants compared with Sanger-derived sequences. After these steps, the data approaches 100% consensus accuracy (Table 1). Identical results were found by using the R9.4.1 pore through samples with  $C_t$  values through 33.2. The larger deletions in some of the samples with  $C_t$  values >33.2 (Table 1) do not appear to be sequencing errors because they are also detected as minor populations within higher-titer samples.

In the MiSeq data, we observed a similar trend in percent genome coverage at 100x depth, and a slightly lower percentage mapped reads compared with Nanopore data (Figure 1, panels A, B). Increased read depth using the MiSeq potentially enables increased sample throughput. However, the number of available unique dual indices limits actual throughput.

For the nested, singleplex PCR panel, we amplified the same serial dilutions with each nested primer set (Appendix). The endpoint dilution for full-genome coverage is a  $C_t \approx 35$  (Figure 1, panel B). At the  $C_t$  37 dilution, we observed major amplicon dropout; at this dilution, there are <10 copies of the genome on average/reaction.

These protocols enabled rapid sequencing of initial clinical cases of infection with SARS-CoV-2 in the United States. For these cases, we amplified the virus genome by using the singleplex PCR and sequenced the amplicons by using the MinION and Sanger instruments to validate MinION consensus accuracy. The MinION produced full-length genomes in <20 min of sequencing, and Sanger data was available the following day.

We used the multiplex PCR strategy for subsequent SARS-CoV-2 clinical cases ( $n = 167$ ) with  $C_t$  values ranging from 15.7 to 40 (mean 28.8, median

**Table 2.** Comparison of input, time, and cost requirements for sequencing 1 or 96 specimens of severe acute respiratory syndrome coronavirus 2

Method	Input, $\mu\text{L}^*$	1 sample		96 samples	
		Turnaround time	Approximate cost/sample $\dagger$	Turnaround time	Approximate cost/sample $\dagger$
Multiplex/MinION	10	6–8 h	\$528.70	8–10 h	\$35.88
Multiplex/MiSeq	10	30–68 h $\ddagger$	\$1,443.29	30–68 h $\ddagger$	\$57.87
Singleplex/Sanger	190	16–18 h	\$354.40	17–19 d	\$354.40

\*Assumes a process with 200  $\mu\text{L}$  of resuspended respiratory specimen (from a total of 2 mL), extracted, and eluted into 100  $\mu\text{L}$ . See Appendix (<https://wwwnc.cdc.gov/EID/article/26/10/20-1800-App1.pdf>) for details.  
 $\dagger$ Includes specific enzyme and reagent costs; excludes common laboratory supplies and labor costs.  
 $\ddagger$ Varies according to the sequencing kit used.

29.1). In cases with a  $C_t < 30$ , we observed an average of 99.02% specific reads and 99.2% genome coverage at  $>20\times$  depth (Figure 2, panels A, B). Between  $C_t$  30 and 33, genome coverage varied by sample, and decreased dramatically at higher  $C_t$  values, analogous to the isolate validation data. For these samples, we multiplexed 20–40 barcoded samples/flowcell. Enough data are obtained with 60 min of MinION sequencing for most samples, although for higher titer samples, 10–20 min of sequencing is sufficient (Figure 2, panel C).

Up-to-date primer sequences, protocols, and analysis scripts are available on GitHub ([https://github.com/CDCgov/SARS-CoV-2\\_Sequencing/tree/master/protocols/CDC-Comprehensive](https://github.com/CDCgov/SARS-CoV-2_Sequencing/tree/master/protocols/CDC-Comprehensive)). Data from this study is deposited in the National Center for Biotechnology Information Sequence Read Archive (BioProjects PRJNA622817 and PRJNA610248).

## Conclusions

Full-genome sequencing is a critical tool in understanding emerging viruses. Initial sequencing of SARS-CoV-2 showed limited genetic variation (9,10). However, some signature variants have been useful for describing the introduction and transmission dynamics of the virus (11; T. Bedford et al., unpub. data, <https://doi.org/10.1101/2020.04.02.20051417>; X. Deng et al., unpub. data, <https://doi.org/10.1101/2020.03.27.20044925>; M. Worobey et al., unpub. data, <https://doi.org/10.1101/2020.05.21.109322>).

We provide 2 validated PCR target-enrichment strategies that can be used with MinION, MiSeq, and Sanger platforms for sequencing SARS-CoV-2 clinical specimens. These strategies ensure that most laboratories have access to  $\geq 1$  strategies.

The multiplex PCR strategy is effective at generating full genome sequences up to  $C_t$  33. The singleplex, nested PCR is effective up to  $C_t$  35, varying based on sample quality. The turnaround time for the multiplex PCR MinION protocol is  $\approx 8$  hours from nucleic acid to consensus sequence and that for Sanger sequencing is  $\approx 14$ –18 hours (Table 2). The multiplex

PCR protocols offer an efficient, cost-effective, scalable system, and add little time and complexity as sample numbers increase (Table 2). Results from this study suggest multiplex PCR might be used effectively for routine sequencing, complemented by singleplex, nested PCR for low-titer virus samples and confirmation sequencing.

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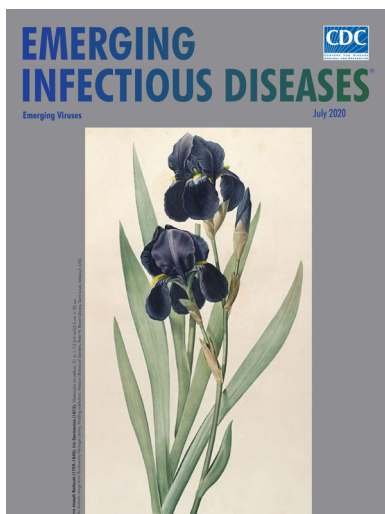
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# Rapid, Sensitive, Full-Genome Sequencing of Severe Acute Respiratory Syndrome Coronavirus 2

## **Appendix**

The following pages describe 2 validated protocols for generating high-quality, full-length severe acute respiratory syndrome coronavirus 2 genomes from primary samples. One protocol uses multiplex reverse transcription PCR, followed by MinION or MiSeq sequencing; the other uses singleplex, nested reverse transcription PCR and Sanger sequencing.

# Protocols for SARS-CoV-2 sequencing

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**Pathogen Discovery Team  
NCIRD/DVD/RVB  
Centers for Diseases Control and Prevention**

# Table of Contents

Disclaimers .....	4
Application Notes .....	5
Singleplex nested RT-PCR .....	6
Protocol Notes .....	6
Required Reagents .....	6
Procedure .....	6
1. First round of RT-PCR .....	6
2. Second round of semi-nested or nested PCR .....	7
Sanger sequencing .....	8
Required Reagents .....	8
Procedure .....	8
Multiplex PCR .....	9
Protocol Notes .....	9
Required reagents .....	9
Procedure .....	9
1. Generate primer pools .....	9
2. First-strand synthesis .....	9
3. Multiplex PCR .....	10
Nanopore Sequencing .....	11
Protocol Notes .....	11
Procedure .....	11
1. Barcode amplicons .....	11
2. Prepare Nanopore Ligation-based Library .....	12
3. Load MinION and sequence .....	13
4. Generate consensus sequences from MinION data .....	14
5. Quality control and analysis suggestions .....	15
Illumina Library Preparation and Sequencing .....	16
Protocol Notes .....	16
Required Reagents .....	16
Procedure for Library Preparation .....	16
1. Fragmentation and End Repair .....	16
2. Adapter Ligation .....	17
3. PCR enrichment of Adapter-Ligated DNA .....	17



4. Sizing and quantitation.....	18
MiSeq sequencing .....	20
Protocol Notes.....	20
Required Reagents .....	20
Procedure .....	20
1. Dilute and Pool Libraries .....	20
2. Denature Libraries .....	20
3. Load and Run MiSeq.....	21
4. Generation of consensus sequences from MiSeq data.....	22
Appendix A – Singleplex PCR Primers.....	23
Appendix B – Sequencing Primers.....	25
Appendix C – Plate Setup for Nested PCR and Sanger Sequencing.....	27
Appendix D – Multiplex PCR Primers .....	29
Appendix E – AMPure XP bead clean-up.....	32
Appendix F – Quantitation using Qubit .....	33
Required reagents .....	33
Procedure .....	33
Appendix G – CENTRI-SEP 96 Protocol .....	34

## Disclaimers

The findings and conclusions in this report have not been formally disseminated by the Centers for Disease Control and Prevention and should not be construed to represent any agency determination or policy.

The protocols described here are for research purposes only and should not be used in place of approved diagnostic testing.

# Application Notes

## Validation specimen submission extraction, and quantitation

1. For clinical specimens, CDC requested that submitting labs submit upper respiratory swabs in 2-3 mL viral transport medium (VTM), according to the guidelines detailed at: <https://www.cdc.gov/coronavirus/2019-ncov/lab/guidelines-clinical-specimens.html>.
2. Extraction of respiratory specimens in was performed using the QIAamp Viral RNA Mini Kit (QIAGEN). 200 uL of specimen VTM was used for each extraction and eluted from the column in 100 uL RNase-free water.
3. Samples for validation were quantitated with CDC N2 qRT-PCR assay, detailed at <https://www.cdc.gov/coronavirus/2019-ncov/lab/rt-pcr-panel-primer-probes.html>

## General Guidelines

1. Multiplex PCR Protocol is effective for Ct < 33.
2. Singleplex PCR is effective for some samples up to Ct 35. These are also useful for fill in reactions.
3. For the full Singleplex/Sanger protocol, two nucleic acid extractions (400 uL raw sample) will be needed.

# Singleplex nested RT-PCR

## Protocol Notes

To complete this protocol, 190  $\mu$ L of extracted template is needed. For samples between Ct 27 and 35, two rounds of nested RT-PCR are recommended; for samples up to Ct 27, one round of RT-PCR is recommended. The resulting PCR products can be individually proceeded with Sanger sequencing, or they can be pooled for Oxford Nanopore or Illumina sequencing, depending on the number of samples and availability of sequencing platforms.

**See Appendix C for recommended plate setups.**

## Required Reagents

Company	Product	Catalog number
Thermo (Invitrogen)	Superscript III one-step RT-PCR with Platinum Taq High Fidelity DNA polymerase	12574035
Sigma Aldrich (Roche)	Protector RNase inhibitor	3335402001
Takara	LA Taq DNA polymerase with GC buffer	RR02AG
	Nuclease Free water	
	50 $\mu$ M Primers	

## Procedure

### 1. First round of RT-PCR

- 1.1. Prepare the first-round master mix as below. Please note, the protocol is generic as all 38 primer pairs require the same master mix (see Appendix A). For each SARS-CoV-2 sample to be sequenced, 38 individual PCR reactions are required.

Component	Volume ( $\mu$ L)
Water	1.75
2x Buffer (2.4mM MgSO <sub>4</sub> )	12.5
5mM MgSO <sub>4</sub>	4.5
50 $\mu$ M Primer For	0.25
50 $\mu$ M Primer Rev	0.25
RNase Inhib. 40U/ $\mu$ L	0.25
SSIII / Platinum Taq high fidelity	0.5
Pre-mix	20
Template (RNA)	5
Total	25

- 1.2. Add 5 $\mu$ L of RNA template to each of the 38 PCR reactions. Spin tubes/plates down and proceed to PCR.
- 1.3. Perform first round PCR with the cycling parameters as below.

60°C	1 minutes	Decrease 0.5 C°/sec
50°C	30 minutes	
94°C	15 seconds	40 cycles
55°C	15 seconds	
72°C	1 minutes	
72°C	7 minutes	
4°C	Hold	

## 2. Second round of semi-nested or nested PCR

- 2.1. After first round RT-PCR is complete, prepare the master mix for 2<sup>nd</sup> round of semi-nested or nested PCR as below. Please note, the protocol is generic as all 38 second round primer pairs require the same master mix. Primer information is located in Appendix A. For the 2<sup>nd</sup> round of semi-nested- or nested PCR, there are 38 individual PCR reactions for each sample to be sequenced.

Component	Volume (uL)
Water	5.75
2× GCBuffer I	12.5
dNTP Mixture (2.5 mM each)	4
50uM Primer For	0.25
50uM Primer Rev	0.25
TaKaRa LA Taq™ (5 units/μl)	0.25
Pre-mix	23
Template (1R product)	2
Total	25

- 2.2. Add 2 uL of the corresponding first round PCR product to the second round PCR master mix. Spin tubes/plates down and proceed to PCR.
- 2.3. Perform second round PCR with the cycling parameters as below.

94°C	3 minutes	
94°C	15 seconds	30 cycles
55°C	15 seconds	
72°C	1 minutes	
72°C	7 minutes	
4°C	Hold	

- 2.4. Following the completion of second round PCR, run 3 uL of all 38 PCR reactions on 1% agarose gels or fragment analyzer to check for amplification.

# Sanger sequencing

## Required Reagents

Company	Product	Catalog number
Thermo (Applied Biosystems)	ExoSap-It	78201.1.ML
Thermo (Applied Biosystems)	BigDye v3.1 cycle sequencing kit	4337455
Princeton Separations	Centri-sep 96 well plates	CS-963
	Nuclease Free water	
	5 uM Primers	

## Procedure

1. Transfer 10 uL of each PCR reaction to new tubes/plate for ExoSap cleanup. Add 4 uL ExoSap-It to each PCR reaction (10 uL) and incubate at 37°C for 15 minutes, followed by 80°C for 15 minutes on a thermocycler.
2. Prepare sequencing master mix as below.  
Sequencing primers for each amplicon are listed in Appendix B

Component	Volume (uL)
Water	5.5
2x Buffer	2
5uM Primer	1
BigDye 3.1 enzyme	1
Pre-mix	9.5
Template (2R PCR product)	0.5
Total	10

3. Add 0.5 uL of corresponding ExoSap cleaned PCR product to each sequencing reaction mix. Spin tubes/plates down and proceed to sequencing PCR.
4. Perform sequencing PCR with the parameters listed below:

96°C	2 minutes	
96°C	30 seconds	30 cycles
50°C	15 seconds	
60°C	3 minutes	
4°C	Hold	

5. Following sequencing PCR, clean-up of sequencing reactions is performed with Centri-Sep 96-well plates following the manufacturer's instructions (Appendix G) with one addition. 20 uL nuclease free water is added to the 96-well collection plate prior to the final spin.
6. The 96-well collection plate with the cleaned sequencing sample plus water is loaded onto the ABI sequencer.
7. Sequencer 5.4 is used for data analysis of Sanger PCR data.

# Multiplex PCR

## Protocol Notes

This protocol uses 10 uL of template for each sample. The pooled, multiplexed PCR products can be followed with nanopore sequencing or Illumina MiSeq sequencing depending on the number of samples and available sequencing platforms. We have been able to sequence full genomes reliably under Ct 30, and depending on the sample, up to Ct 33.

This protocol was adapted from Quick J et al. *Nat Protoc.* 2017 Jun;12(6):1261-1276.

## Required reagents

Company	Product	Catalog number
Thermo Fisher (Invitrogen)	SuperScript IV 1 <sup>st</sup> strand synthesis system	18091200
NEB	NEBNext Q5 Hot Start HiFi PCR Master Mix	M0543L
	Nuclease Free water	
	Primers	

## Procedure

### 1. Generate primer pools

- 1.1. Prepare primers as 50 uM primer stocks.
- 1.2. Add an equal volume of each 50 uM primer stock to six 1.5mL Eppendorf tubes labeled as pool 1, 2, 3, 4, 5, and 6. Primers for each pool are listed in Appendix D.
- 1.3. Prepare 10 uM working concentration by diluting each pool 1:5 with nuclease free water.

### 2. First-strand synthesis

- 2.1. Mix the following components.

Component	Volume(uL)
RNA (template)	10
Random primer 25uM	2
dNTPs	1
Total	13

- 2.2. Denature the template-primer-dNTP mix at 65°C for 5 minutes.
- 2.3. Place on ice for 5 minutes.
- 2.4. Add the following components to the template-primer-dNTP mix:

Component	Volume (uL)
5x SSIV buffer	4
0.1 M DTT	1
RNAse inhibitor	1
SSIV RT (200 units/uL)	1
Total	20

- 2.5. Incubate in a thermal cycler at the following temperatures:  
25°C 10 minutes, 50°C for 10 minutes, 85°C for 10 minutes, hold at 4°C.
- 2.6. Spin down. Can be stored at -20°C
- 2.7. Add 1 uL RNase H and incubate at 37°C for 20 minutes

### 3. Multiplex PCR

- 3.1. Mix the following components in 6 wells of a PCR plate or strip tube.

Component	Volume (uL)
NEBNext Q5 Hot Start HiFi PCR Master Mix	15
PCR grade water	10.2
Primer pool 1, 2, 3, 4, 5, or 6 (10uM)	1.8
Total	27

- 3.2. Add 3 uL of cDNA from above to each tube.
- 3.3. Run the following PCR program:

98°C	30 seconds	
98°C	15 seconds	40 cycles
65°C	5 minutes	
4°C	Hold	

Note: fewer cycles may be used, but 40 cycles is used to maximize detection of lower-titer samples.

- 3.4. Optional: Run a 2% agarose gel for each multiplexed PCR reaction pool 1, 2, 3, 4, 5, and 6 to check for specific bands of the correct size (0.4-0.6 kb).
- 3.5. Pool 20 uL from each of 6 tubes of multiplexed PCR reactions in a 0.3 mL tube in a PCR strip or a well in PCR plate (the total volume is 120 uL).
- 3.6. Add 1X ratio (120 uL) of AMPure XP beads to the PCR product pools.
- 3.7. Purify according to standard AMPure protocol (see Appendix E).
- 3.8. Elute in 80 uL water.
- 3.9. Quantitate 1 uL of cleaned PCR products using Qubit dsDNA HS kit (Appendix F).
- 3.10. Optional: Run a 2% agarose gel and load 3 uL of cleaned PCR products to check for specific bands of the correct size (0.4-0.6 kb).



# Nanopore Sequencing

## Protocol Notes

This protocol takes advantage of the multiplexing density afforded by the “PCR Barcoding Expansion 1-96” kit. This protocol is derived from Oxford Nanopore’s protocols available at <http://community.nanoporetech.com>.

Required reagents for Nanopore barcoding and sequencing:

Company	Product	Catalog number
NEB	NEBNext Ultra II End-repair/dA tailing module	E7546
NEB	Blunt/TA ligase master mix	M0367
NEB	NEBNext Quick Ligation Module	E6056
TaKaRa	TaKaRa LA Taq DNA Polymerase with GC Buffer	RR02AG
Beckman Coulter	Agencourt Ampure XP Beads	A63880/A63881
Oxford Nanopore Technologies	Nanopore Ligation Sequencing Kit (1D)	SQK-LSK109
Oxford Nanopore Technologies	PCR Barcoding Expansion 1-96	EXP-PBC096
Oxford Nanopore Technologies	SpotON Flow Cell (R9.4.1)	FLO-MIN106D
Oxford Nanopore Technologies	MinION	MinION Mk1B

## Procedure

### 1. Barcode amplicons

1.1. Mix the following components:

Component	Volume (uL)
500 ng amplicon DNA	25
Ultra II end-prep reaction buffer	3.5
Ultra II end-prep enzyme mix	1.5
Total	30

1.2. Incubate at 20°C for 10 minutes, 65°C for 5 minutes, hold at 4°C.

1.3. Add 1X ratio (30 uL) AMPure XP beads.

1.4. Purify according to standard AMPure protocol (Appendix E).

1.5. Elute the DNA target from the beads with 17 uL water.

1.6. Optional: quantitate 1 uL of cleaned end-prep DNA using Qubit dsDNA HS kit (Appendix F)

1.7. Mix the following components:

Component	Volume (uL)
Cleaned end-prep DNA	15
Barcode Adapter	10
Blunt/TA ligase master mix	25
Total	50

1.8. Incubate at 20°C for 10 minutes.

1.9. Add 1X ratio (50 uL) AMPure XP beads.

1.10. Purify according to standard AMPure protocol (Appendix E).

1.11. Elute the DNA in 12 uL water.

- 1.12. Transfer eluate into new PCR plate or well
- 1.13. Quantitate 1  $\mu$ L of ligated DNA according to the protocol (Appendix F).
- 1.14. Mix the following components:

Component	Volume( $\mu$ L)
30ng adapter-ligated DNA	x
PCR Barcode primer (one of BC1-BC96)	1
2x GC Buffer I	25
dNTP mix (10mM)	8
TaKaRa LA Taq (5U/ $\mu$ L)	0.5
Water	50 – x
Total	50

- 1.15. Mix by pipetting and spin down
- 1.16. Run the following PCR program:

95°C	3 minutes	
95°C	15 seconds	18 cycles
62°C	15 seconds	
72°C	1 minutes	
72°C	7 minutes	
4°C	Hold	

- 1.17. Add 1X ratio (50  $\mu$ L) of AMPure XP beads.
- 1.18. Purify according to standard AMPure protocol (Appendix E).
- 1.19. Elute the DNA target from the beads with 25  $\mu$ L water.
- 1.20. Quantitate 1  $\mu$ L cleaned, barcoded PCR products with Qubit dsDNA HS kit (Appendix F).

## 2. Prepare Nanopore Ligation-based Library

- 2.21. Pool the barcoded PCR products equally by mass.
- 2.22. Prepare LSK109 ligation-based libraries by mixing the following components:

Component	Volume( $\mu$ L)
1 $\mu$ g pooled barcoded sample	x
DNA CS	1
Ultra II End-prep reaction buffer	7
Ultra II End-prep enzyme mix	3
Water	49-x
Total	60

- 2.23. Incubate at 20°C for 10 minutes, 65°C for 5 minutes, hold at 4°C.
- 2.24. Add 1X ratio (60  $\mu$ L) of AMPure XP beads.
- 2.25. Purify according to standard AMPure protocol (Appendix E).
- 2.26. Elute the DNA target from the beads with 62  $\mu$ L water.
- 2.27. To ligate sequencing adapters, mixing the following components:

Component	Volume(uL)
End-repaired DNA from previous step	60
Ligation buffer (LNB)	25
NEBNext Quick T4 DNA Ligase	10
Adapter Mix (AMX)	5
Total	100

- 2.28. Incubate 10 minutes at 20°C
- 2.29. Add 0.8X ratio (80 uL) of AMPure XP beads
- 2.30. Purify according to standard AMPure protocol (Appendix E).
- 2.31. Elute the DNA target from the beads with 15 uL water
- 2.32. Quantitate 1 uL clean, prepared library with Qubit dsDNA HS kit (Appendix F).

### 3. Load MinION and sequence

- 3.33. Set up the MinION flow cell and host computer, including MinKNOW software.
- 3.34. Open the MinKNOW GUI from the desktop icon and establish a local connection.
- 3.35. Inset flow cell into MinION.
- 3.36. Click “Check Flow Cells” at the bottom of the screen then click “Start test.” Check the number of active pores available. When the check is complete, it is reported in the Notification panel. Check to ensure it has enough pores for a good sequencing run (warranty for flow cells: 800 nanopores or above checked within 5 days of receipt).
- 3.37. Thaw the Sequencing Buffer (SQB), Loading Beads (LB), Flush Tether (FLT) and one tube of Flush Buffer (FB) at room temperature before placing the tubes on ice.
- 3.38. Thoroughly mix the Sequencing Buffer (SQB) and Flush Buffer (FB) tubes by vortexing,
- 3.39. Spin down the Flush Tether (FLT) tube, mix by pipetting, and return to ice.
- 3.40. Open the lid of the nanopore sequencing device and slide the flow cell's priming port cover clockwise 90 degrees. (The following steps are demonstrated at <https://youtu.be/CC11Jlydgrc>)
- 3.41. Set a P1000 pipette to 200 uL, insert the tip into the priming port, turn the wheel until the dial shows 220-230 uL, or until you can see a small volume of buffer entering the pipette tip. Do not remove more than this.
- 3.42. Visually check that there is continuous buffer from the priming port across the sensor array.
- 3.43. Prepare the flow cell priming mix: add 30 uL of thawed and mixed Flush Tether (FLT) directly to the tube of thawed and mixed Flush Buffer (FB), and mix by pipetting up and down.
- 3.44. Load 800 uL of the priming mix into the flow cell via the priming port, avoiding the introduction of air bubbles.
- 3.45. Wait for 5 minutes.
- 3.46. Thoroughly mix the contents of the Loading Beads (LB) by pipetting.
- 3.47. Prepare library for loading my mixing:

Component	Volume (uL)
Sequencing Buffer (SQB)	37.5
Loading Beads (LB), mixed immediately before use	25.5
150-200 ng DNA Library	12
Total	75

- 3.48. Gently lift the SpotON sample port cover to make the SpotON sample port accessible.
- 3.49. Load 200 µl of the priming mix into the flow cell via the priming port (not the SpotON sample port), avoiding the introduction of air bubbles.
- 3.50. Mix the prepared library gently by pipetting up and down just prior to loading.

- 3.51. Add 75 uL of sample to the flow cell via the SpotON sample port in a dropwise fashion. Ensure each drop flows into the port before adding the next.
- 3.52. Gently replace the SpotON sample port cover, making sure the bung enters the SpotON port, close the priming port and replace the MinION lid.
- 3.53. Start the sequencing run using the MinKNOW software.

## 4. Generate consensus sequences from MinION data

There are many considerations for generating high-quality consensus data from the MinION. Here are some suggestions for basecalling based on our experience.

Software:

Software	Source URL
Guppy 3.4.1+	<a href="https://community.nanoporetech.com/downloads">https://community.nanoporetech.com/downloads</a>
Medaka 0.11.5	<a href="https://github.com/nanoporetech/medaka">https://github.com/nanoporetech/medaka</a>
Minimap2 2.17 (r941)	<a href="https://github.com/lh3/minimap2">https://github.com/lh3/minimap2</a>
SAMtools 1.9	<a href="http://www.htslib.org/">http://www.htslib.org/</a>
BCFtools 1.9	<a href="http://www.htslib.org/">http://www.htslib.org/</a>
BAMClipper	<a href="https://github.com/tommyau/bamclipper">https://github.com/tommyau/bamclipper</a>
cutadapt 2.3+	<a href="https://github.com/marcelm/cutadapt">https://github.com/marcelm/cutadapt</a>
vcf_mask_lowcoverage.pl	<a href="https://github.com/CDCgov/SARS-CoV-2_Sequencing">https://github.com/CDCgov/SARS-CoV-2_Sequencing</a>
IGV	<a href="http://software.broadinstitute.org/software/igv/">http://software.broadinstitute.org/software/igv/</a>

Example commands below have user-supplied variable names bold. You will need to customize the details to your environment.

### 4.1. Basecalling

Basecalling may also be done using MinKNOW software. If so, you may skip the Guppy basecalling step.

```
# Run Guppy
guppy_basecaller --input_path $rundir --save_path $outputdir -r \
  --config na_r9.4.1_450bps_hac.cfg --barcode_kits EXP-PBC096 \
  --trim_barcodes --require_barcodes_both_ends
# Combine all the output fastq files
mkdir $outputdir/fastq
find $outputdir -name "*.fastq" |while read infile; do
  if [[ $i =~ barcode|unclassified ]]; then
    outfile=$(grep -Eo "barcode..|unclassified" <<< $infile).fastq
    outfile="fastq/$outfile"
    cat $infile >> $outfile
  fi
done
```

### 4.2. Filter on quality and length

Filtering out low quality sequence, as well as unexpectedly long and short reads helps tremendously on off-target mapping affecting consensus quality.

```
cutadapt -j $threads -m 300 -M 1200 -q 15 -o $fastqfiltered $fastqfile
```

### 4.3. Mapping

Download reference sequence from GenBank: MN908947.3

```
minimap2 -L -a -x map-ont -t 12 MN908947.fasta $ fastqfiltered > $samfile  
samtools view -b $samfile | samtools sort - -o $bamfile  
samtools index $bamfile
```

### 4.4. Clip primers

This step requires a BEDPE file describing the positions of the primers. It is available at

[https://github.com/CDCgov/SARS-CoV-2\\_Sequencing](https://github.com/CDCgov/SARS-CoV-2_Sequencing)

BAMClipper by default will output at file with the suffix “primerclipped.bam.”

Clipping by position allows only primers near the beginning of a read to be trimmed (rather than genuine sequence in the middle of a read), and it is faster than sequence-based trimming (e.g. Porechop).

```
cd $outputdir  
bamclipper.sh -b $bamfile -p SC2_200324.bedpe -n 12 -u 80 -d 80
```

### 4.5. Generate VCF and consensus sequences

Medaka is very lenient with calling variants. We generally require a variant quality score of  $\geq 30$  and depth of coverage  $\geq 20$  to call a variant. Below 20X coverage, we call an ‘N.’

The script to automate the filtering is available at [https://github.com/CDCgov/SARS-CoV-2\\_Sequencing](https://github.com/CDCgov/SARS-CoV-2_Sequencing)

```
# Generate Medaka VCF File  
medaka consensus --model r941_min_high_g344 --threads 12 \ $primerclippedbamfile  
$primerclippedbamfile.hdf  
medaka variant MN908947.fasta $primerclippedbamfile.hdf $vcf  
# Filter variants and generate consensus sequence  
vcf_mask_lowcoverage.pl --bam $primerclippedbamfile \  
--reference MN908947.fasta --vcf $vcf --consout $consensusfasta \  
--depth 20 --qual 30
```

## 5. Quality control and analysis suggestions

- 5.6. Watch out for 1-base insertions/deletions. Though consensus calling has improved considerably, there are residual errors. There are several stretches in SARS-CoV-2 that have homopolymers long enough to be problematic
- 5.7. Do not ignore other deletions. There have been several deletions reported (3, 9, 15, 33bp, 384bp, etc), so keep in mind the difference between a potential real indel and missing amplicon or nanopore error.
- 5.8. IGV can be useful for examining the “believability” of variants However, some of these 1-2bp indels appear in the reads, but they cannot be confirmed by Illumina or Sanger sequencing. These are either unlucky PCR bias or systematic sequencing error.

# Illumina Library Preparation and Sequencing

## Protocol Notes

Starting Material: 100 pg–250 ng DNA. We recommend that the DNA be in 1X TE (10 mM Tris pH 8.0, 1 mM EDTA), however, 10 mM Tris pH 7.5–8, low EDTA TE or water are also acceptable. If the input DNA is less than 26 µl, add TE (provided) to a final volume of 26 µl. This protocol is adapted from the NEBNext Ultra II FS protocol, which can be found in its entirety at <http://www.neb.com>.

For sizing, other devices, such as the 2100 BioAnalyzer, 5200 FragmentAnalyzer, QIAxcel, or LabChipGX may also be used. These vary in quantitation accuracy, so fluorometric quantitation with Qubit (or similar instrument) or qPCR is recommended.

## Required Reagents

Company	Product	Catalog number
New England Biolabs (NEB)	NEBNext Ultra II FS DNA Library Prep Kit for Illumina	E7805S/E7805L
New England Biolabs (NEB)	NEBNext® Multiplex Oligos for Illumina (96 Unique Dual Index Primer Pairs)	E6440S/E6440L
Beckman Coulter	Agencourt Ampure XP Beads	A63880/A63881
	10mM Tris-HCl, pH 8.0	
	Molecular biology grade ethanol	
	Nuclease-free water	
Agilent	High Sensitivity D1000 screen tape	5067-5584
Agilent	High Sensitivity D1000 reagents	5067-5585

## Procedure for Library Preparation

### 1. Fragmentation and End Repair

- 1.1. Ensure that the Ultra II FS Reaction Buffer is completely thawed. If a precipitate is seen in the buffer, pipette up and down several times to break it up, and quickly vortex to mix. Place on ice until use.
- 1.2. Vortex the Ultra II FS Enzyme Mix 5-8 seconds prior to use and place on ice.
- 1.3. Add the following components to a 0.2 ml thin wall PCR tube on ice

Component	Volume (µL)
NEBNext Ultra II FS Enzyme Mix (yellow tube)	2
NEBNext Ultra II FS Reaction Buffer (yellow tube)	7
DNA (pooled PCR amplicons)	26
Total	35

- 1.4. Vortex the reaction for 5 seconds and briefly spin down. Place in a thermocycler with the heated lid set to  $\geq 75^{\circ}\text{C}$  and run the following program:

37°C for 7 minutes, 65°C for 30 minutes, 4°C hold indefinitely

## 2. Adapter Ligation

- 2.1. Determine dilution for adapter if necessary, see table below. Dilute the NEBNext Adapter for Illumina (red tube) in 10 mM Tris-HCl, pH 8.0 with 10 mM NaCl as indicated below.

Input DNA in the End Prep reaction	Adapter dilution (volume of adapter: total volume)	Working adapter concentration
250 ng - 101 ng	No dilution	15 uM
100 ng – 5 ng	10-fold (1:10)	1.5 uM
Less than 5 ng	25-fold (1:25)	0.6 uM

- 2.2. Add the following components directly to the FS reaction mixture from 1.1(35 uL):

Component	Volume (uL)
NEBNext Ultra II Ligation Master Mix (red tube)	30
NEBNext Ultra II Ligation enhancer (green tube)	1
NEBNext adapter for Illumina	2.5

Notes:

- Mix the Ultra II Ligation Master Mix by pipetting up and down several times prior to adding to the reaction.
  - The Ligation master mix and ligation enhancer can be mixed ahead of time and is stable for at least 8 hours at 4°C. Do not pre-mix the adapter prior to use in the adapter ligation step.
  - The NEBNext adapter is provided in NEBNext Multiplex Oligos for Illumina (96 Unique Dual Index Primer Pairs)
- 2.3. Set a pipette to 50 uL and pipette entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.  
Note: The NEBNext Ultra II Ligation master mix is very viscous. Care should be taken to ensure adequate mixing of the ligation reaction as incomplete mixing will result in reduced ligation efficiency. The presence of a small amount of bubbles will not interfere with performance.
- 2.4. Incubate at 20°C for 15 minutes in a thermocycler with the heated lid off.
- 2.5. Add 3 uL of USER enzyme (red tube) to the ligation mixture.  
Note: This step is only required for use with NEBNext adapters. USER enzyme is provided in NEBNext Multiplex Oligos for Illumina (96 Unique Dual Index Primer Pairs)
- 2.6. Mix well and incubate at 37°C for 15 minutes in a thermocycler with the heated lid set to ≥47°C
- 2.7. Add 57uL (0.8X) re-suspended AMPure XP beads to the ligation reaction (87uL).
- 2.8. Follow steps in the AMPure XP bead clean-up section (Appendix E).
- 2.9. Elute the DNA target from the beads by adding 17 uL of 10mM Tris-HCl or 0.1X TE.
- 2.10. Transfer 15 uL to a new PCR tube for amplification.

## 3. PCR enrichment of Adapter-Ligated DNA

- 3.1. Add the following components to a sterile strip tube:

Component	Volume (uL)
Adapter ligated DNA fragments (from above)	15
Unique dual index primer pair*	10
NEBNext Ultra II Q5 master mix (blue tube)	25
Total volume	50

\*The primers are provided in NEBNext® Multiplex Oligos for Illumina® (96 Unique Dual Index Primer Pairs). Please refer to the NEB #E6440 manual for valid barcode combination and tips for setting up PCR reactions

- 3.2. Set a pipette to 40 uL and then pipette the entire volume up and down at least 10 times to mix thoroughly. Perform a quick spin to collect all liquid from the sides of the tube.
- 3.3. Place tube on a thermocycler and perform PCR amplification using the following PCR cycling conditions:

Cycle step	Temperature	Time	# of cycles
Initial denaturation	98°C	30 seconds	1
Denaturation	98°C	10 seconds	3-15*
Annealing/extension	65°C	75 seconds	
Final extension	65°C	5 minutes	1
Hold	4°C	∞	

\*Follow the recommendations for cycle number listed in the table below.

#### Cycle recommendations

Input DNA in the end prep reaction	# of cycles required to generate a library yield of:	
	100 ng	1 ug
250 ng	2-3	3-4
100 ng	3-4	4-5
50 ng	4-5	5-6
10 ng	6-7	8-9
5 ng	7-8	9-10
1 ng	8-10	11-12
0.5 ng	9-10	12-13
0.1ng	12-13	N/A

- 3.4. Add 0.9X AMPure XP beads to the PCR reactions (45uL).
- 3.5. Follow steps in the AMPure XP bead clean-up section (Appendix E).
- 3.6. Elute DNA target from beads into 33 uL 0.1X TE.
- 3.7. Transfer 30 uL supernatant to a new PCR tube. Libraries can be store at -20C°.
- 3.8. Check size distribution of libraries and quantitate library concentration.

#### 4. Sizing and quantitation

- 4.1. Allow TapeStation reagents to equilibrate at room temperature for 30 minutes prior to use.
- 4.2. Vortex reagents well before use.
- 4.3. To prepare ladder, mix 2 uL high sensitivity D1000 sample buffer with 2 uL high sensitivity D1000 ladder.



- 4.4. To prepare sample, mix 2 uL high sensitivity D1000 sample buffer with 2 uL sample.
- 4.5. Spin down, then vortex using IKA vortexer and adapter at 2000 rpm for 1 minute.
- 4.6. Spin down to position the sample at the bottom of the tube.
- 4.7. Load samples into the 2200 TapeStation instrument and follow the software procedure for analysis.
- 4.8. Quantitate 1 uL library sample with Qubit dsDNA HS kit (Appendix F).

# MiSeq sequencing

## Protocol Notes

This procedure requires Illumina-style libraries that have been quality-controlled and quantitated using the recommended procedures (i.e. TapeStation and Qubit or qPCR). Exact loading concentrations may vary by machine or lab-dependent factors. For more details on loading and running the MiSeq, consult the more detailed manuals at <http://www.illumina.com>.

## Required Reagents

Company	Product	Catalog number
Illumina	MiSeq reagent kit v3	MS-102-3003
Illumina	PhiX control kit v3	FC-110-3001
	NaOH	
	Nuclease-free water	

## Procedure

### 1. Dilute and Pool Libraries

- 1.1. Calculate the molar concentration of each library to be diluted using average size from the TapeStation and mass from Qubit, using the following equation:

$$\frac{\text{concentration (ng/uL)}}{660\text{g/mol} \times \text{avg library fragment size}} \times 10^6 \text{ uL/L} = \text{concentration (nM)}$$

- 1.2.
- 1.3. Make a 4nM dilution of each library.
- 1.4. Combine equal volumes of each diluted library into a new tube. This is the 4nM library pool.

### 2. Denature Libraries

- 2.1. Make a fresh dilution of 0.2N NaOH by combining the following volumes in a microcentrifuge tube:
  - 800 uL laboratory-grade water
  - 200 stock 1.0N NaOH
- 2.2. Remove HT1 from freezer and thaw at room temperature. Store at 2°C to 8°C until you are ready to dilute denatured libraries.
- 2.3. Combine the following volumes in a microcentrifuge tube:
  - 5 uL 4nM library
  - 5 uL 0.2N NaOH
- 2.4. Vortex briefly and then centrifuge at 280 x g for 1 minute.
- 2.5. Incubate at room temperature for 5 minutes.
- 2.6. Add 990 uL pre-chilled HT1 to the tube containing denatured library. The result is 1 mL of a 20pM denatured library.
- 2.7. Dilute the 20pM library to the desired concentration, see table below:

Concentration	6 pM	8 pM	10 pM	12 pM	15 pM	20 pM
20 pM library	180 uL	240 uL	300 uL	360 uL	450 uL	600 uL
Pre-chilled HT1	420 uL	360 uL	300 uL	240 uL	150 uL	0 uL

- 2.8. Invert to mix and then pulse centrifuge.

- 2.9. Dilute stock PhiX to 4nM by combining:
  - 2 uL 10 nM PhiX library
  - 3 uL 10 mM Tris-Cl, pH 8.5 with 0.1% Tween 20
- 2.10. Denature the PhiX control by adding the following volumes in a microcentrifuge tube:
  - 5 uL 4nM PhiX library
  - 5 uL 0.2N NaOH
  - Remaining 4nM PhiX can be frozen and reused
- 2.11. Vortex briefly to mix and centrifuge at 280 x g for 1 minute.
- 2.12. Incubate at room temperature for 5 minutes.
- 2.13. Dilute denatured PhiX library to 20 pM by adding 990 uL pre-chilled HT1 to the PhiX tube. Invert to mix.
- 2.14. If using a MiSeq reagent kit v2, dilute 20 pM PhiX library to 12.5 pM by adding the following volumes in a microcentrifuge tube:
  - 375 uL 20pM denatured PhiX library
  - 225 uL pre-chilled HT1
- 2.15. Combine library and PhiX control according to the table below:

Denatured and diluted PhiX	30 uL
Denatured and diluted library	570 uL

- 2.16. Set aside on ice until you are ready to load it onto the reagent cartridge.

### 3. Load and Run MiSeq

- 3.1. Thaw frozen reagents overnight at 4°C overnight or in a RT water bath.
- 3.2. Mix reagents thoroughly by inverting several times. Inspect the bottom of reagent cartridge to ensure all liquids return to the bottom of each tube without any air bubbles.
- 3.3. Using a 1000 uL pipette tip, piece the foil on position 17.
- 3.4. Using a fresh 1000 uL pipette tip, transfer the denatured and library (with PhiX spiked) into position 17.
- 3.5. Generate Sample Sheet using MiSeq Experiment Manager.
- 3.6. Load MiSeq according to onscreen instructions in the MiSeq Control software.

## 4. Generation of consensus sequences from MiSeq data

The Illumina MiSeq provides very high-quality data, and consensus sequenced may be generated by variety of methods, including commercial tools such as Geneious and CLC Genomics Workbench. The procedure outlined here is a suggestion using free, open source tools.

Software	Source URL
cutadapt 2.3+	<a href="https://github.com/marcelm/cutadapt">https://github.com/marcelm/cutadapt</a>
bowtie2	<a href="https://github.com/BenLangmead/bowtie2">https://github.com/BenLangmead/bowtie2</a>
seqtk	<a href="https://github.com/lh3/seqtk">https://github.com/lh3/seqtk</a>
SAMtools 1.9	<a href="http://www.htslib.org/">http://www.htslib.org/</a>
BCFtools 1.9	<a href="http://www.htslib.org/">http://www.htslib.org/</a>
IGV	<a href="http://software.broadinstitute.org/software/igv/">http://software.broadinstitute.org/software/igv/</a>

- 4.1. Trim reads for quality (Q25+) and for adapters on both ends. Then trim primer sequences (a hard 30 bases on each end), keeping only sequenced that are at least 75 bases. For reads <150 bases, this will need to be modified.

```
cutadapt -j $threads -g GTTTCACGTCACGATA -G GTTTCACGTCACGATA \
-a TATCGTGACTGGGAAAC -A TATCGTGACTGGGAAAC -g ACACCTCTTCCCTACACGACGCTCTTCCGATCT \
-G ACACCTCTTCCCTACACGACGCTCTTCCGATCT -a AGATCGGAAGAGCACACGTCTGAACTCCAGTCA \
-A AGATCGGAAGAGCGTCGTGTAGGGAAAGAGTGT -n 3 -m 75 -q 25 \
--interleaved $read1 $read2 | cutadapt -j $threads --interleaved -m 75 -u 30 \
-u -30 -U 30 -U -30 -o $read1.trim.fastq -p $read2.trim.fastq -
```

- 4.2. Map reads to reference sequence

```
bowtie2-build MN908947.fasta MN908947
bowtie2 --sensitive-local -p $threads -x MN908947 \
-l $read1.trim.fastq -2 $read2.trim.fastq -S $samfile
samtools view -b $samfile | samtools sort -o $bamfile
samtools index $bamfile
```

- 4.3. Call variants, generate consensus sequence. This will call positions covered by at least 100 reads.

```
samtools mpileup -aa -d 8000 -uf MN908947.fasta $bamfile |\
bcftools call -Mc |tee -a $vcf | \
vcfutils.pl vcf2fq -d 100 -D 100000000|\
seqtk seq -A - | sed '2~2s/[actg]/N/g' > $consensusfasta
```

# Appendix A – Singleplex PCR Primers

Amplicon	1st round	Sequence	Size	2nd round	Sequence	Size
PCR1	1F_209_1	GTTGCAGCCGATCATCAGCAC	756	W1_2L_368	TGGAGGAGGTCTTATCAGAGGC	597
	SC2M1-2_RIGHT_965	GTTCACGGCAGCAGTATACACC		SC2M1-2_RIGHT_965	GTTCACGGCAGCAGTATACACC	
PCR2	W1_2F_00826_1	AACAACCTCTGTGGCCCTGATG	904	W1_2F_00850_2	TACCCTCTTGAGTGCAATTAAG	853
	W1_2R_01730_1	TCCACAAAAGCACTTGTGGAAGC		W1_2R_01703_2	AAAGATGCCAAAATAATGGCG	
PCR3	W1_3F_01573_1	GGTGTGTGGAGAAGGTTCCG	938	W1_3F_01596_2	AGGTCTTAATGACAACCTCTTG	894
	W1_3R_02511_1	TGTGGGAAGTGTTCCTCCCTC		W1_3R_02490_2	TAAGAAGATAATTTCTTTTGGG	
PCR4	W1_4F_02387_1	CATTTGTCACGCACTCAAAGGG	942	W1_4F_02404_2	AAGGGATTGTACAGAAAGTGTG	903
	W1_4R_03329_1	GTCTGAACAACCTGGTGAAGTTCC		W1_4R_03307_2	CCATCTCTAATTGAGGTTGAAC	
PCR5	W1_5F_03185_1	AGCAAGAAGAAGATTGGTTAGATGATG	1014	W1_5F_03208_2	GATGATAGTCAACAACTGTTGG	868
	W1_5R_04199_1	ATTTCAAGTAGTCCACCAGCC		W1_5R_04175_2	TTAGTAGGTATAACCACAGCAG	
PCR6	W1_6F_04054_1	CATCCAGATTCTGCCACTCTTG	968	W1_6F_04073_2	TTGTTAGTGACATTGACATCAC	932
	W1_6R_05022_1	CATGTCCACAACCTTGCCTGTG		W1_6R_05005_2	TGTGGAGGTTAATGTTGTCTAC	
PCR7	W1_7F_04884_1	TCCTACCACATTCACCTAGATGG	933	W1_7F_04904_2	ATGGTGAAGTTATCACCTTTG	893
	W1_7R_05817_1	AGCACCGTCTATGCAATACAAAG		W1_7R_05797_2	AAGTTTCTTTAGAAGTTATATG	
PCR8	W1_8F_05676_1	TGTTATGATGTCAGCACCACTG	977	W1_8F_05699_2	CTCAGTATGAACCTTAAGCATGG	933
	W1_8R_06653_1	ACAGCAGCTAAACCATGAGTAGC		W1_8R_06632_2	GCAAGGGTTTTCAACCTAATAC	
PCR9	W1_9F_06522_1	TACAGAAGAGGTTGGCCACAC	954	W1_9F_06543_2	AGATCTAATGGCTGCTTATGTAG	919
	W1_9R_07476_1	ACAACCGTCTACAACATGCAC		W1_9R_07462_2	CATGCACATAACTTTCCATAC	
PCR10	W1_10F_07326_1	TGCAGTACATTTTATTAGTAATCTTGG	999	W1_10F_07356_2	TATGTGTTAATAATTAATCTTG	952
	W1_10R_08325_1	GTCACGGGGTGTGATGTTTTTC		W1_10R_08308_2	TTTCAACTTTGTTATAGGTGAGC	
PCR11	W1_11F_08170_1	GCAGCTCGGCAAGGGTTTGTGG	952	W1_11F_08184_2	GTTTGTGATTGAGATGTAGAAAC	922
	W1_11R_09122_1	CGTGTGTCAGGGCGTAAACTTTC		W1_11R_09106_2	AACTTTCATAAGCAACAGAACC	
PCR12	W1_12F_08996_1	CAGCTTGTGTTTTGGCTGCTG	980	W1_12F_09017_2	AATGTACAATTTTAAAGATGC	949
	W1_12R_09976_1	GAGCCTTTGCGAGATGACAAC		W1_12R_09966_2	GAGATGACAACAAGCAGCTTC	
PCR13	W1_13F_09831_1	GTATCTAAAGTTGCGTAGTGATG	1005	W1_13F_09850_2	GATGTGCTATTACCTCTTACGC	966
	W1_13R_10836_1	AACGGCAATTCAGTTTGAGC		W1_13R_10816_2	CAGAAAGGGTCTTAGTATGTC	
PCR14	W1_14F_10686_1	TGTTATAAATGGAGACAGGTGG	984	W1_14F_10708_2	TTTCTCAATCGATTACCACAAC	949
	W1_14R_11670_1	GCGGTTGAGTAAACAAAAGAGGC		W1_14R_11657_2	CAAAAAGAGCCAAAGTAAACAAG	
PCR15	W1_15F_11527_1	GCCAGAGGTATTGTTTTATGTGTGT	911	W1_15F_11527_2	GCCAGAGGTATTGTTTTATGTGTGT	889
	W1_15R_12438_1	GGGAACACAACCATCTCTTGC		W1_15R_12416_2	TTGTTGATAATGTTGTTGAGTGC	
PCR16	W1_16F_12311_1	CTAGATCTGAGGACAAGAGGGC	929	W1_16F_12327_2	GAGGGCAAAAGTTACTAGTGC	892
	W1_16R_13240_1	ACGATGCACCACCAAGGATTC		W1_16R_13219_2	CTTGATCCATATTGGCTTCCGG	
PCR17	W1_17F_13112_1	ATCTAGCTAGTGGGGACAACC	930	W1_17F_13126_2	GGACAACCAATCACTAATTGTG	902
	W1_17R_14042_1	AATACCAGCATTTCGCATGGCA		W1_17R_14028_2	GCATGGCATCACAAGTGTAC	
PCR18	W1_18F_13873_1	TACTTGTACATACAATTGTTGTGATG	914	W1_18F_13873_2	TACTTGTACATACAATTGTTGTGATG	914
	18R_14809_1	GATAGTAGTCATAATCGCTGATAGCAG		W1_18R_14787_1	TAGCAGCATTACCATCTGAGC	
PCR19	W1_19F_14655_1	GCTTTTCAAACCTGTCAAACCCGG	902	W1_19F_14670_2	AAACCCGGTAATTTAACAAG	879
	W1_19R_15557_1	TGCATTAACATTGGCCGTGAC		W1_19R_15549_2	CATTGGCCGTGACAGCTTGAC	
PCR20	W1_20F_15429_1	AGTGAAATGGTCATGTGTGGCG	971	W1_20F_15441_2	ATGTGTGGCGTTCACTATATG	939
	W1_20R_16400_1	ACAACCTGGAGCATTGCAAAAC		W1_20R_16380_2	CATACGGATTAAACAGACAAGAC	
PCR21	21F_16221_1	GCATACAGTCTTACAGGCTGTTGG	919	W1_21F_16291_2	GCATACGTAGACCATTCTTATG	849
	21R_17140_1	CAGAAGGGTAGTAGAGAGCTAGGC		21R_17140_1	CAGAAGGGTAGTAGAGAGCTAGGC	
PCR22	W1_22F_17065_1	ATTCTACACTCCAGGGACCACC	970	W1_22F_17082_2	CCACTGGTACTGGTAAGAGTC	930
	W1_22R_18035_1	TAAAGTTGCCACATTCCTACGTGG		W1_22R_18012_2	GAATTTCAAGACTTGTAATTTG	
PCR23	W1_23F_17881_1	CCACTGAAACAGCTCACTCTTG	1019	W1_23F_17901_2	TGTAATGTAACAGATTTAATG	978
	W1_23R_18900_1	TAACAAGCACTCGTGGACAGC		W1_23R_18879_2	CTAGACACCTAGTCATGATTGC	
PCR24	W1_24F_18767_1	TGTTCAACAATGGGTTTTACAGG	910	W1_24F_18786_2	ACAGGTAACCTACAAGCAACC	879
	W1_24R_19677_1	CCTGTTGTCCATCAAAGTGTCCC		W1_24R_19665_2	CAAAGTGTCCCTTATTACAAC	

PCR25	W1_25F_19546_1 25R_20572_1	CAGCTGGCTTTAGCTTGTGGG CAACCTTAGAACTACAGATAAACTCTTG	936	W1_25F_19546_1 W1_25R_20482_1	CAGCTGGCTTTAGCTTGTGGG GATGAACCTGTTTGCGCATCTG	936
PCR26	W1_26F_20343_1 W1_26R_21315_1	CATAGTCAGTTAGGTGGTTTAC CTATTTGTTCCGCTGGTTTGCC	972	W1_26F_20356_2 W1_26R_21300_2	GTGGTTTACATCTACTGATTGG GTTTGCCAAGATAATTACATCC	944
PCR27	27F_21136_1 27R_22218_1	AAGCTAGCTCTTGGAGGTTCCG CCCTGAGGGAGATCACGCAC	926	W1_27F_21204_2 W1_27R_22099_2	CTCATGGGACACTTCGCATGGTGG CAAGGTCCATAAGAAAAGGCTG	895
PCR28	W1_28F_21976_1 W1_28R_22993_1	CCATTTTTGGGTGTTTATTACC TGCTACCGCCTGATAGATTTTC	1017	W1_28F_21996_2 W1_28R_22975_2	CCACAAAACACAAAAGTTGG TTTCAGTTGAAATATCTCTCTC	979
PCR29	W1_29F_22847_1 W1_29R_23812_1	TTACAGGCTGCGTTATAGCTTGG TGCTGCATTAGTTGAATCACC	965	W1_29F_22864_2 W1_29R_23795_2	GCTTGGAAATTCTAACATCTTG TCACCACAAATGTACATTGTAC	931
PCR30	W1_30F_23681_1 W1_30R_24625_1	ACTCTAATAACTCTATTGCCATACCCAC CAGAAGCTCTGATTTCTGCAGC	944	W1_30F_23704_2 W1_30R_24610_2	CCCACAAATTTTACTATTAGTG CTGCAGCTCTAATTAATTGTTG	906
PCR31	W1_31F_24492_1 W1_31R_25491_1	AAATGATATCCTTTACGCTTGCACAAAG TTGCACTAGCGCAACAAAATC	999	W1_31F_24514_2 W1_31R_25476_2	GACAAAGTTGAGGCTGAAGTGC CAAAATCTGAAGGAGTAGCATC	962
PCR32	W1_32F_25348_1 W1_32R_26367_1	CCAGTGCTCAAAGGAGTCAAATTAC ACGCACACAATCGAAGCGCAG	1019	W1_32F_25357_2 W1_32R_26358_2	AAAGGAGTCAAATTACATTACAC ATCGAAGCGCAGTAAGGATGGC	1001
PCR33	W1_33F_26222_1 W1_33R_27128_1	ACAAGCTGATGAGTACGAACTTATG TGCCAATCCTGTAGCGACTGTATGC	906	W1_33F_26241_2 W1_33R_27115_2	CTTATGTACTCATTGTTTCGG GCGACTGTATGCAGAAAACC	874
PCR34	W1_34F_26988_1 W1_34R_28006_1	TAGGACGCTGTGACATCAAGG AGGACACGGGTCATCAACTAC	1018	W1_34F_26999_2 W1_34R_27992_2	GACATCAAGGACCTGCCTAAAG CAACTACATATGGTTGATGTTG	993
PCR35	35F_27834_1 35th_R2_28855	ATCTTTTGGTTCTCACTTGAAGTGC TGAAGTGTTCGACTACGTGATG	1021	W1_35F_27875_1 35th_R2_28855	TGTCACGCCTAAACGAACATG TGAAGTGTTCGACTACGTGATG	980
PCR36	W1_36F_28694_1 W1_36R_29724_1	CACCAAAGATCACATTGGCAC TGTGGTGGCTCTTTCAAGTCC	1030	W1_36F_28716_2 W1_36R_29724_2	CCGCAATCCTGCTAACAAATGC TGTGGTGGCTCTTTCAAGTCC	1008
PCR37	W1_37F_29551_1 W1_37R_29873_2	AGGCAGATGGGCTATATAAACG TTTTGTCAATCTCCTAAGAAGC	322	W1_37F_29596_2 W1_37R_29873_2	TATAGTCTACTCTTGTGCAGAATG TTTTGTCAATCTCCTAAGAAGC	280
PCR38	SC2M1-1_LEFT2_1 SC2M1-1_RIGHT2_495	TTAAAGTTTATACCTTCCCAGG CGAGCATCCGAACGTTTGATGA	495	0_1b W1_1R_490	TTAAAGTTTATACCTTCCCAGGTA CATCCGAACGTTTGATGAACAC	490

## Appendix B – Sequencing Primers

### Sequencing primer to amplicon matrix

PCR Product	Sequencing primers			
PCR1	W1_2L_368	SC2M1-2_RIGHT_965	SC2M1-2_LEFT_445	SC2M1-1_RIGHT_574
PCR2	W1_2F_00850_2	W1_2R_01703_2	W1_4F_1067*	W1_3R_1206*
PCR3	W1_3F_01596_2	W1_3R_02490_2	W1_6L_1819	W1_5R_1969
PCR4	W1_4F_02404_2	W1_4R_03307_2	W1_9L_2948*	W1_8R_3094*
PCR5	W1_5F_03208_2	W1_5R_04175_2	W1_11L_3638*	W1_10R_3792*
PCR6	W1_6F_04073_2	W1_6R_05005_2	W1_13L_4307*	W1_12R_4522*
PCR7	W1_7F_04904_2	W1_7R_05797_2	W1_15L_5159*	W1_14R_5299*
PCR8	W1_8F_05699_2	W1_8R_06632_2	SC2M1-16_LEFT_6030	SC2M1-15_RIGHT_6172
PCR9	W1_9F_06543_2	W1_9R_07462_2	W1_20L_6877*	W1_19R_7009*
PCR10	W1_10F_07356_2	W1_10R_08308_2	W1_22L_7625*	W1_21R_7771*
PCR11	W1_11F_08184_2	W1_11R_09106_2	W1_25L_8669*	W1_24R_8794*
PCR12	W1_12F_09017_2	W1_12R_09966_2	W1_27L_9308*	W1_26R_9459
PCR13	W1_13F_09850_2	W1_13R_10816_2	W1_29R_10593*	W1_30L_10448*
PCR14	W1_14F_10708_2	W1_14R_11657_2	W1_32L_11111*	W1_31R_11251*
PCR15	W1_15F_11527_2	W1_15R_12416_2	W1_34L_11808*	W1_33R_11948*
PCR16	W1_16F_12327_2	W1_16R_13219_2	W1_37L_12878*	W1_35R_12700*
PCR17	W1_17F_13126_2	W1_17R_14028_2	W1_39L_13600*	W1_38R_13741*
PCR18	W1_18F_13873_1	W1_18R_14787_1	W1_41L_14342*	W1_40R_14503
PCR19	W1_19F_14670_2	W1_19R_15549_2	W1_43L_14960*	W1_42R_15108*
PCR20	W1_20F_15441_2	W1_20R_16380_2	W1_46L_16004*	W1_44R_15773*
PCR21	W1_21F_16291_2	21R_17140_1	W1_48L_16735*	W1_46R_16490*
PCR22	W1_22F_17082_2	W1_22R_18012_2	W1_50L_17424*	W1_49R_17553*
PCR23	W1_23F_17901_2	W1_23R_18879_2	W1_53L_18503*	W1_52R_18667*
PCR24	W1_24F_18786_2	W1_24R_19665_2	W1_55L_19277*	W1_54R_19405*
PCR25	W1_25F_19546_1	W1_25R_20482_1	W1_57L_20013*	W1_56R_20146*
PCR26	W1_26F_20356_2	W1_26R_21300_2	W1_59L_20656*	W1_58R_20796*
PCR27	W1_27F_21204_2	W1_27R_22099_2	W1_61L_21411*	W1_60R_21562
PCR28	W1_28F_21996_2	W1_28R_22975_2	W1_64L_22457	W1_63R_22612*
PCR29	W1_29F_22864_2	W1_29R_23795_2	W1_66L_23182*	W1_65R_23308*
PCR30	W1_30F_23704_2	W1_30R_24610_2	W1_69L_24259*	W1_67R_24002*
PCR31	W1_31F_24514_2	W1_31R_25476_2	W1_71L_24935*	W1_70R_25075*
PCR32	W1_32F_25357_2	W1_32R_26358_2	SC2M1-66_LEFT_25665	SC2M1-65_RIGHT_25790
PCR33	W1_33F_26241_2	W1_33R_27115_2	SC2M1-68_LEFT_26454	SC2M1-67_RIGHT_26590
PCR34	W1_34F_26999_2	W1_34R_27992_2	SC2M1-71_LEFT_27650	SC2M1-69_RIGHT_27432
PCR35	W1_35F_27875_1	35th R2_28855	W1_81L_28414*	SC2M1-71_RIGHT_28203
PCR36	W1_36F_28716_2	W1_36R_29724_2	SC2M1-75_LEFT_29344	SC2M1-74_RIGHT_29469
PCR37	W1_37F_29596_2	W1_37R_29873_2		
PCR38	0_1b	W1_1R_490		

\*Primer sequence in table below.

SC2M1 primers are sourced from the multiplex primer set (Appendix D). Others are primers from Appendix A.

## Additional Sequencing Primer Sequences

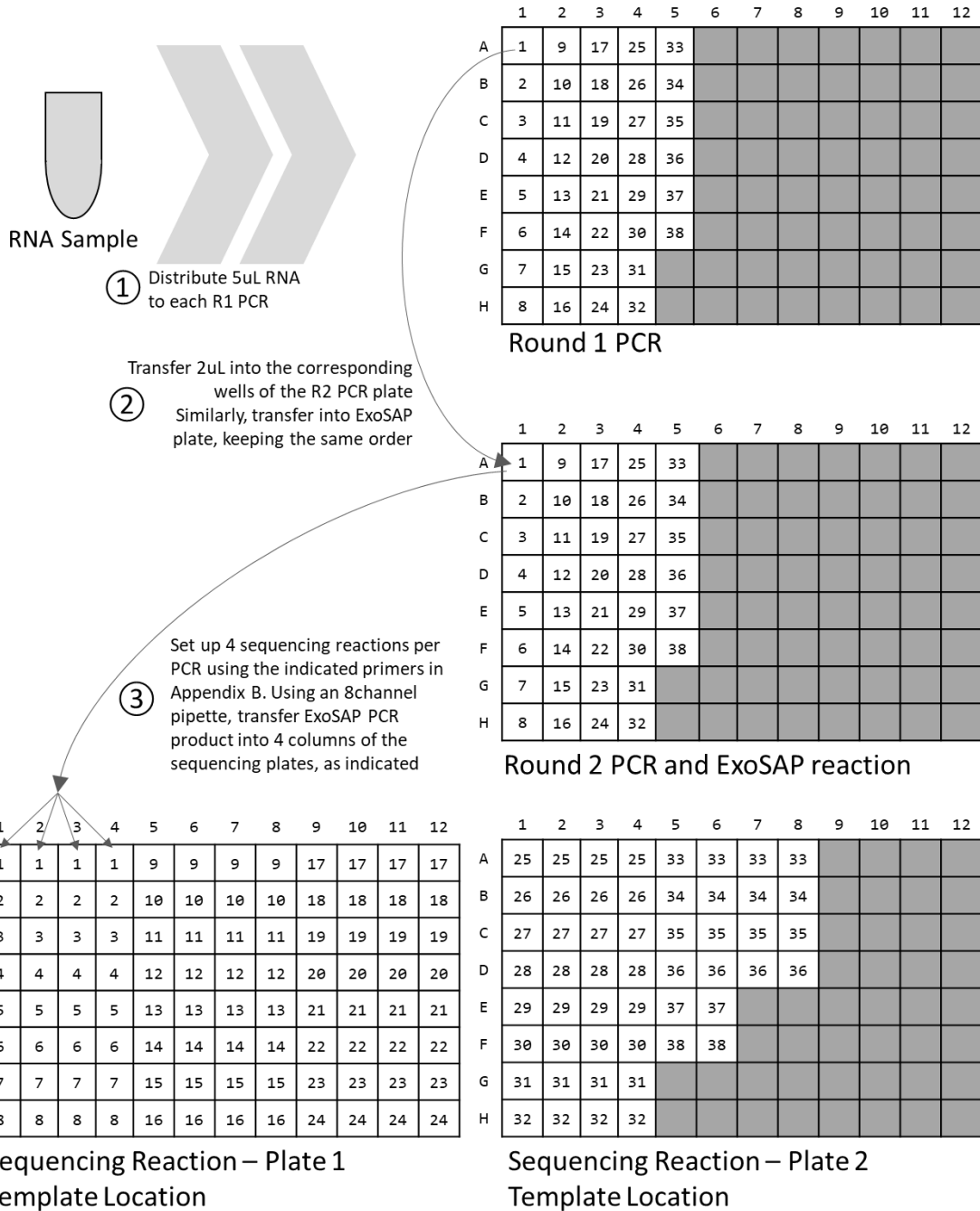
Name	Sequence	Nmae	Sequence
W1_4F_1067	GGGAATGTCCAAATTTTGATTTCC	W1_41L_14342	TTTGATGACAGATGCATTCTGC
W1_3R_1206	TGGTTGCATTCAATTTGGTGACG	W1_43L_14960	TAAATGGGGTAAGGCTAGACTTTATTATG
W1_9L_2948	TTGATTTAGATGAGTGGAGTATGGCTAC	W1_42R_15108	CGGTGCGAGCTCTATTCTTTGC
W1_8R_3094	ATGGCTCAAACCTCTTCTTCTCAC	W1_46L_16004	GATTGAACGGTTCGTGCTTTAGC
W1_11L_3638	GTGAAGACATTCAACTCTTAAGAGTGC	W1_44R_15773	GCTAGCCACTAGACCTTGAGATGC
W1_10R_3792	AGCTAAGTAGACATTTGTGCGAAC	W1_48L_16735	GGGAAGTTGGTAAACCTAGACCAC
W1_13L_4307	GAAGTGTCTTGGAAATTTGCGAG	W1_46R_16490	AGCACACAATGGAAAATAATGGG
W1_12R_4522	CACCATAATCAACCACACCCCTC	W1_50L_17424	GTGTACATTGGCGACCCCTGCTC
W1_15L_5159	TTGAGTACTACCACACAATGATCC	W1_49R_17553	TTCCGAGGAACATGTCTGGACC
W1_14R_5299	CAGTGGCAAGATAACAGTTGTTATC	W1_53L_18503	AGGACTTCTTTGGAATGTAGTGGC
W1_20L_6877	AGTGTCCGTAATTTTGTCTAGAGGC	W1_52R_18667	CATAGACAACAGGTGCGCTCAG
W1_19R_7009	CAGCGGTTGAGTAGATTAAGAACC	W1_55L_19277	TTGTGATGGTGGCAGTTTGTATG
W1_22L_7625	ATTGTGATACATTTGTGCTGGTAGTAC	W1_54R_19405	CCATGAGACTCACATGGACTGTC
W1_21R_7771	GATGGATGGAACCATTTCACTG	W1_57L_20013	GACTTATTTAGAAATGCCCGTAATGGTG
W1_25L_8669	TTCAAGTGAATCATAGGATACAAGG	W1_56R_20146	AACTGTGTTTTTACGGCTTCTCC
W1_24R_8794	TACCACCACGCTGGCTAAACC	W1_59L_20656	AATCTAGTCAAGCGTGCAACC
W1_27L_9308	CAGGAGTTTTCTGTGGTGTAGATGC	W1_58R_20796	ATTTTGCACATTCATCATTATGCC
W1_29R_10593	GTTACCTTCTAAGTCTGTGCCAGC	W1_61L_21411	CTTAAATTAAGGGGACTGCTGTTATG
W1_30L_10448	CCAATTTCACTATTAAGGGTTCATTCC	W1_63R_22612	AAACAGATGCAAATCTGGTGGC
W1_32L_11111	TGGGTATTATTGCTATGTCTGCTTTTG	W1_66L_23182	TTCAACTTCAATGGTTTAAACAGGCAC
W1_31R_11251	TACGCATCACCAACTAGCAGG	W1_65R_23308	CAAGTGTCTGTGGATCACGGAC
W1_34L_11808	TGGCAAACCTTGATCAAAGTAGC	W1_69L_24259	ATGCAAATGGCTTATAGTTTAAATGG
W1_33R_11948	TGTAACGGACACATTGAGCCC	W1_67R_24002	TTGCTTGGTTTTGATGGATCTGG
W1_37L_12878	TCTATACAGAACTGGAACACCTTG	W1_71L_24935	ACTGTGATGTTGTAATAGGAATTGTCAAC
W1_35R_12700	AGACATCTGCTAGTGAACAGG	W1_70R_25075	TGCCAGAGATGTCACCTAAATCAAC
W1_39L_13600	GTCGCTTCCAAGAAAAGGACG	W1_81L_28414	AATACTGCGTCTTGGTTCACCG
W1_38R_13741	AAGTCATGTTTAGCAACAGCTGG		



# Appendix C – Plate Setup for Nested PCR and Sanger Sequencing

Primers are added to each PCR reaction (PCR1-PCR38) prior to adding RNA. The layout stays the same until sequencing reactions are run.

We recommend making PCR primer plates (R1 and R2) in the same format so that primers may be added by multichannel pipetting.



We recommend making sequencing primer plates as shown below, so that primers may be rapidly added to the sequencing reactions. Primer sequences may be found in Appendices A and D.

W1_2L_368	SC2M1-2- RIGHT_965	SC2M1- 2_LEFT_44 5	SC2M1-1- RIGHT_574	W1_9F_ 06543_2	W1_9R_ 07462_2	W1_20L_ 6877	W1_19R_ 7009	W1_17F_ 13126_2	W1_17R_ 14028_2	W1_39L_ 13600	W1_38R_ 13741
W1_2F_ 00850_2	W1_2R_ 01703_2	W1_4F_10 67	W1_3R_ 1206	W1_10F_ 07356_2	W1_10R_ 08308_2	W1_22L_ 7625	W1_21R_ 7771	W1_18F_ 13873_1	W1_18R_ 14787_1	W1_41L_ 14342	W1_40R_ 14503
W1_3F_ 01596_2	W1_3R_ 02490_2	W1_6L_ 1819	W1_5R_ 1969	W1_11F_ 08184_2	W1_11R_ 09106_2	W1_25L_ 8669	W1_24R_ 8794	W1_19F_ 14670_2	W1_19R_ 15549_2	W1_43L_ 14960	W1_42R_ 15108
W1_4F_ 02404_2	W1_4R_ 03307_2	W1_9L_ 2948	W1_8R_ 3094	W1_12F_ 09017_2	W1_12R_ 09966_2	W1_27L_ 9308	W1_26R_ 9459	W1_20F_ 15441_2	W1_20R_ 16380_2	W1_46L_ 16004	W1_44R_ 15773
W1_5F_ 03208_2	W1_5R_ 04175_2	W1_11L_ 3638	W1_10R_ 3792	W1_13F_ 09850_2	W1_13R_ 10816_2	W1_29R_ 10593	W1_30L_ 10448	W1_21F_ 16291_2	21R_ 17140_1	W1_48L_ 16735	W1_46R_ 16490
W1_6F_ 04073_2	W1_6R_ 05005_2	W1_13L_ 4307	W1_12R_ 4522	W1_14F_ 10708_2	W1_14R_ 11657_2	W1_32L_ 11111	W1_31R_ 11251	W1_22F_ 17082_2	W1_22R_ 18012_2	W1_50L_ 17424	W1_49R_ 17553
W1_7F_ 04904_2	W1_7R_ 05797_2	W1_15L_ 5159	W1_14R_ 5299	W1_15F_ 11527_2	W1_15R_ 12416_2	W1_34L_ 11808	W1_33R_ 11948	W1_23F_ 17901_2	W1_23R_ 18879_2	W1_53L_ 18503	W1_52R_ 18667
W1_8F_ 05699_2	W1_8R_06 632_2	SC2M1-16- LEFT_6030	SC2M1-15- RIGHT_617 2	W1_16F_ 12327_2	W1_16R_ 13219_2	W1_37L_ 12878	W1_35R_ 12700	W1_24F_ 18786_2	W1_24R_ 19665_2	W1_55L_ 19277	W1_54R_ 19405

### Sequencing Reaction – Plate 1 Sequencing Primer Location

W1_25F_ 19546_1	W1_25R_ 20482_1	W1_57L_ 20013	W1_56R_ 20146	W1_33F_ 26241_2	W1_33R_ 27115_2	SC2M1- 68_LEFT_ 26454	SC2M1- 67_RIGHT_ 26590				
W1_26F_ 20356_2	W1_26R_ 21300_2	W1_59L_ 20656	W1_58R_ 20796	W1_34F_ 26999_2	W1_34R_ 27992_2	SC2M1- 71_LEFT_ 27650	SC2M1- 69_RIGHT_ 27432				
W1_27F_ 21204_2	W1_27R_ 22099_2	W1_61L_ 21411	W1_60R_ 21562	W1_35F_ 27875_1	35th R2_ 28855	W1_81L_ 28414	SC2M1- 71_RIGHT_ 28203				
W1_28F_ 21996_2	W1_28R_ 22975_2	W1_64L_ 22457	W1_63R_ 22612	W1_36F_ 28716_2	W1_36R_ 29724_2	SC2M1- 75_LEFT_ 29344	SC2M1- 74_RIGHT_ 29469				
W1_29F_ 22864_2	W1_29R_ 23795_2	W1_66L_ 23182	W1_65R_ 23308	W1_37F_ 29596_2	W1_37R_ 29873_2						
W1_30F_ 23704_2	W1_30R_ 24610_2	W1_69L_ 24259	W1_67R_ 24002	0_1b	W1_1R_ 490						
W1_31F_ 24514_2	W1_31R_ 25476_2	W1_71L_ 24935	W1_70R_ 25075								
W1_32F_ 25357_2	W1_32R_ 26358_2	SC2M1- 66_LEFT_ 25665	SC2M1- 65_RIGHT_ 25790								

### Sequencing Reaction – Plate 2 Sequencing Primer Location

## Appendix D – Multiplex PCR Primers

### Pool 1

PCR	Name	Sequence
1	SC2M1-1_LEFT_31	ACCAACCAACTTTCGATCTCTTGT
	SC2M1-1_RIGHT_574	TGTCTCACCACACTACGACCGTAC
5	SC2M1-5_LEFT_1706	TCTGCTCCACAAGTGCTTTTGT
	SC2M1-5_RIGHT_2266	ACAGGTGACAATTTGTCACCG
9	SC2M1-9_LEFT_3306	TGGAACCTACACCAGTTGTTTCAGAC
	SC2M1-9_RIGHT_3878	CAGCGATCTTTTGTCAACTTGCT
13	SC2M1-13_LEFT_4885	TCCTACCACATTCCACCTAGATGG
	SC2M1-13_RIGHT_5400	GCACAAAAGTTAGCAGCTTCACC
17	SC2M1-17_LEFT_6408	CTGAAGAAGTAGTGGAAAATCCTACCA
	SC2M1-17_RIGHT_6903	GCCTCTAGACAAAATTTACCACACT
21	SC2M1-21_LEFT_8004	TTGGTGATAGTGCAGGAGTTGC
	SC2M1-21_RIGHT_8553	CCACCCCTAAGTGCTATCTTTGTTGT
25	SC2M1-25_LEFT_9551	CCAGTTTACTATTCTTACCTGGTGT
	SC2M1-25_RIGHT_10061	AACCACCTGCAAAAACAGCTGA
29	SC2M1-29_LEFT_11047	AGTCCAGAGTACTCAATGGCTTTTGT
	SC2M1-29_RIGHT_11541	ACAATACCTCTGGCCAAAACATGA
33	SC2M1-33_LEFT_12557	ATCCAACAGGTTGTAGATGCAGAT
	SC2M1-33_RIGHT_13136	TTGGTTGTCCCCACTAGCTAG
37	SC2M1-37_LEFT_14103	TTTCATACAAAACAGCCAGGT
	SC2M1-37_RIGHT_14641	GTGCAGCTACTGAAAAGCACGT
41	SC2M1-41_LEFT_15637	AGAAATAGAGATGTTGACACAGACTTTGT
	SC2M1-41_RIGHT_16208	GCCTCATAAACTCAGGTTCCCA
45	SC2M1-45_LEFT_17317	AATGCATTGCCTGAGACGACAG
	SC2M1-45_RIGHT_17903	CAAGAGTGAGCTGTTTCACTGGT
49	SC2M1-49_LEFT_18897	TGTTAAGCGTGTGACTGGACT
	SC2M1-49_RIGHT_19484	GCACCACCTAAATTGCAACGTG
53	SC2M1-53_LEFT_20554	TCTGTAGTTTCTAAGGTTGTCAAAGTGA
	SC2M1-53_RIGHT_21144	AGTAGCTTTTGTGTATAAACCCACA
57	SC2M1-57_LEFT_22203	GTGATCTCCCTCAGGTTTTTTCG
	SC2M1-57_RIGHT_22697	ACTTAAAAGTGAAAATGATGCGGAA
61	SC2M1-61_LEFT_23737	AATTCTACCAGTGTCTATGACCAAGAC
	SC2M1-61_RIGHT_24231	GCACCAAAGGTCCAACCAAG
65	SC2M1-65_LEFT_25214	CTAGGTTTTATAGCTGGCTTGATTGC
	SC2M1-65_RIGHT_25790	CATTTCCAGCAAAGCCAAAGCC
69	SC2M1-69_LEFT_26877	CTTCTCAACGTGCCACTCCATG
	SC2M1-69_RIGHT_27432	AGCGAGTGTATCAGTGCCAAG
73	SC2M1-73_LEFT_28525	TGGCTACTACCGAAGAGCTACC
	SC2M1-73_RIGHT_29045	GCTTCTTAGAAGCCTCAGCAGC

### Pool 2

PCR	Name	Sequence
2	SC2M1-2_LEFT_445	TTTGCCCAACTTGAACAGCCC
	SC2M1-2_RIGHT_965	GTTACAGGCAGCAGTATACACC
6	SC2M1-6_LEFT_2138	AAACCCGTCTTGTATTGGCTTG
	SC2M1-6_RIGHT_2642	TTTCGAGCAACATAAGCCCGTT
10	SC2M1-10_LEFT_3715	AGCTGGTATTTTTGGTGCTGACC
	SC2M1-10_RIGHT_4262	CCTGACCCGGGTAAGTGGTTAT
14	SC2M1-14_LEFT_5258	ACTTCTATTAATGGGCAGATAACAAGT
	SC2M1-14_RIGHT_5818	AGCACCGTCTATGCAATACAAAGT
18	SC2M1-18_LEFT_6748	AAACCGTGTTTGACTAATTATATGCCTT
	SC2M1-18_RIGHT_7255	TGCCAAAACCACTCTGCAACT
22	SC2M1-22_LEFT_8407	CGTTAAAGATTTTATGTCATTGTCTGAACA
	SC2M1-22_RIGHT_8913	TGCAAAAAGTCACCATTAGTTGTGC
26	SC2M1-26_LEFT_9903	AGTACAAGTATTTTAGTGGAGCAATGGA
	SC2M1-26_RIGHT_10451	TGGGCCCTCATAGCACATTGGTA
30	SC2M1-30_LEFT_11400	TGAATGCTTGACACTCGTTTATAAAGTT
	SC2M1-30_RIGHT_11944	CTGGACACATTGAGCCACAAT
34	SC2M1-34_LEFT_13006	TGCCACAGTACGTCTACAAGCT
	SC2M1-34_RIGHT_13501	GTGTAAGACGGGCTGCACCTTAC
38	SC2M1-38_LEFT_14480	ACTTCAGAGAGCTAGGTTGTGACA
	SC2M1-38_RIGHT_15027	TGCGAAAAGTGCACTTTGATCCT
42	SC2M1-42_LEFT_16065	GGAGTATGCTGATGCTTTTCATTTGTAC
	SC2M1-42_RIGHT_16648	CGGTTTCTGCTGCAAAAAGCTT
46	SC2M1-46_LEFT_17752	TGGAGAAAAGCTGCTTTTATTTCACCT
	SC2M1-46_RIGHT_18275	GCTTCTTCGCGGGTGATAAACA
50	SC2M1-50_LEFT_19311	TGCATTCCACACACCAGCTTTT
	SC2M1-50_RIGHT_19866	ATTAGCAGCAATGTCCACACCC
54	SC2M1-54_LEFT_20990	TGATTGGTGATTGTGCAACTGTACA
	SC2M1-54_RIGHT_21562	TGTTCTGTTAGTTGTTAACAGAACATCA
58	SC2M1-58_LEFT_22563	ACTTGTGCCCTTTTGGTGAAGT
	SC2M1-58_RIGHT_23128	TGCTGGTGCAATGAGAAGTTCA
62	SC2M1-62_LEFT_24095	GCTGCTAGAGACCTCATTGTGTC
	SC2M1-62_RIGHT_24623	AAGCTCTGATTTCTGCAGCTCT
66	SC2M1-66_LEFT_25665	CTCACACCTTTTGCTGTTGCT
	SC2M1-66_RIGHT_26224	GTGCTTACAAAGGCACGCTAGT
70	SC2M1-70_LEFT_27254	TTATGAGGACTTTTAAAGTTTCCATTGGA
	SC2M1-70_RIGHT_27808	AGCAGAAAAGCTAAAAGCACA
74	SC2M1-74_LEFT_28918	TGATGCTGCTTCTGCTTTGCTG
	SC2M1-74_RIGHT_29469	TCTGCAGCAGGAAGAAGTCA
78	SC2M1-52_LEFT2_20349	AGTCAGTTAGGTGTTTACATCTACTGA
	SC2M1-52_RIGHT2_20798	TTTTGCGACATTCATTATGCCT

Pool 3

PCR	Name	Sequence
3	SC2M1-3_LEFT_827 SC2M1-3_RIGHT_1395	AACAACCTTCTGTGGCCTGATG TCTGAATTGTGACATGCTGGACA
11	SC2M1-11_LEFT_4126 SC2M1-11_RIGHT_4658	GGGTGATGTTGTTCAAGAGGGT ACCGAGCAGCTTCTTCCAAATT
15	SC2M1-15_LEFT_5677 SC2M1-15_RIGHT_6172	TGTTATGATGTCAGCACCCACTG AGCCACCACATCACCATTTAAGT
19-2	SC2M1-19b_LEFT_7235 SC2M1-19_RIGHT_7694	TGCAGAGTGGTTTTGGCATATATTCT ACTGTAGTGACAAGTCTCTCGCA
23	SC2M1-23_LEFT_8778 SC2M1-23_RIGHT_9330	TTAGCCAGCGTGGTGGTAGTTA TCTACACCACAGAAAACCTCTGGT
27	SC2M1-27_LEFT_10318 SC2M1-27_RIGHT_10837	GCTTAAGGTTGATACAGCCAATCCT AACGGCAATTCAGTTTGAGCA
31	SC2M1-31_LEFT_11810 SC2M1-31_RIGHT_12335	GGCAAACCTTGATCAAAGTAGCC TTGCCCTCTTGTCTCAGATCT
35	SC2M1-35_LEFT_13366 SC2M1-35_RIGHT_13861	AAACACAGTCTGTACCCTCTGC TGTCACAATTACCTTCATCAAAATGCC
39	SC2M1-39_LEFT_14888 SC2M1-39_RIGHT_15391	ACGATGGTGGCTGTATTAATGCT GGTGTGACAAGCTACAACACGT
43	SC2M1-43_LEFT_16518 SC2M1-43_RIGHT_17087	AAATACATGTGTTGGTAGCGATAATGTT GGTGGTCCCTGGAGTGTAGAAT
47	SC2M1-47_LEFT_18148 SC2M1-47_RIGHT_18668	GGTTTATGTGTTGACATACCTGGCA CATAGACAACAGGTGCGCTCAG
51	SC2M1-51_LEFT_19725 SC2M1-51_RIGHT_20255	TGATGGTGTGATGAGAATTGTTTGAA TCAATTTCCATTGACTCCTGGGT
55	SC2M1-55_LEFT_21421 SC2M1-55_RIGHT_21916	AGGGGTACTGCTGTTATGCTTTAAA AAGTAGGGACTGGGTCTTCGAA
59	SC2M1-59_LEFT_22986 SC2M1-59_RIGHT_23519	CCGGTAGCACACCTTGTAATGG CCCCATTAAACAGCCTGCACG
63	SC2M1-63_LEFT_24493 SC2M1-63_RIGHT_25003	AAATGATATCCTTTACAGTCTTGACAAA TGAGTCTAATTCAGGTTGCAAAGGA
67	SC2M1-67_LEFT_26096 SC2M1-67_RIGHT_26590	AAAATTGTTGATGAGCCTGAAGAACA ACTAGGTTCCATTGTTCAAGGAGC
71	SC2M1-71_LEFT_27650 SC2M1-71_RIGHT_28203	TGTTTCATCAGACAAGAGGAAGTTCA ACGAAACAACGCACTACAAGACT
75	SC2M1-75_LEFT_29344 SC2M1-75_RIGHT_29848	TGACGCATACAAAACATTTCCAC AAAATCACATGGGGATAGCACTACT

Pool 4

PCR	Name	Sequence
4	SC2M1-4_LEFT_1262 SC2M1-4_RIGHT_1840	ACGGGCGATTTTGTAAAAGCCA TCACCAATATTCACGGCACCTTT
8	SC2M1-8_LEFT_2932 SC2M1-8_RIGHT_3461	ACTTACACCACCTGGGCATTGATT CTGCAACACCTCTCCATGTTT
12	SC2M1-12_LEFT_4519 SC2M1-12_RIGHT_5017	TGGTGCTAGATTTTACTTTTACACCAGT CACAACTTGCCTGGAGGTTA
16	SC2M1-16_LEFT_6030 SC2M1-16_RIGHT_6544	ACGCAAGCTTCGATAATTTAAGTTTGT TGTGTGGCCAACCTCTCTGTGA
20	SC2M1-20_LEFT_7560 SC2M1-20_RIGHT_8128	GGTCCTTTTATGCTATGCTAATGGAGG TGCAAGTTACAGTCTCGAGTT
24	SC2M1-24_LEFT_9203 SC2M1-24_RIGHT_9734	GATTCTGAGTACTGTAGGCACGG AGAACCAATAGAAATGCTTTGTGGAAA
28	SC2M1-28_LEFT_10697 SC2M1-28_RIGHT_11209	GGAGACAGGTGGTTTCTCAATCG AGCTACAGTGGCAAGAGAAGGT
32	SC2M1-32_LEFT_12201 SC2M1-32_RIGHT_12719	AGTTGAAGAAGCTTTGAATGTGGCT TCTGTGCTAGTGAACAGGACT
36	SC2M1-36_LEFT_13727 SC2M1-36_RIGHT_14232	GCTGTTGCTAAACATGACTTCTTTAAGT AGGCTTTGTTAAGTCAGTGTCAACA
40	SC2M1-40_LEFT_15264 SC2M1-40_RIGHT_15771	TGTAGAAAAACCTCACCTTATGGG AGCCACTAGACCTTGAGATGCA
44	SC2M1-44_LEFT_16948 SC2M1-44_RIGHT_17458	CCTACACTAGTGCCACAAGAGC GTGCAGTAATTGAGCAGGGTC
48	SC2M1-48_LEFT_18506 SC2M1-48_RIGHT_19038	GACTTCCTTGGAAATGAGTGCCT ACCAATGCTGGAAGAACTGGG
52	SC2M1-52_LEFT_20124 SC2M1-52_RIGHT_20698	TGGAGAAGCCGTAACACACAGT GATTAGGCATAGCAACACCCGG
56	SC2M1-56_LEFT_21775 SC2M1-56_RIGHT_22345	TGGGACCAATGGTACTAAGAGGT ACCAGCTGTCCAACCTGAAGAA
60	SC2M1-60_LEFT_23379 SC2M1-60_RIGHT_23876	ACCAGGTTGCTGTTCTTTATCAGG CAGCTATTCAGTTAAAGCACGGT
64	SC2M1-64_LEFT_24858 SC2M1-64_RIGHT_25369	GCACACACTGGTTTGTAAACACAA TTTGACTCCTTTGAGCACTGGC
68	SC2M1-68_LEFT_26454 SC2M1-68_RIGHT_27004	TCCTGATCTTCTGGCTAAACGAACT ATGTCACAGCGTCTAGATGGT
72	SC2M1-72_LEFT_28066 SC2M1-72_RIGHT_28649	TTGAATTGTGCGTGGATGAGGC TAGCACCATAGGGAAGTCCAGC
76	SC2M1-1_LEFT2_1 SC2M1-1_RIGHT2_495	TTAAAGGTTTATACCTTCCCAGG CGAGCATCCGAACGTTTGTATGA

Pool 5

PCR	Name	Sequence
5W	W1_5L_1457	GTAAGGGTGGTCGCACTATTGC
	W1_5R_1969	TTGTTATAGCGGCTTCTGTAAAAAC
81*	SC2M1-19a_LEFT_6957	TGGTTTTACTATTAAGTGTTCGCTAGGT
	SC2M1-19a_RIGHT_7393	TCGGGGCCATTTGTACAAGATT
81*	SC2M1-19a_LEFT_6957	TGGTTTTACTATTAAGTGTTCGCTAGGT
	SC2M1-19a_RIGHT_7393	TCGGGGCCATTTGTACAAGATT
83	SC2M1-21a_LEFT_7984	AGGCATTAGTGTCTGATGTTGGTG
	SC2M1-21a_RIGHT_8384	TGACTTTTTGCTACCTGCGCAT
28W	W1_28L_9659	TCACACCTTAGTACCTTTCGGATAAC
	W1_28R_10207	GGTTAAGCATGTCTTCAGAGGTGC
85	SC2M1-34a_LEFT_12994	GTAGTTTAGCTGCCACAGTACGT
	SC2M1-34a_RIGHT_13399	AACCTTCCACATACCGCAGAC
42	SC2M1-42_LEFT_16065	GGAGTATGCTGATGCTTTTCATTTGTAC
	SC2M1-42_RIGHT_16648	GCGTTTTCTGCTGCAAAAAGCTT
89	SC2M1-49a_LEFT_18711	CCTGTTGGCATCATTCTATTGGATTT
	SC2M1-49a_RIGHT_19112	GTCACTACAAGGTGTGCATCA
91	SC2M1-50a_LEFT_19181	TGCCTATTTGGAATTGCAATGTGCG
	SC2M1-50a_RIGHT_19569	AAACCCACAAGCTAAAGCCAGC
93	SC2M1-51a_LEFT_19661	TTTGATGGACAACAGGGTGAAGT
	SC2M1-51a_RIGHT_20098	GCTTGTTGGGACCTACAGATGG
60W	W1_60L_21029	GGATCTCATTATTAGTGATATGTACGACCC
	W1_60R_21562	TTGTTTCGTTTAGTTGTTAAACAAGAACATC
64W	W1_64L_22457	CAAAGTGTACGTTGAAATCCCTCACTG
	W1_64R_22993	TGCTACCGCCTGATAGATTTC
98	SC2M1-67a_LEFT_25910	GGCACAACAAGTCTATTCTGAAC
	SC2M1-67a_RIGHT_26276	CGTACCTGTCTTCCGAAACG
100	SC2M1-69a_LEFT_26846	TGTGGTCATTCAATCCAGAACTAACA
	SC2M1-69a_RIGHT_27226	ACCTGAAAGTCAACGAGATGAAACA
102	SC2M1-70a_LEFT_27252	TTATGAGGACTTTTAAAGTTTCCATTTGGA
	SC2M1-70a_RIGHT_27644	AGGTGAAACTGATCTGGCAGCT
71	SC2M1-71_LEFT_27650	TGTTTCATCAGACAAGGAAAGTTCA
	SC2M1-71_RIGHT_28203	ACGAACAACGCACTACAAGACT
95	0_1b	TTAAAGGTTTATACCTTCCCAGGTA
	W1_1R_490	CATCCGAACGTTTGATGAACAC
36	SC2M1-36_LEFT_13727	GCTGTTGCTAAACATGACTTCTTTAAGT
	SC2M1-36_RIGHT_14232	AGGCTTTGTTAAGTCAGTGTCAACA

Pool 6

PCR	Name	Sequence
6W	W1_6L_1819	AGGTGCCTGGAATATTGGTGAAC
	W1_6R_2345	ATGATAGAGTCAGCACACAAGC
7ab	SC2M1-7a_LEFT_2491	AGGGAGAAACACTTCCCACAGA
	SC2M1-7b_RIGHT_3165	AGCAGAAGTGGCACCAAATTC
84	SC2M1-21b_LEFT_8240	TCAATCTGACATAGAAGTTACTGGCG
	SC2M1-21b_RIGHT_8618	GCAGCAACAAAAAGGAACACAAGT
26W	W1_26L_8999	CTTGTTGTTTTGGCTGTGAATG
	W1_26R_9459	CATAAAATAGTAGGCAAGGCATGTTACTAC
40W	W1_40L_13986	CGCCAAGCTTTGTTAAAAACAGTAC
	W1_40R_14503	TGTACAACACCTAGCTCTCTGAAGTG
86	SC2M1-34b_LEFT_13245	CTGTACTGCCGTTGCCACATAG
	SC2M1-34b_RIGHT_13620	CGTCCTTTTCTTGGAAAGCGACA
90	SC2M1-49b_LEFT_18955	TGCGGCTGTAGAAAGGTTCAA
	SC2M1-49b_RIGHT_19331	AAAAGCTGGTGTGGAATGCA
92	SC2M1-50b_LEFT_19395	AGTCTCATGAAAAACAAGTAGTGCA
	SC2M1-50b_RIGHT_19820	TGGTACTGGTTAATGTTGCGCT
94	SC2M1-51b_LEFT_19957	AACGATTTGTGCCACTCACT
	SC2M1-51b_RIGHT_20373	TCAGTAGATGTAACACCTAACTGACT
99	SC2M1-67b_LEFT_26128	TCACACAATCGACGGTTCATCC
	SC2M1-67b_RIGHT_26541	GTACCGTTGGAATCTGCCATGG
101	SC2M1-69b_LEFT_27080	TAGCAGGTGACTCAGGTTTTGC
	SC2M1-69b_RIGHT_27443	AAGCTCACAAAGTAGCGAGTGTT
9	SC2M1-9_LEFT_3306	TGGAACCTACACGATTTGTTCAAGC
	SC2M1-9_RIGHT_3878	CAGCGATCTTTTGTCAACTTGCT
75	SC2M1-75_LEFT_29344	TGACGCATACAAAACATTCCCAC
	SC2M1-75_RIGHT_29848	AAAATCACATGGGGATAGCACTACT
70b	SC2M1-70b_LEFT_27497	TCTTCTGGAACATACGAGGGCA
	W1_34R_28006_1	AGGACACGGGTCACTCAACTAC
95	0_1b	TTAAAGGTTTATACCTTCCCAGGTA
	W1_1R_490	CATCCGAACGTTTGATGAACAC
19-3	SC2M1-19b_LEFT_7235	TGCAGAGTGGTTTTTGGCATATATTCT
	W1_21R_7771	GATGGATGGAACATTCTTCACTG
62-2	SC2M1-62a1_LEFT_23993	ACCAAGCAAGAGGTCAATTTATTGAAGA
	W1_30R_24625_1	CAGAAGCTCTGATTTCTGCAGC

\*This amplicon is intentionally doubled due to its reduced efficiency—add 2 parts of this primer set to the pool.

## Appendix E – AMPure XP bead clean-up

Bead-based clean-ups are done at several steps throughout the protocols presented. This covers the basic clean-up steps, make sure to check the specific protocol for the ratio of beads to use.

Depending on the number of samples, the AMPure XP bead clean-up takes about 30-40 minutes.

Required reagents for bead-based clean-up

Company	Product	Catalog number
Beckman Coulter	Agencourt AMPure XP beads	A63882
	10mM Tris-HCl pH 8.0	

1. Allow AMPure XP beads to warm to room temperature for at least 30 minutes before using.
2. Vortex AMPure XP beads to re-suspend.
3. Add appropriate ratio of re-suspended AMPure XP beads to the ligation reaction. Mix well by pipetting up and down at least 10 times.
4. Incubate for 5 minutes at room temperature.
5. Place the tube/plate on an appropriate magnetic stand to separate the beads from the supernatant. If necessary, quickly spin the sample to collect the liquid from the sides of the tube or plate wells before placing on the magnetic stand.
6. After the solution is clear (about 5 minutes), carefully remove and discard the supernatant. Be careful not to disturb the beads that contain DNA targets (do not discard beads).
7. Add 200  $\mu$ L of freshly prepared 80% Ethanol to the tube/plate while in the magnetic stand.
8. Incubate at room temperature for 30 seconds, then carefully remove and discard the supernatant.
9. Add another 200  $\mu$ L of freshly prepared 80% Ethanol to the tube/plate while in the magnetic stand.
10. Incubate at room temperature for 30 seconds, then carefully remove and discard the supernatant.
11. Air dry the beads for 2 minutes while the tube/plate is on the magnetic stand and with the lid(s) open.  
Caution: Do not over dry the beads. This may result in lower recovery of DNA target.
12. Remove the tube/plate from the magnet. Elute the DNA target from the beads by adding appropriate volume of 10mM Tris-HCl or water.
13. Mix well by pipetting up and down or on a vortex mixer. Incubate for 2 minutes at room temperature. If necessary, quickly spin the sample to collect liquid from the sides of the tube or plate wells before placing on the magnetic stand.
14. Place the tube/plate on the magnetic stand.
15. After the solution is clear (about 5 minutes), transfer to a new tube.

## Appendix F – Quantitation using Qubit

Quantitation is done at several various steps throughout the protocols included and this protocol can be used anytime quantitation is indicated.

### Required reagents

Company	Product	Catalog number
Thermo Fisher	dsDNA HS assay kit	Q32854
Thermo Fisher	dsDNA BR assay kit	Q32850
Thermo Fisher	Qubit assay tubes	Q32856

Note: depending on the sample, either the high sensitivity (HS) or broad range (BR) kit may be used, the protocols are the same the only difference is the reagents.

Quantitation takes about 10-20 minutes depending on the number of samples.

### Procedure

1. Set up the required number 0.5 mL Qubit assay tubes for standards and samples. Note: the standards require two tubes.
2. Label tube lids. Do not label the side of the tube as this could interfere with the sample read.
3. Prepare the Qubit working solution by diluting the Qubit dsDNA HS reagent 1:200 in Qubit dsDNA HS buffer. Use a clean plastic tube each time you prepare Qubit working solution. Do not mix the working solution in a glass container.

The final volume in each tube must be 200 uL. Each standard tube requires 190 uL of Qubit working solution and each sample tube requires anywhere from 180-199 uL. Prepare sufficient Qubit working solution to accommodate all standards and samples.

4. Add 190 uL of Qubit working solution to each of the tubes used for standards.
5. Add 10 uL of each qubit standard to the appropriate tube, mix by vertexing 2-3 seconds.
6. Add Qubit working solution to individual assay tube, mix by vertexing 2-3 seconds.

Your sample can be anywhere from 1-20 uL. Add a corresponding volume of Qubit working solution to each assay tube: anywhere from 180-199 uL.

7. Add each sample to the assay tubes containing the correct volume of Qubit working solution, then mix by vertexing 2-3 seconds. The final volume in each tube should be 200 uL.
8. Allow all tubes to incubate at room temperature for 2 minutes.
9. Sample concentration can now be measured on the Qubit Fluorometer.

## Appendix G – CENTRI-SEP 96 Protocol



### **CENTRI-SEP 96 Protocol**

CENTRI•SEP 96 plates must be allowed to equilibrate to room temperature before use. We recommend that the plates be removed from the refrigerator at the same time the sequencing reactions are initiated. This will allow sufficient time for the plates to warm.

1. Remove the adhesive foil from the bottom and then from the top of the CENTRI-SEP 96 plate.
2. Stack the CENTRI-SEP 96 plate on top of a 96-well wash plate and centrifuge at 1500 x g for 2 minutes. Use an external timer and start timing when the rotor has reached the set speed. Discard the liquid in the wash plate. The gel matrix in the wells should appear opaque at this point.
3. Transfer the samples (20  $\mu$ L or less) to the individual wells in the CENTRI-SEP 96 plate, taking care to place the samples in the centers of the gel beds.
4. Stack the CENTRI-SEP 96 plate on top of a 96-well collection plate and centrifuge at 1500 x g for 2 minutes.
5. Remove the 96-well collection plate containing the cleaned samples and dry in a speed-vac equipped with the appropriate rotor. Alternatively the plate can be sealed for storage.