

JAN 09, 2024

L-4 LEECH TESTING

REDI-NET
Consortium¹

¹REDI-NET Consortium



REDI-NET Consortium

University of Notre Dame, Naval Medical Research Center, Wal...

DISCLAIMER

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Protocol status: Working We use this protocol and it's working

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ABSTRACT

This protocol details standard operating procedure for leech testing.

GUIDELINES

OBJECTIVE

To outline the procedures for properly using the Oxford Nanopore Sequencing platforms (GridION or MinION Mk1C) to sequence gDNA and TNA extracted from collected leech samples.

SUMMARY/SCOPE

This SOP provides guidance on procedures of Oxford Nanopore sequencing to generate sequencing reads for downstream data analysis and pathogen detection.

RESPONSIBLE PERSON

Principal Investigator, Study Coordinator, Entomology Component Lead, Managers

PROTOCOL integer ID:

89963

Keywords: gDNA
PREPARATION, TNA
PREPARATION, cDNA
SYNTHESIS, Purification of
double-stranded cDNA,
SEQUENCING LIBRARY
PREPARATION

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0093

USAMRAA

Grant ID: HT9425-23-C-0059

Note

NOTE: All study procedures must be conducted in compliance with national and local policies for the prevention and control of COVID-19 infection.

MAINTENANCE OF EQUIPMENT

CAUTION ON RNA HANDLING:

- 1. RNases are very stable and difficult to inactivate and only minute amounts are sufficient to destroy RNA.
- 2. Care should be taken to avoid inadvertently introducing RNases into the samples during or after the purification procedure.
- 3. Clean the work surfaces with RNA Zap to remove nucleases, then wipe the surfaces with 70% to 100% molecular biology grade ethanol to remove additional contaminants.

HANDLING ENZYMATIC REACTIONS

Reagents containing enzymes should be handled On ice before mixed and transferred to the assigned activation temperature.

REFERENCES

REDI-NET Overview Summary

Double-stranded cDNA synthesis (NEB first and second strand cDNA synthesis protocols):

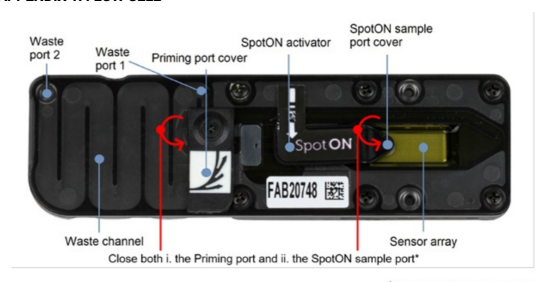
- NEBNext Ultra II RNA First Strand synthesis manual E7771
- NEBNext Ultra II Non-directional RNA Second Strand synthesis manual E6111
- ezdnase_Pl

Oxford Nanopore Manufacturer's protocols:

- Ligation sequencing gDNA Native Barcoding Kit 96 V14 (SQK-NBD114.96)-minion.
- ligation-sequencing-gdna-native-barcoding-v14-sqk-nbd114-96-NBE_9171_v114_revG_15Sep2022-minion
- ligation-sequencing-gdna-native-barcoding-v14-sqk-nbd114-96-NBE_9171_v114_revG_15Sep2022-gridion

APPENDICES

APPENDIX 1. FLOW CELL



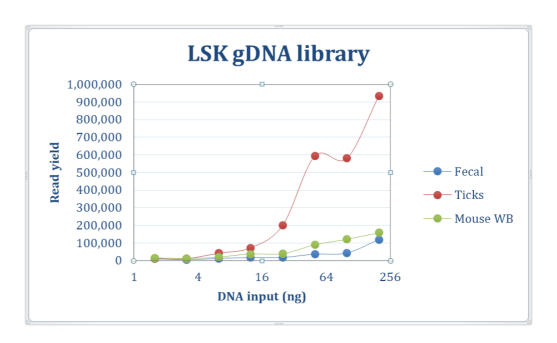
*Both ports are shown in a closed position

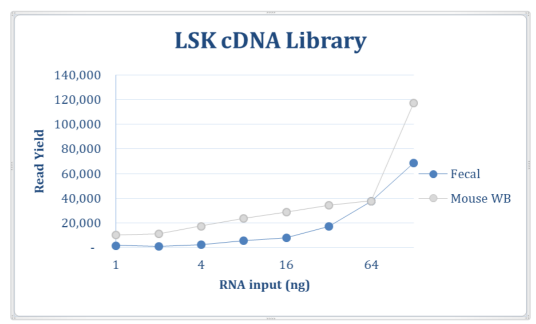
APPENDIX 2. cDNA END-PREP MASTER MIX PREPARATION

A	В	С
Component	Volume for 1 reaction	Volume for n+1 reactions
cDNA sample	20 μΙ	20 μΙ
Nuclease-free water	30 μΙ	μl
Ultra II End-prep reaction buffer	7 μΙ	µl
Ultra II End-prep enzyme mix	3 μΙ	µl
Final total volume	60 µl	µl

APPENDIX 3. EXPECTED OUTCOMES

The DNA or RNA inputs vs the sequencing read yields.





MATERIALS

EQUIPMENT AND MATERIALS

Note

NOTE: If product number is listed, please ensure use of this or equivalent product.

A	В
Equipment	Mfg / Product #
Oxford Nanopore GridION or MinION Mk1C device	Oxford Nanopore Technologies, GRD-CapEx or Oxford Nanopore Technologies, M1CCapEx
Computer monitor (with HDMI port or Display port), mouse and keyboard	Locally sourced
MinKNOW - software equipped already in the GridION and MinION Mk1C device	Oxford Nanopore Technologies
Ice bucket with ice	Locally sourced
Qubit fluorometer	ThermoFisher, Q33238 or equivalent
DynaMag-2 magnet	Invitrogen, 12321D or equivalent
DynaMag-96 Side Magnet	Invitrogen, 12331D or equivalent
Hula sample mixer	ThermoFisher, 15920D
Microplate centrifuge	Locally sourced
Timer	Locally sourced
Thermal cycler	Locally sourced
96-well PCR plate holder	Locally sourced
P1000 pipette and tips	Locally sourced
P200 pipette and tips	Locally sourced
P20 pipette and tips	Locally sourced
P10 pipette and tips	Locally sourced
P10 8-channel pipette	Locally sourced
P300 8-channel pipette	Locally sourced

A	В	С	
Material	Description	Mfg / Product #	
200 ng DNA from a sample	Per sample from SOP B- 2 (gDNA Preparation)	REDI-NET DNA sample	
20 ul eluents from negative control extraction	From SOP B-2 (gDNA Preparation)	REDI-NET negative control	

A	В	С
100 ng DNA from positive control extraction	From SOP B-2 (gDNA Preparation)	REDI-NET positive control
160 ng RNA from a sample	Per sample from SOP B-2 (TNA preparation)	REDI-NET RNA sample
40 ng RNA from positive control extraction	from SOP B-2 (TNA preparation)	REDI-NET negative control
8 µl total nucleic acid negative control extraction	From SOP B-2 (TNA preparation)	REDI-NET positive control
10 µl total nucleic acid	Per sample from SOP B- 2 (TNA Preparation)	REDI-NET TNA sample
10 µl total nucleic acid from negative control extraction	From SOP B-2 (TNA Preparation)	REDI-NET negative control
10 µl total nucleic acid from positive control extraction	from SOP B-2 (TNA Preparation)	REDI-NET positive control
Native Barcoding Kit 96 V14	(Sequencing Library Preparation)	Oxford Nanopore, SQK- NBD114.96
ezDNase	(cDNA synthesis)	ThermoFisher, Invitrogen 11766051
NEBNext Ultra II RNA First Strand Synthesis Module	(cDNA synthesis)	New England Biolabs, E7771L
NEBNext Ultra II Non- Directional RNA Second Strand Synthesis Module	(cDNA synthesis)	New England Biolabs, E6111L
Random primer mix (Random hexamer and poly-T mixture)	(cDNA synthesis)	New England Biolabs, S1330
USB Dithiothreitol (DTT), 0.1M Solution	(cDNA synthesis)	ThermoFisher,707265ML
Agencourt AMPure XP beads	(Sequencing Library Preparation)	Beckman Coulter, A63881
NEBNext End repair / dA- tailing Module	(Sequencing Library Preparation)	New England Biolabs, E7546L
NEBNext FFPE Repair Mix	(Sequencing Library Preparation)	New England Biolabs, M6630L
NEB Blunt/TA Ligase Master Mix	(Sequencing Library Preparation)	New England Biolabs, M0367L
NEBNext Quick Ligation Module	(Sequencing Library Preparation)	New England Biolabs, E6056L
R10.4.1 flow cells	Flow cells for sequencing experiment (consumable)	Oxford Nanopore, FLO- MIN114
low DNA binding tubes	1.5 mL (<i>consumable</i>)	Eppendorf, 022131021 or equivalent
low DNA binding tubes	2.0 mL (<i>consumable</i>)	Eppendorf, 022431048 or equivalent

A	В	С
PCR tubes	0.2 mL thin-walled (<i>consumable</i>)	Eppendorf, 951010006 or equivalent
PCR plate	96 well, low DNA binding, semi-skirted with heat seals (<i>consumable</i>)	Eppendorf, 0030129504 or equivalent
BRAND Self-adhesive Plate Sealing Film	Aluminum (<i>consumable</i>)	Fisher Scientific, 13-882-329
Clear Adhesive Film	For PCR plate sealing	ThermoFisher, 4306311
Qubit Assay Tubes	For Qubit DNA/RNA measurement (consumable)	Thermo Fisher, Q32856
Qubit 1X dsDNA HS Assay Kit	(consumable)	ThermoFisher, Q33230
Qubit RNA HS Assay Kit	(consumable)	ThermoFisher, Q32852
Nuclease-free water	To prepare ethanol dilutions (<i>consumable</i>)	Locally sourced
Freshly prepared 80% ethanol in nuclease-free water	Prepared from 100% molecular biology grade ethanol (<i>consumable</i>)	Locally sourced
Freshly prepared 70% ethanol in nuclease free water	Prepared from 100% molecular biology grade ethanol (<i>consumable</i>)	Locally sourced
Data sheets	REDI-NET DCS B-4 Testing	REDI-NET Data Portal

Equipment	
Qubit Fluorometer	NAME
Fluorometer	TYPE
Invitrogen	BRAND
Q33238	SKU
https://www.thermofisher.com/order/catalog/product/Q33238#/Q3323	38 LINK

Equipment	
DynaMag™-2 Magnet	NAME
Magnet	TYPE
DynaMag™	BRAND
12321D	SKU
https://www.thermofisher.com/order/catalog/product/12321D	LINK

Equipment	
Hula mixer	NAME
Mixer	TYPE
Invitrogen	BRAND
15920D	SKU
Any rotator mixer	SPECIFICATIONS

- Native Barcoding Kit 96 V14 Oxford Nanopore

 Technologies Catalog #SQK-NBD114.96
- 🔀 ezDNase™ Enzyme **Thermo Fisher Catalog #11766051**
- NEBNext Ultra II RNA First Strand Synthesis Module 96 rxns New England Biolabs Catalog #E7771L
- NEBNext Ultra II Non-Directional RNA Second Strand Synthesis Module 100 rxns New England Biolabs Catalog #E6111L
- igotimes Random primer mix New England Biolabs Catalog #S1330S
- USB Dithiothreitol (DTT) 0.1M Solution Thermo Fisher Scientific Catalog #707265ML

- Agencourt AMPure XP beads Beckman Coulter Catalog #A63881
- NEBNext Ultra II End Repair/dA-Tailing Module 96 rxns **New England**Biolabs Catalog #E7546L
- NEBNext FFPE DNA Repair Mix 96 rxns New England Biolabs Catalog #M6630L
- Blunt/TA Ligase Master Mix 250 rxns New England Biolabs Catalog #M0367L
- NEBNext Quick Ligation Module 100 rxns New England Biolabs Catalog #E6056L
- Nanopore Flow Cell R10.4.1 Oxford Nanopore
 Technologies Catalog #FLO-MIN114
- DNA LoBind Tubes 2.0 ml Eppendorf Catalog #022431048
- Eppendorf PCR
 Tubes Eppendorf Catalog #951010006
- 96 well LoBind PCR plates Semiskirted Eppendorf Catalog #0030129504
- NEBNext Microbiome DNA Enrichment Kit 6 rxns New England Biolabs Catalog #E2612S

the

- RNaseOUT™ Recombinant Ribonuclease Inhibitor **Thermo Fisher** Scientific Catalog #10777019
- BRAND™ Self-adhesive Plate Sealing Film **Fisher**Scientific Catalog #13-882-329
- MicroAmp™ Clear Adhesive Film **Thermo Fisher** Scientific Catalog #4306311

Qubit assay tubes Thermo Fisher Scientific Catalog #Q32856

Qubit 1X dsDNA High Sensitivity Assay Kit **Thermo Fisher**Scientific Catalog #Q33230

Qubit RNA HS (High Sensitivity) assay Thermo Fisher Scientific Catalog #Q32852

SAFETY WARNINGS

•

RISKS AND PERSONAL PROTECTION

Gloves should be worn all the time when handling samples.

BEFORE START INSTRUCTIONS

BEFORE START

- 1. Check the DNA and RNA concentrations in each sample of total nucleic acid (TNA) extraction.
- 2. If the concentrations are detectable, choose the sequencing approach following the table below.
- 3. If DNA or RNA concentration is not detectable, prepare the library for the detectable one.
- Use sections <u>gDNA PREPARATION</u> and <u>TNA PREPARATION</u> for gDNA and TNA preparation, respectively, then subject the prepared gDNA and TNA to Section <u>SEQUENCING LIBRARY PREPARATION</u>.

A	В	С	D	E
		DNA concentration (ng/ul)		
		< 1 ng/ul	1-10 ng/ul	> 10 ng/ul
	< 4 ng/ul	TNA	DNA	DNA
RNA concentration (ng/ul)	4-20 ng/ul	TNA	TNA	TNA
	>20 ng/ul	TNA	TNA	TNA

gDNA PREPARATION

1



When the RNA concentration of the sample is lower than the detectable range of the Qubit High Sensitivity Assay (< Δ 0.01 ng/ μ I), the sample is subjected to gDNA sequencing. The cDNA synthesis can be skipped.

- When the DNA concentration > $\boxed{\bot}$ 10 ng/ μ I, calculate the required volume of $\boxed{\bot}$ 200 ng DNA, then transfer the volume to a new 200 μ I PCR tube or a well of a 96-well PCR plate. Adjust the volume with nuclease-free water to a final volume of $\boxed{\bot}$ 20 μ L.
- 3 Prepare Δ 100 ng gDNA from positive control extraction in Δ 20 μL nuclease-free water in a new 200μl PCR tube or a well of a 96-well PCR plate.
- 4 Transfer Δ 20 μL negative control extraction to a new tube or a well of a 96-well PCR plate.
- 5 All samples are subjected to section SEQUENCING LIBRARY PREPARATION.

TNA PREPARATION

- To prepare TNA for sequencing both cDNA and gDNA, cDNA needs to be prepared separately and then mixed with TNA from the original sample.
- 7 Prepare cDNA following section <u>cDNA SYNTHESIS</u> (positive control and negative control included) until step 40.
- 8 Transfer Δ 10 μ L double-stranded cDNA of section cDNA SYNTHESIS step 40 to a new 200 μ l PCR

- 9 Subject the Δ 20 μL double-stranded cDNA/TNA mixture to section <u>SEQUENCING LIBRARY</u> PREPARATION.

Note

NOTE: Twenty-four samples must be pooled in one sequencing run to make the most out of a sequencing flow cell. For collecting 24 samples, the samples from gDNA and TNA preparations can be placed in the same 96-well plate for End-prep and Barcode Ligation before pooling, then pooled for the subsequent steps of library preparation.

cDNA SYNTHESIS: DNase treatment

- Prepare 40 ng RNA from positive control extraction and adjust the volume to final 40 ng RNA from positive control extraction and adjust the volume to final 40 ng with nuclease-free water in a new 200µl PCR tube or a well of a 96-well PCR plate.
- Transfer Δ 8 μL negative control extraction to a new tube or a well of a 96-well PCR plate.
- Remove contaminated DNA (~ 15 mins): Thaw total nucleic acid, 10x ezDNase Buffer, and DTT on 7m 5s ice at Room temperature. Vortex 10x ezDNase Buffer and DTT briefly, spin down by centrifugation for 00:00:05, and place On ice ezDNase is not frozen and should be placed On ice before use. Set up thermal cycler programs: 37 °C, 00:02:00, and 55 °C, 00:05:00.
- 13.1 Mix the following components in an RNase-free tube or plate. For processing multiple samples, make



a master mix for 10× ezDNase buffer and ezDNase with 10% overage. Aliquot the master mix into the wells of a 96-well plate, then add TNAs.

A	В
Component	Volume
10× ezDNase Buffer	1 μΙ
ezDNase	1 μΙ
RNA from step 6	8 μΙ
Total volume	10 μΙ

13.2 Gently mix the samples then centrifuge the tube (Include a reaction for extraction positive control and negative control of each batch nucleic acid extraction).

13.3 Incubate the sample for 00:02:00 at 37 °C

2m



13.4 Add $\mathbb{Z}_{1 \mu L}$ 100mM DTT into the reaction tube.



13.5 Incubate the sample at \$\ 55 \circ\$ for \ \ 00:05:00 \ to inactivate the enzyme.

F....



Chill the tube On ice to bring the sample to Room temperature, then spin down and place the tube On ice.

cDNA SYNTHESIS: First strand cDNA Synthesis (~ 1hr)

14 <u>BEFORE START</u>: Thaw 60 μM stock Random Primer Mix (NEB, S1330S) at Room temperature . DC 5s NOT USE the Random Primer provided by the NEBNext First Strand Synthesis Module. Thaw Random

Primer Mix solution, NEBNext First Strand Reaction Buffer, NEBNext Second Strand Reaction Buffer at Room temperature then place on ice. Vortex the vials briefly, spin done by centrifugation for 00:00:05, and place on ice. First and Second Strand Enzyme Mix are not frozen, should be briefly centrifuged and placed on ice before use.

15

Add the following reagents into the ezDNase-treated RNA from step 13.6. For processing multiple samples, make a master mix for the 60 μ M Random Primer Mix and nuclease-free water with 10% overage.

A	В
Component	Volume
ezDNase treated RNA	10 μΙ
60 μM Random Primer	1 μΙ
Nuclease free water	3 μΙ
Total volume	14 μΙ

16

Mix gently, spin down and incubate at 65 °C for 00:05:00 . Chill on ice, spin down again 5m and place on ice.



Add the following components in the indicated order, if multiple reactions will be processed at the same time, make a master mix with a 10% overage:

A	В
Component	Volume
NEBNext First Strand Synthesis Reaction Buffer	4 μΙ
NEBNext First Strand Synthesis Enzyme Mix	2 μΙ
Total volume	20 μΙ

18 Mix gently and spin down.

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19 Incubate the tube for 600:10:00 at 8 25 °C followed by 600:15:00 at 8 42 °C



20 Terminate the reaction by heating at \$\ \bigset* 70 °C for \ \cdot 00:15:00 15m



- 21 Place the tube § On ice or pre-chilled freezer block.
- 22 Continue immediately with the second strand synthesis reaction as described below.

cDNA SYNTHESIS: Second strand cDNA Synthesis (~ 1hr)

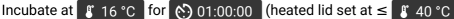
23 Pipette the following components directly into the first strand reaction tube (with A 20 µL mixture) § On ice in the indicated order, if multiple reactions will be processed at the same time, make a master mix with a 10% overage:

A	В
Component	Volume
5x NEBNext Second Strand Synthesis Read	etion Buffer 5 µl
NEBNext Second Strand Synthesis Enzyme	e Mix 2.5 μl
Nuclease-free water	22.5 µl
Final total volume	50 μΙ

24 Mix gently and centrifuge briefly.



25m







25

26 Proceed with cDNA purification or store the reaction mixture at 8 -20 °C before the subsequent cDNA purification (the double-stranded cDNA is ready to be shipped to Gold Labs if necessary).

cDNA SYNTHESIS: Purification of double-stranded cDNA (~ 15...

27

Note

NOTE: Before starting, prepare fresh 70% ethanol in nuclease-free water sufficient for your samples. (500 μl per sample).

Resuspend the AMPure XP beads by vortexing.

- 28 Transfer the sample (\bot 50 μ L) to a clean 1.5ml low DNA binding tube.
- 29 Add A 40 µL of resuspended AMPure XP beads to the reaction and mix by flicking the tube.



30

Incubate on a Hula mixer (rotator mixer) for 00:05:00 at Room temperature

31

Spin down the sample and pellet on the magnet. Keep the tube on the magnet, and using a pipette, discard the supernatant.

- Keep the tube on the magnet and wash the beads with Δ 200 μL of freshly prepared 70% ethanol without disturbing the pellet. Remove the ethanol using a pipette and discard.
- Repeat the previous step X1.
- Spin down and place the tube back on the magnet. Pipette off any residual ethanol. Allow to dry for ~ 30s 00:00:30 , but do not dry the pellet to the point of cracking.
- Remove the tube from the magnetic rack and resuspend the pellet in \mathbb{Z} 13 μL nuclease-free water.
- Incubate on a Hula mixer (rotator mixer) for 00:10:00 at Room temperature .
- 37 Spin down and pellet beads on magnet until the eluate is clear and colorless.
- Remove and retain $\underline{\mathbb{L}}$ 11 μL of eluate into a clean 1.5ml low DNA binding tube.
- Optional: Analyze Δ 1 μL of the purified double-stranded cDNA for quantity using Qubit fluorometer and Qubit 1X dsDNA HS Assay Kit.

40

Subject \blacksquare 10 μ L purified double-stranded cDNA for section SEQUENCING LIBRARY PREPARATION.



Note

STOP POINT: The synthesized double-stranded cDNA can be stored at sequencing.

SEQUENCING LIBRARY PREPARATION

Before starting, prepare fresh 70% ethanol in nuclease-free water sufficient for your samples

10m

1 mL per sample). Program the thermal cycler or use a heat block for 96 well plate: 20 °C for

00:05:00 and 6 65 °C for 00:05:00 . Thaw Ultra II End-prep reaction buffer, NEBNext FFPE

DNA Repair Buffer, Barcode Plate(from SQK-NBD114.96 Kit), and Blunt/TA Ligase Master Mix 6 On ice

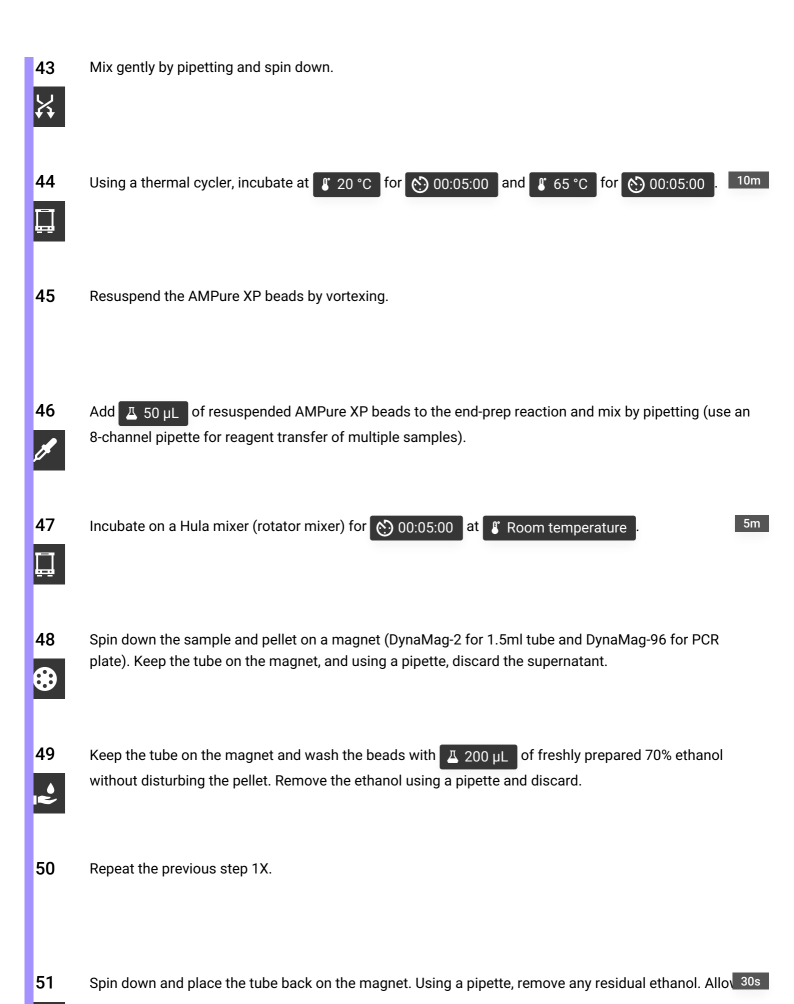
After fully thaw, mix by vortex, spin down briefly, and place 6 On ice . Check that there is no precipitate present (the Blunt/TA Master Mix can sometimes form a precipitate). Spin down Ultra II End-prep enzyme mix and place 6 On ice .

SEQUENCING LIBRARY PREPARATION: End-prep (~ 50 minutes)



Mix the following reagents in a 0.2ml PCR tube. To process 24 samples, prepare a master mix by multiplying gradients except for cDNA by 24 with a 10% overage. Aliquot the master mix into a 96-well plate, then add cDNA or TNA (see Appendix 2 for master mix preparation):

А	В
Component	Volume
DNA/TNA sample	20 μΙ
Nuclease-free water	4 μΙ
Ultra II End-prep reaction buffer	1.75 µl
Ultra II End-prep enzyme mix	1.5 µl
NEBNext FFPE DNA Repair Buffer	1.75 µl
NEBNext FFPE DNA Repair Mix	1 μΙ
Final total volume	30 μΙ





dry for \sim \bigcirc 00:00:30 , but do not dry the pellet to the point of cracking.

52

Remove the tube from the magnetic rack and resuspend the pellet in \square 12 μ L nuclease-free water. Incubate for \bigcirc 00:02:00 at \square Room temperature .



Pellet the beads on a magnet until the eluate is clear and colorless.

54

Remove and retain $\boxed{\text{L}}$ 11 μL of eluate into a clean 1.5ml low DNA binding tube.

SEQUENCING LIBRARY PREPARATION: Barcode ligation (~ 25 m

55

Add the reagents in the order given below, mixing by flicking the tube between each sequential addition:



Note

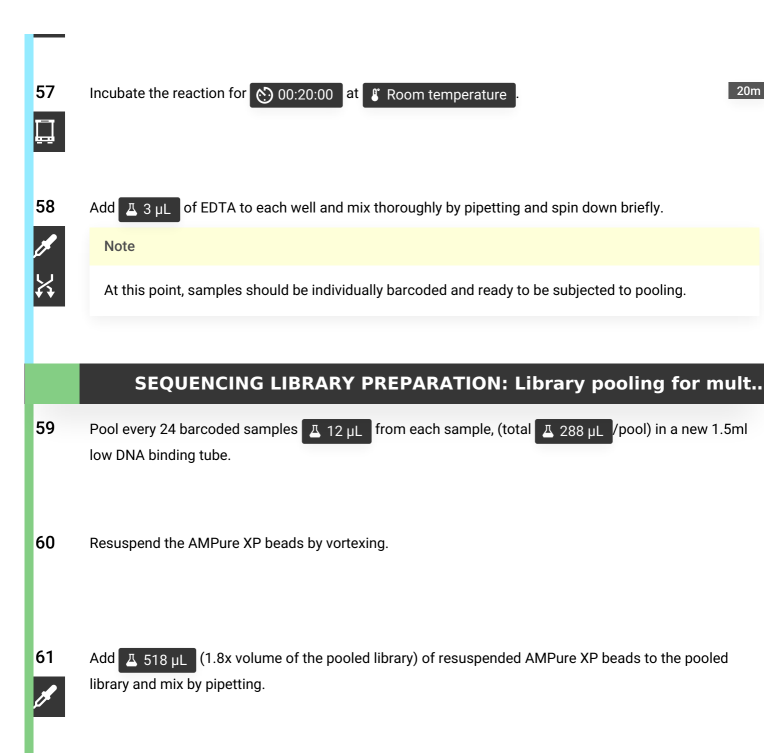
NOTE: When working on 24 End-prepped gDNA/TNA, set up the reactions in a low DNA binding 96-well plate. The Native barcodes can be transferred by an 8-channel pipette directly punching through the sealing foil with tips of the barcode plate. Please reseal the used wells with trimmed adhesive foil. Each well provides sufficient volume for two barcoding ligations.

A	В
Component	Volume
End-prepped DNA	10 μΙ
Native Barcode (pick one form Native Barcoding Expansion 1-96)	2 μΙ
Blunt/TA Ligase Master Mix	12 μΙ
Final total volume	24 μΙ

56

Mix gently by flicking the tube and spin down.





Spin down the sample and pellet on a magnet. Keep the tube on the magnet for 00:05:00, and us 5m

Incubate on a Hula mixer (rotator mixer) for (5) 00:10:00 at

62

63

Room temperature

10m

20m

64 Keep the tube on the magnet and wash the beads with Z 700 µL of freshly prepared 80% ethanol without disturbing the pellet. Remove the ethanol using a pipette and discard. 65 Repeat the previous step 1X. 66 Spin down and place the tube back on the magnet. Using a pipeete, remove any residual ethanol. Allo 30s to dry for $\sim (5)$ 00:00:30 , but do not dry the pellet to the point of cracking. 67 Remove the tube from the magnetic rack and resuspend the pellet in A 35 µL nuclease-free water 10m Incubate for (5) 00:10:00 at (8 37 °C) temperature. 68 Spin down and pellet the beads on a magnet until the eluate is clear and colorless. 69 Remove and retain A 35 uL of eluate into a clean 1.5ml low DNA binding tube.

SEQUENCING LIBRARY PREPARATION: Adapter ligation (\sim 45 m.

- BEFORE STARTING: Thaw Short Fragment Buffer (SFB), Elution Buffer (EB), and NEBNext Quick Ligation Reaction Buffer (5×) at Room temperature, mix by vortexing, spin down, and place On ice.

 Check that the contents or each tube are clear of any precipitate. Spin down the T4 Ligase and the Native Adapter (NA), and place On ice.
 - Taking the pooled and barcoded DNA, perform adapter ligation as follows, mixing by flicking the tube between each sequential addition.

71

A	В
Pooled barcoded sample	30 μΙ
Native Adapter (NA)	5 μΙ
NEBNext Quick Ligation Reaction Buffer (5×)	10 μΙ
Quick T4 DNA Ligase	5 μΙ
Final total volume	50 μΙ

Mix gently by flicking the tube, and spin down.



20m



- **74** Resuspend the AMPure XP beads by vortexing.
- Add Δ 90 μL of resuspended AMPure XP beads to the reaction and mix by pipetting.



76 Incubate on a Hula mixer (rotator mixer) for 🕙 00:10:00 at 🖁 Room temperature .



- 77 Place on the magnetic rack, allow beads to pellet and pipette off supernatant.
- Add Δ 125 μL of the Short Fragment Buffer (SFB) to the beads. Close the tube lid and resuspend the

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0	beads by flicking the tube. Return the tube to the magnetic rack, allow beads to pellet and using a pipette
	discard the supernatant.

- **79** Repeat the previous step X1.
- Spin down and place the tube back on the magnet. Using a pipette, remove any residual supernatant.
- Remove the tube from the magnetic rack and resuspend the pellet in 🔼 13 µL of Elution Buffer (EB).
- Incubate on at 37 °C for 00:10:00 at Room temperature, agitate the sample for 10s e 2 min.
- Pellet beads on magnet until the eluate is clear and colorless.
- Remove and retain \square 13 μ L of eluate into a clean 1.5ml low DNA binding tube.
- Quantify Δ 1 μ L of eluted sample using a Qubit fluorometer and Qubit 1X dsDNA HS Assay Kit (recovery aim ~ Δ 430 ng in total).
- Make up the library to Δ 12 μ L at 10-20 fmol.

Put the library On ice until ready to load or store the library at -20 °C for future sequencing.

Priming and loading the SpotON Flow Cell

88 Check the number of pores in your flow cell.

Note

NOTE: before starting the flow cell pore checking, check the hardware following the manufacturer's guidance.

- **88.1** Turn on GridION (or MinION Mk1C) device. Make sure all the connections for the display, mouse, keyboard, and internet are ready.
- Depending on the number of pooled samples, get one to four new flow cells from the fridge and check the expiration date.
- 88.3 Double-click the MinKNOW icon shown on the desktop to initiate the program.
- 88.4 Use Oxford Nanopore Community username and password to login.
- **88.5** Select the device shown on the screen.
- 88.6 Open the lid of GridION (or MinION Mk1C) and insert the flow cells under the clips, press down the

88.7	The Sequencing Overview tab should show the flow cell not checked in each position in use.
88.8	Navigate to the Start tab and select Flow Cell Check .
88.9	Select the flow cells to assign the flow cell type FLO-MIN114 from the dropdown menu.
38.10	Click Start to begin the flow cell check.
38.11	Record the port number and date of checking on the original package of the flow cell. The flow cell with less than 800 pores should not be used for the sequencing. If the flow cell is not expired, contact Oxford Nanopore Company for customer service.
38.12	If the flow cell is going to be used immediately, keep it on the GridION or MinION Mk1C sequencer for priming. Otherwise put the flow cell back to the original pouch, store at \$\mathbb{g} 4 \cdot \mathbb{C}\$ for next day use. The opened flow cell should be used within one week.
	Priming and loading the SpotON Flow Cell: Flow cell priming
89	BEFORE STARTING: Thaw the Sequencing Buffer (SB), Library Beads (LIB), Flow Cell Tether (FCT) and one tube of Flow Cell Flush (FCF) at Room temperature . Mix SB by tapping or pipetting (DO NOT Vortex) and vortex the
	other tubes. Spin down tubes at 🕴 Room temperature .

flow cell to ensure good thermal and electrical contact.

- 90 Check the air bubble of priming pore.
- 91 Slide open the GridION lid (or MinION Mk1C) and insert flow cell with minimum 800 pores.
- 92 Slide the priming port cover clockwise to open the priming port.

Note

NOTE: Please see **Appendix 1** for the positions of the flow cell ports.

- After opening the priming port, check for a small air bubble under the cover. Draw back a small volume (20-30 µl) to remove any bubbles:
- 93.1 Set a P1000 pipette to 200 μ l. Insert the tip into the priming port. Turn the wheel until the dial shows 220-230 μ l, or until you can see a small buffer volume entering the pipette tip.

Note

- Prepare the flow cell priming mix and prime flow cells.
- 94.1 Prepare flow cell priming mix with components as follows, mix by inverting the tube and pipetting.

A	В
Component	Volume

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A	В
Bovine Serum Albumin (BSA) (50 mg/ml)	5 μΙ
Flow Cell Tether (FCT)	30 μΙ
Flow Cell Flush (FCF)	1170 µl
Final total volume	1205 μΙ

- Load \square 800 μ L of the priming mix into each flow cell via the priming port, avoiding the introduct 5m of air bubbles. Wait for \bigcirc 00:05:00
- **95** Prepare the library for loading.

Note

<u>IMPORTANT</u>: The Library Beads (LIB) tube contains a suspension of beads. These beads settle very quickly. It is vital that they are mixed immediately before use.

- **95.1** Thoroughly mix the contents of the Library Beads (LIB) by pipetting.
- **95.2** In a new tube, prepare each library for loading as follows:

А	В
Component	Volume
Sequencing Buffer (SB)	37.5 μl
Library Beads (LIB)	25.5 μΙ
DNA library	12 μΙ
Final total volume	75 μl

96 Complete the flow cell priming.

96.1 Gently lift the SpotON sample port cover to make the SpotON sample port accessible. 96.2 Load A 200 uL of the priming mix into the flow cell via the priming port (not the SpotON sample port), avoiding the introduction of air bubbles. 97 Loading samples. 97.1 Mix the prepared library gently by pipetting up and down just prior to loading. 97.2 Add A 75 µL 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 drop. 97.3 Gently replace the SpotON sample port cover, making sure the bung enters the SpotON port, close the priming port and replace the GridION lid. Priming and loading the SpotON Flow Cell: Data acquisition and 98 Double-click the MinKNOW icon displayed on the desktop to initiate the program. 99 Use Oxford Nanopore Community username and password to login or continue as Guest.

00 01 Go to the Start tab, and click the Start Sequencing option to choose the running parameters. 101.1 Type in the Experiment Name using the scheme, [YYYY_MM_DD_Approach(gDNA or TNA)_Sample type (soil, water,... etc.)] 101.2 Type in Sample ID (same as experiment name) 101.3 Choose flow cell FLO-MIN114 from the drop-down menu. 02 Click Continue to Kit Selection to move to the next page. 102.1 Click the kit **SQK-NBD114-96** from the Kit Selection menu. 03 Click **Continue to Run Options** to choose run parameters. 2d 103.1 Set run length to 48:00:00 and minimum read length 200 bp. Let adaptive sampling be

Select the device shown on the screen.

	unchecked.
04	Click Continue to Analysis to choose basecalling and Barcoding parameters.
104.1	In the Basecalling options, checkup the basecalling with configuration: High accuracy basecalling.
104.2	In the Barcoding options, turn on the Trim barcodes and Mid-read barcoding filtering.
104.3	Do not turn on the Alignment option.
05	Click Continue to output to the next page.
105.1	Select the output data location, format, and filtering options. Check up the box for Raw reads in POD5 format and Basecalled reads in FASTQ format. Keep the filter score as the system default.
06	Click Continue to final review to proceed.
07	Review the settings listed in the Run Setup page. Correct any errors. Select Start to run the experiment.

The system will automatically navigate the Sequencing Overview when sequencing starts.

48 hrs later, check the sequencing data. Use 1 mL pipette to remove 1 mL waste solution in the waste channel via waste port 1 (see Appendix 1). Remove the flow cells on the device, put it back in the original package, and turn off the device.