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Amplicon sequencing for *Salmonella* Paratyphi A detection

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Typhoid Environmental ...



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We are still developing and optimizing this protocol

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Abstract

This protocol is for direct detection of Salmonella Paratyphi A in wastewater. The wastewater extract can be pre-screened by qPCR for possible presence of Salmonella using triplex qPCR as described in

<https://doi.org/10.1093/infdis/jiad427>



Materials

☒ NEBNext Ultra II End Repair/dA-Tailing Module - 96 rxns **New England Biolabs Catalog #E7546L**

☒ Blunt/TA Ligase Master Mix - 250 rxns **New England Biolabs Catalog #M0367L**

☒ Agencourt AmPure XP beads **Catalog #A63880**

☒ Ultrapure BSA **Ambion Catalog #AM2616**

☒ LongAmp Taq 2X Master Mix - 100 rxns **New England Biolabs Catalog #M0287S** or

☒ LongAmp Taq 2X Master Mix - 500 rxns **New England Biolabs Catalog #M0287L**

☒ Nanopore Flow Cell R10.4.1 **Oxford Nanopore Technologies Catalog #FLO-MIN114**

☒ Native barcoding kit (96) **Oxford Nanopore Technologies Catalog #SQK-NBD114.96**

☒ ONT Native barcoding sequencing kit v14 (24) **Oxford Nanopore Technologies Catalog #SQK-NBD114.24**

☒ Fresh 80% Ethanol

☒ Nuclease-Free Water

☒ Qubit dsDNA Broad Range assay kit (500 assays) **Invitrogen - Thermo Fisher Catalog #Q32853**

☒ Qubit assay tubes **Thermo Fisher Scientific Catalog #Q32856**

☒ Ultrapure BSA **Ambion Catalog #AM2616**

☒ NEBNext Quick Ligation Module - 20 rxns **New England Biolabs Catalog #E6056S** or

☒ NEBNext Quick Ligation Module - 100 rxns **New England Biolabs Catalog #E6056L**

☒ ONT Flow Cell Wash Kit **Oxford Nanopore Technologies Catalog #EXP-WSH004** or EXP-WSH004-XL (48 washes)

Sterile low retention filter tips- for P1000, P200 and P10 pipettes.

200ul strip tubes/96 well plates

1.5 ml microcentrifuge tubes

Primers - high purity

Equipment

- Qubit Fluorometer- Q33238 (Invitrogen)- to quantify DNA and library
- Benchtop vortex - for mixing reagents
- Minifuge - for spinning down reagents
- Hula mixer/rotator- for mixing during the bead clean-up step.
- Plate centrifuge, if you will be using PCR plates instead of strips
- Thermocycler- for PCR and end-prep DNA preparation
- P10 to P1000- single channel pipettes
- P10 and P200- multichannel pipettes
- Laboratory timer
- Benchtop minicooler/ ice bucket
- Magnetic stand for 96 well plate



- Magnetic rack for 1.5 ml tube
- High powered GPU laptop, requirements listed on the ONT website at:
https://community.nanoporetech.com/requirements_documents/minion-it-reqs.pdf

Primer panel

- 1 The following primers have been designed to identify some *S. Paratyphi* A genotypes and chromosomal mutations for AMR. These primers can be ordered lyophilised from your preferred oligo supplier.

1.1 Genotyping panel_version1

	A	B	C	D	E
	Forward primer	Forward sequence	Reverse primer	Reverse sequence	Product size
	2.4.4_v2_F	CCCACTTT CGTGTGAC GGTA	2.4.4_v2_R	ATGATGAA GTCCAGTT GCTG	808
	2.4.5_v2_F	CAGCATTT TCCAGCTC CTCAC	2.4.5_v2_R	ATTTATTGT CGCCCCCG GTT	1348
	1.1_2.4.3_v2_F	AAGTATCC CGGATAAT GCAGC	1.1_2.4.3_v2_R	GGAGAACG TAAATTCG GCGC	1024
	2.4.9_v2_F	AAACGCGC AAAACCTCT CACC	2.4.9_v2_R	CGTTGAGT GGCCTGAA ATGG	1125
	2.4_1_2_v2_F	CATCGATTT GCTGGAGC GTG	2.4_1_2_v2_R	CACTCAGG CGACACAG TCCT	1255
	2.4.3_v2_F	GAGCCGAA ATCAACGT CGCCG	2.4.3_v2_R	GGAACCTCG GCGGTTAT CGAT	1573
	2.1_v2_F	TGGTGCTC ATTGTCTC GCTT	2.1_v2_R	AGACCGCT GCAATCAA AACC	866
	1.2.2_2.4.1_V1_F	CCTTGTTG TGGGTTTC TGCG	1.2.2_2.4.1_V1_R	CCTTTTGG CCGAGATC AAGG	1369
	1.2.2_2.3_V1_F	TCATGCTG AATCACCA CCCGC	1.2.2_2.3_V1_R	GCTAATCT GCCGCAAC CATG	1358
	2.2_1_2_V1_F	AATTGCCA GATGAAGA AGTCG	2.2_1_2_V1_R	CACATCAT CGTGCAGC CCGA	908
	acrB_v2_F	ACACAGGA AGACGACG	acrB_v2_R	AAAGTGCT GGATGAGG	908

A	B	C	D	E
	ATTAGC		TCACG	
2.0.1.1.2.1_V1_F	TCTTCGTA CAACCGCG TCAA	2.0.1.1.2.1_V1_R	TGGCAGGA GTAGACGT ATGC	1579
2.4.2_V1_F	GCAAACCG CAGTCCCA GGAAG	2.4.2_V1_R	AGGCGGTA ATCGAAAG GGCT	510
2.0.1.2.4.6_V1_F	ATGGCGCT GTACCACT CAAA	2.0.1.2.4.6_V1_R	CCTCAGGT GTTTTAGA CCCA	1510
1.2.2.2.4.8_V1_F	ATGGCGGA GGTGTGCA AATA	1.2.2.2.4.8_V1_R	CCGGCGAA AATTGTTG AAGT	1193
1.2.2.3.2_V1_F	AACTGTCG GCATCGGT TCTGT	1.2.2.3.2_V1_R	GGGGTAAC ACAACATC CCTG	1308
2.3.3_V1_F	CCACTGGT GATCTCCG TACCC	2.3.3_V1_R	GTGAAGCC GAAGAAAC TGAA	903
2.3.4_V1_F	CCATGAAC CATGAACC TTGAC	2.3.4_V1_R	TTCCTGCT GCGACCTC TTCA	1209
2.4.7_V1_F	CTACGTTC AAATCGAT GATCG	2.4.7_V1_R	CTGGGGCT ATTTCCGG GCGA	515
2.3.1_V1_F	ACTATTGC CAACCCAA GCCAG	2.3.1_V1_R	CAGTTGGG TGAGCAGG TTCG	1247
gyrA_F	TGACGCCT TCTTCGTA CTCAC	gyrA_R	CTGAAGCT GATCGCCG ATAAAC	2127
parC_v7_F	TGGCACAA TCACTAAA CGCG	parC_v7_R	GCGACGTA CTGGGTAA GTAT	708

2 Primer reconstitution

The primers are received in lyophilized form and need to be reconstituted before use. Prepare a stock solution of 100µM primers.

2.1 To reconstitute the lyophilised primers, use the nmole information on the sheet received with the primers.

Convert the nmol to µmol and then divide by the 100 µmol/L.

For example: for a primer with 24 nmoles, to make 100µM stock solution:



$24 \text{ nmol} / 1000 = 0.024 \text{ } \mu\text{mol}$
 $0.024 \mu\text{mol} / 100 \mu\text{mol/L} = 0.00024 \text{ L}$
 $0.00024 \text{ L} \times 1000 = 0.24 \text{ ml or } 240 \text{ } \mu\text{l}$
So add 240 μl to make a 100 μM solution.

In simple words, multiply the nmol value by 10. Example: for 24 nmol $\times 10 = 240 \text{ } \mu\text{l}$ to make a 100 μM solution.

- 2.2 Add the required volume of nuclease free water to the primer vial, pulse vortex and spin down. This is the primer stock with 100 μM concentration. Store at -20°C for long-term storage.

3 **Primer Dilution**

Using the 100 μM stock, prepare a 10 μM working stock of each primer

- 3.1 In a fresh tube add 5 μl of 100 μM primer stock to 45 μl nuclease-free water to give 50 μl of 10 μM primer.
Store at 4°C for frequent usage or -20°C for long-term storage.

4 **Primer pooling**

Pool the 10 μM concentration of forward and reverse primers listed in 1.1 in equal proportions into a single tube, pulse vortex and spin down. Use this pool for PCR reactions.

5 **PCR reaction**

- 5.1 Thaw the primer pool and LongAmp taq 2x Mastermix on ice.
Pulse vortex and spin down the primer panels, then return to ice.
Do not vortex the LongAmp taq, mix by flicking or pipetting and spinning down before placing back on ice.
- 5.2 Prepare the master mix as follows for the number of samples, plus two controls, and one more extra reaction to account for the pipetting error.

	A	B
	Reagent	Volume for 1 reaction
	2x LongAmp Taq	12.5 μl

	A	B
	Primer pool	2 µl
	Water	5.5 µl
	Total	20 µl

- 5.3 Dispense 20 µl of master mix per reaction into 0.2 ml PCR tubes.
Add 5 µl of sample DNA.
Mix well by pipetting and spin down.

6 PCR Controls

- 6.1 Positive control: ATCC 9150-Salmonella Paratyphi A strain DNA (or use appropriate control strain that is available)
- 6.2 Negative control: 5µl nuclease-free water for a no template control

7 Thermocycler conditions/ program

Set up the thermocycler conditions as follows:

	A	B	C	D
	Step	Temperature	Time	Cycles
	I	94°C	30 seconds	1 cycle
	II	94°C	30 seconds	40 cycles
		58°C	30 seconds	
		65°C	2 minutes 40 seconds	
	III	65°C	10 minutes	1 cycle
	IV	10°C	∞	

8 Store the amplicons at 4°C for a short duration or -20°C for long-term storage if not proceeding with library preparation immediately.

9 **Gel electrophoresis (Optional)**

You may check the amplification of PCR targets by running the PCR products on a 1% agarose gel or on a TapeStation using a D5000 DNA screen tape.

PCR clean-up and quantification

10 PCR clean-up

- 10.1 Prepare the AMPure XP beads for use; resuspend by vortexing.
- 10.2 Add the required volume of resuspended AMPure XP beads to the reaction and mix by pipetting or flicking the tube. The volume is calculated as the 0.8x PCR reaction volume. For example: For 25 ul of PCR reaction = $0.8 \times 25 = 20$ ul of beads.
- 10.3 Incubate on a rotator for 5 minutes at room temperature. If you do not have a rotator, you can flick the tube every 2 minutes to ensure the beads are still in suspension.
- 10.4 Spin down the sample and pellet on a magnet. Keep the tube on the magnet, and pipette off the supernatant.
- 10.5 Keep on the magnet, wash beads with 100 µL of freshly prepared 80% ethanol without disturbing the pellet.
- 10.6 Remove the 80% ethanol using a pipette and discard.
- 10.7 Repeat steps 10.5 & 10.6.
- 10.8 Spin down briefly (~5 seconds) and place the tube back on the magnet.
- 10.9 Allow to air dry for 1-2 minutes.

- 10.10 Remove the tube from the magnetic rack and resuspend the pellet in 20µl of nuclease free water, mix by gently flicking or pipetting.
- 10.11 Incubate for 2 minutes at room temperature.
- 10.12 Pellet beads on magnet until the eluate is clear and colourless.
- 10.13 Still on the magnet, remove 20µl eluate and store it in a clean tube. Avoid disturbing the pelleted beads. If you find the beads keep getting drawn up the pipette tip, try removing 18µl instead.

Primer panel

11 Quantification

The PCR products are quantified using the
ReagentQubit dsDNA BR Assay KitThermo Fisher ScientificCatalog #Q32853
ReagentQubit assay tubesThermo Fisher ScientificCatalog #Q32856
Invitrogen Qubit 4 Fluorometer catalog #Q33238

- 11.1 Standards The Qubit dsDNA kit requires 2 standards for calibration-- Standard #1 and Standard #2.
- 11.2 Label the tube lids. Do not label the side of the tube as this could interfere with the sample reading.
- 11.3 Prepare Qubit working solution for the required number of samples and standards as follows Each sample: Qubit dsDNA BR Reagent 1 µL Qubit dsDNA BR Buffer 199 µL.
- 11.4 Aliquot Qubit working solution to each tube: standard tubes require 190µL of Qubit working solution sample tubes require 198µL of Qubit working solution.
- 11.5 Add 10µL of the standard to the appropriate tube.
- 11.6 Add 2µL of each cleaned PCR product to the appropriate tube. The final volume in each tube must be 200µL once the sample/standard has been added.
- 11.7 Mix each tube vigorously by vortexing for 3–5 seconds.

- 11.8 Allow all tubes to incubate at room temperature for 2 minutes, then proceed to “Read standards and samples.
- 11.9 On the Home screen of the Qubit Fluorometer, press DNA, then select 1X dsDNA broad range as the assay type. The Read standards screen is displayed. Press Read Standards to proceed.
- 11.10 Insert the tube containing Standard #1 into the sample chamber, close the lid, then press Read standard. When the reading is complete (~3 seconds), remove Standard #1.
- 11.11 Insert the tube containing Standard #2 into the sample chamber, close the lid, then press Read standard. When the reading is complete, remove Standard #2.
- 11.12 The instrument displays the results on the Read standard screen. Then press run samples.
- 11.13 On the assay screen, select the sample volume and units: Press the + or – buttons on the wheel, or anywhere on the wheel itself, to select the sample volume added to the assay tube (2µL). From the unit dropdown menu, select the units for the output sample concentration (in this case choose ng/µL).
- 11.14 Insert a sample tube into the sample chamber, close the lid, then press Read tube. When the reading is complete (~3 seconds), remove the sample tube and carefully record the calculated sample concentration.
- 11.15 The top value (in large font) is the calculated concentration of the original sample.

The bottom value is the dilution concentration. For information on interpreting the sample results, refer to the Qubit Fluorometer User Guide.
- 11.16 Repeat step 9.14 until all samples have been read.
- 11.17 All negative controls should ideally be ‘too low’ to read on the Qubit machine, but MUST be < 1ng per ul. If your negative controls >1ng per ul, considerable contamination has occurred and you must redo previous steps.
- 12 **200fmol calculation**
The nanopore protocol recommends using 200fmol of your sample DNA in the first step of library preparation (End-preparation)



- 12.1 Based on the DNA concentration obtained from Qubit, transfer 200fmol of DNA into a fresh tube and add nuclease free water to total 12.5ul. You can use the spreadsheet attached to help calculate the volume required of each sample. We have based the ng required of each sample using the average amplicon length for the primer panel to calculate the ng required for 200fmol product (143 ng for the genotyping panel)



200fmol calculation.xlsx 10.3KB

- 12.2 Use this 12.5ul for the End-preparation reaction.

Preparation for sequencing using ONT Native barcodes

13 End-preparation

- 13.1 From the ONT kit (SQK-NBD114.24 or .96) thaw AMPure XP beads (AXP), mix by vortexing, then keep at room temperature. Thaw the NEBnext Ultra II End Repair reagents on ice, flick or invert the tubes to mix, then spin down.
- 13.2 To the 12.5µl of prepared 200fmol amplicon DNA, add the following:

	A	B	C
	1	Reagent	Volume (µl)
	2	Ultra II End-prep Reaction Buffer	1.75
	3	Ultra II End-prep Enzyme Mix	0.75
	4	Total	2.5

You may make up a master mix of these reagents for the number of samples (+1 for pipetting error) and aliquot 2.5µl for each sample.

- 13.3 Gently mix by pipetting or flicking the tube then spin down (5 seconds).
- 13.4 Incubate in a thermal cycler at 20°C for 5 minutes then 65°C for 5 minutes



13.5 The end prep DNA can be stored at 4°C overnight and protocol can be paused at this point.

14 Native barcode ligation

14.1 From the ONT kit:

Thaw the EDTA at room temperature, mix by vortexing, spin down, then place on ice. Thaw the required Native barcodes at room temperature (a different barcode for each sample), mix by flicking or pipetting, spin down, then place on ice.

Thaw the Blunt/TA ligase master mix at room temperature, mix by inverting and flicking well, spin down, then place on ice.

14.2 Select a different barcode for each sample and note this down in your sample spreadsheet (the same spreadsheet used to calculate sample volumes).

Add the following reagents to a clean 0.2ml tube:

	A	B
	Reagent	Volume
	Native barcode	2.5µl
	Blunt/TA Ligase Master Mix	10µl
	End-prepped DNA	7.5ul

Mix by gently pipetting or flicking then spin down (5 seconds).

14.3 Incubate at room temperature for 20 minutes.

14.4 To each sample add 4µl EDTA (blue cap), mix by pipetting and then spin each tube down.

14.5 Pool all samples into a single 1.5ml tube and note down the final volume.

You can estimate this by multiplying the volume of each sample (22µl if clear cap EDTA, 24µl if blue cap EDTA) by the total number of samples.

- 14.6 Add 0.4x AMPure XP beads to the pooled reaction and mix by flicking the tube.

For example, if the pool volume is 200µl, add 80µl AMPure XP beads.

Incubate at room temperature for 10 minutes, flicking the tube every 2-3 minutes to maintain the bead suspension.

- 14.7 Spin down and pellet the beads on a magnet until the eluate is clear and colourless (3-5 minutes). Whilst still on the magnet remove and discard the supernatant.

- 14.8 Wash the pellet with 700µl 80% ethanol, without disturbing the pellet remove the ethanol and repeat this step.

- 14.9 Spin down (5 seconds), place the tube back on the magnet and remove any residual ethanol. Allow the pellet to air dry for 30 seconds then remove from the magnet and resuspend in 35µl nuclease free water.

- 14.10 Incubate at 37°C for 10 minutes, gently flicking the tube every couple of minutes to encourage elution.

- 14.11 Pellet the beads on the magnet until clear and colourless then take 30µl into a clean 1.5ml tube.

- 14.12 The samples can be stored at 4°C overnight and protocol can be paused at this point.

15 **Adapter ligation**

- 15.1 Thaw NEBnext Quick ligation buffer, Short Fragment Buffer (SFB - ONT kit), and Elution Buffer (EB) at room temperature, vortex to mix, spin down and place on ice.

Flick gently to mix then spin down the NEBNext Quick T4 ligase enzyme and the Native Adapter (NA - from ONT kit) and place on ice.

- 15.2 To the tube with 30µl cleaned barcoded samples, add the following:



	A	B	C
	1	Reagent	Volume (μl)
	2	Native adapter (NA)	5
	3	NEBNext Quick Ligation buffer	10
	4	Quick T4 DNA ligase	5

Gently flick the tube to mix then spin down (5 seconds).

15.3 Incubate at room temperature for 20 minutes

Note: At this point, you can remove your flow cell (FLO-MIN114) from the fridge to allow it to come to room temperature.

15.4 Resuspend AMPure beads by vortexing then add 20μl to the reaction and mix by gently flicking the tube. Incubate at room temperature for 10 minutes, gently flicking the tube every 2-3 minutes.

Note: At this point you can start up MinKNOW and run the Flow Cell check.

15.5 Spin down and pellet on the magnet. Still on the magnet, remove and discard the supernatant.

15.6 Wash the pellet with 125μl Short Fragment Buffer, flick the tube to resuspend the beads, spin down, then place back on the magnet to pellet the beads. Repeat this step.

15.7 Spin down and place back on the magnet then remove any residual buffer. Resuspend in 15μl of Elution Buffer.

15.8 Incubate at 37°C for 10 minutes, flicking every couple of minutes to aid elution.

During this time, thaw Flow Cell Flush (FCF), Flow Cell Tether (FCT), Sequencing Buffer (SB), and Library Beads (LB) at room temperature, mix by vortexing, then spin down and place on ice.



You can also perform the flow cell check. Insert the MinION device into your laptop, open MinkNOW, insert the room temperature flow cell into the MinION, then select Start, then Flow Cell Check.

- 15.9 Spin down then place back on the magnet to pellet the beads. Remove and retain 15µl in a clean 1.5ml tube and keep on ice until ready to load.
- 15.10 Quantify 1 µl of the library using the Qubit fluorometer to ensure the library is not lost during clean-up steps. This value may be useful later for troubleshooting.

16 Priming the flow cell and loading the library

- 16.1 Prepare the flow cell priming mix by adding the following to a clean 1.5ml tube:

	A	B	C
	1	Reagent	Volume (µl)
	2	Flow Cell Flush (FCF)	1170
	3	Bovine Serum Albumin (BSA; 50mg/ml)	5
	4	Flow Cell Tether (FCT)	30

- 16.2 Open the lid of the nanopore sequencing device and slide the flow cell's priming port cover clockwise so that the priming port is visible. After opening the priming port, check for any bubbles under the cover. Draw back a small volume to remove any bubbles (a few µLs). Visually check that there is continuous buffer from the priming port across the sensor array.
- 16.3 Using a P1000 pipette, slowly load 800µL of the priming mix into the flow cell via the priming port. Leave a small amount of liquid in the end of the pipette tip to ensure you do not introduce air into the flowcell. Leave for 5 minutes.



- 16.4 Mix the contents of the LIB tube by pipetting just before adding to the following library mix in a 1.5ml tube:

	A	B	C
	1	Reagent	Volume (μL)
	2	DNA library	12
	3	Sequencing buffer (SB)	37.5
	4	Library beads (LIB)	25.5

- 16.5 Complete the flowcell priming by opening the SpotOn port cover and carefully loading 200μL of the priming mix into the priming port. As before, leave a small amount of liquid in the bottom of the tip to avoid the introduction of air bubbles. When adding the priming mix, you may see a small amount of liquid come up through the SpotOn port. If you do, pause and allow the liquid to flow back into the flowcell before continuing putting through the priming mix.
- 16.6 Mix the prepared library mix gently by pipetting. Add the library mix to the flowcell via the SpotOn port in a dropwise fashion, allowing each drop to flow into the flowcell before adding the next.
- 16.7 Replace the SpotOn port cover and close the priming port, then replace the lid of your sequencing device

17 Starting the Sequencing Run

Open the ONT MinKNOW software and follow the steps below to set up and start your sequencing run.

- 17.1 Click start, then start sequencing. Create a name for you sequencing run, it is good practise to make this unique and identifiable for if you ever need to revisit the data. The date and an experiment name are recommended. In sample name you can put a number or repeat the experiment name - this is not as important as the run name. Then click continue.
- 17.2 Select the kit used - this is SQK-NBD114.24 or SQK-NBD114.96 depending on whether you have purchased the 24 or 96 barcode kit. Click continue.
- 17.3 In the run length options, set the run time; refer to the attached guide for calculating the sequencing time. Click continue.



Calculating the sequencing time an... 17KB

17.4 In the basecalling options, select high accuracy basecalling. In the barcoding options, make sure barcoding is enabled and toggle to use "barcode at both ends" and "trim barcodes". Click continue until you reach the run overview, where you can double check the selected options, then click start run.

18 **Washing the flow-cell after sequencing**

A Nanopore flow cell can be used multiple times, so it must be washed to remove the library from the previous run.

18.1 Place the tube of Wash Mix (WMX) on ice. Do not vortex the tube and thaw one tube of Wash Diluent (DIL) at room temperature.

18.2 In a clean 1.5 ml Eppendorf DNA LoBind tube, prepare the following Flow Cell Wash Mix: Mix well by pipetting, and place on ice. Do not vortex the tube.

18.3 After the sequencing run, close the ONT MinKNOW software. Detach the MinION device from the laptop and leave the flowcell in the device.

18.4 Before removing the waste fluid, ensure that the flow cell priming port cover and SpotON sample port cover are closed.

18.5 Using a P1000, remove all fluid from the waste channel through waste port 1. As both the flow cell priming port and SpotON sample port are closed, no fluid should leave the sensor array area.

18.6 Rotate the flow cell priming port cover clockwise so that the priming port is visible.

18.7 After opening the priming port, check for a small air bubble under the cover. Draw back a small volume to remove any bubbles.

18.8 Load 200 µl of the prepared Flow Cell Wash Mix into the flow cell priming port, avoiding the introduction of air and incubate for 5 minutes.

18.9 Load the remaining 200 µl of Flow Cell Wash Mix into the flow cell priming port.

18.10 Close the flow cell priming port and incubate for 60 minutes.

- 18.11 Before removing the waste fluid a second time, ensure that the flow cell priming port cover and SpotON sample port cover are closed.
- 18.12 Using a P1000, remove all fluid from the waste channel through waste port 1. As both the flow cell priming port and SpotON sample port are closed, no fluid should leave the sensor array area.
- 18.13 You can now either run a second library on the flow cell straight away or store the flow cell for later use.
- 18.14 To run a second library straight away, follow the flow cell priming instructions in step 16.
- 18.15 To store the flowcell for later use, thaw one tube of Storage Buffer (S) at room temperature, mix contents thoroughly by pipetting and spin down briefly (~30 seconds).
- 18.16 Rotate the flow cell priming port cover clockwise so that the priming port is visible.
- 18.17 After opening the priming port, check for a small air bubble under the cover. Draw back a small volume to remove any bubbles.
- 18.18 Slowly add 500 μ l of Storage Buffer (S) through the flow cell priming port.
- 18.19 Close the priming port.
- 18.20 Using a P1000, remove all fluid from the waste channel through waste port 1. As both the flow cell priming port and SpotON sample port are closed, no fluid should leave the sensor array area.
- 18.21 The flow cell can now be stored at 4-8°C.