

## MiSeq Dual Index Amplicon Sequencing Sample Prep

- 1) Perform the first PCR using the primer constructs  
Illumina adapter-N<sub>4</sub>-*forward primer*  
ACACTCTTTCCCTACACGACGCTCTTCCGATCT-*fwd\_primer*  
and Illumina adapter-reverse primer  
AGACGTGTGCTCTTCCGATCT-*rev\_primer*

Optimization of the protocol may be required, but a starting point is 2.5 µL of each primer (10 µM), 1-10 ng of template DNA, 25 µL of Phusion Mastermix (ThermoScientific) and dH<sub>2</sub>O to a total volume of 50 µL.

Cycling conditions are:

|               |      |       |       |
|---------------|------|-------|-------|
| Denaturation  | 98°C | 30"   |       |
| Denaturation  | 98°C | 10"   | \     |
| Hybridization | ***  | 30"   | - 20x |
| Elongation    | 72°C | ***   | /     |
| Elongation    | 72°C | 2 min |       |

Hybridization conditions are primer-dependent, but should be similar to those used for the primers without adapters. For elongation time, 15 s/kb should suffice.

Kapa HiFi polymerase (Kapa Biosystems) can be considered with the template is particularly refractory to amplification, following the instructions from the manufacturer.

2) A cleaning step must be performed to eliminate loose primers and eventual primer dimers or low molecular weight unspecific products. This can be achieved with magnetic beads, a spin column or a gel. If carrying out this procedure in SciLifeLab Stockholm, a CA-cleaning in the MBS is recommended. See "CA cleaning" protocol.

3) A second PCR is conducted for attaching standard Illumina handles and index primers.

Multiplex\_fwd

AATGATACGGCGACCACCGAGA{TCTACAC}-[i5 index]-ACACTCTTTCCCTACACGACG

Multiplex\_rev

CAAGCAGAAGACGGCATACGAGAT-[i7 index]-

GTGACTGGAGTTCAGACGTGTGCTCTTCCGATCT

The curly brackets denote bases that will not be read in the sequencer. See notes at the bottom for multiplexing indexes combination guidelines.

Each well should contain 23 µL of the clean PCR product above, 25 µL of Phusion Mastermix and 1.5 µL of each primer (10 µM).

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Cycling conditions are as follows:

|               |      |       |   |
|---------------|------|-------|---|
| Denaturation  | 98°C | 30"   | \ |
| Denaturation  | 98°C | 10"   |   |
| Hybridization | 62°C | 30"   |   |
| Elongation    | 72°C | ***   | / |
| Elongation    | 72°C | 2 min |   |

Add one second to the elongation time used in step 1.

- 4) Repeat step (2), twice if necessary. If using magnetic beads, adapt the bead amount for the larger amount of DNA now present. Beads will interfere with the downstream steps, so remove them first by placing the plate in a magnetic stand, leaving it undisturbed for at least 3 minutes and then transferring the reaction product to a fresh plate.
- 5) Quantify sample concentration and length. For concentration, a fluorometric measurement such as Qubit (Invitrogen) or PicoGreen (Invitrogen) should be preferred over an auto-fluorescence method such as NanoDrop (Thermo Scientific).
- 6) Measure the average fragment length. BioAnalyzer (Agilent) HS is recommended, but a simple agarose gel may suffice if you have enough material.
- 7) Pool all samples together for a total volume of at least 15 µL and a concentration of 2-10 nM / barcoded template. The molarity of a sample can be calculated using the formula:  
$$\text{Concentration} \times 10^6 / 656.6 \times \text{Length}$$
- 8) Samples are now ready to be handled by facility personnel. Adding 5% of random DNA, such as PhiX, is recommended for improving amplicon sequencing in Illumina instruments.

## INDEXES

| i7   |          | i5                                                                                                                        |          |
|------|----------|---------------------------------------------------------------------------------------------------------------------------|----------|
| N701 | TAAGGCGA | N501                                                                                                                      | TAGATCGC |
| N702 | CGTACTAG | N502                                                                                                                      | CTCTCTAT |
| N703 | AGGCAGAA | N503                                                                                                                      | TATCCTCT |
| N704 | TCCTGAGC | N504                                                                                                                      | AGAGTAGA |
| N705 | GGACTCCT | N505                                                                                                                      | GTAAGGAG |
| N706 | TAGGCATG | N506                                                                                                                      | ACTGCATA |
| N707 | CTCTCTAC | N507                                                                                                                      | AAGGAGTA |
| N708 | CAGAGAGG | N508                                                                                                                      | CTAAGCCT |
| N709 | GCTACGCT | <b>NB!</b> the i7 barcodes should be reverse complemented in the primers you order, to be read correctly upon sequencing! |          |
| N710 | CGAGGCTG |                                                                                                                           |          |
| N711 | AAGAGGCA |                                                                                                                           |          |
| N712 | GTAGAGGA |                                                                                                                           |          |

These are the standard Nextera indexes. To make sure that the sequencer detects signal in multiple channels, combine them as follows:

| Plex   | i7                                                                                   | i5                                                                                                         |
|--------|--------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------|
| 1      | Any                                                                                  | Any                                                                                                        |
| 2      | [option1] N701 + N702<br>[option2] N702 + N704                                       |                                                                                                            |
| 3      | [option1] N701 + N702 + N704<br>[option2] N703 + N705 + N706                         |                                                                                                            |
| 4 to 5 | [option1] N701 + N702 + N704 + any other<br>[option2] N703 + N705 + N706 + any other |                                                                                                            |
| 6      | N701 + N702 + N704 + N705 + N706                                                     |                                                                                                            |
| 7-12   | [option1] N701 + N702 + N704 + any other<br>[option2] N703 + N705 + N706 + any other | [option1] N501 + N502<br>[option2] N503 + N504<br>[option3] N505+ N506                                     |
| > 12   | N701 + N702 + N704 + N703 + N705 + N706 + any other                                  | [option1] N501 + N502 + any other<br>[option2] N503 + N504 + any other<br>[option3] N505+ N506 + any other |

If doing twelve samples or less, consider also using the single index protocol.