



### **VERSION 2**

APR 13, 2023

( HTTM: Illumina library preparation V.2

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#### **ABSTRACT**

Part of the HTTM protocol dedicated to the preparation of Illumina sequencing libraries.

# OPEN ACCESS

dx.doi.org/10.17504/protocol s.io.n2bvj8oowgk5/v2

#### **External link:**

https://doi.org/10.1371/journa l.pone.0283990

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#### **MANUSCRIPT CITATION:**

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

**MATERIALS** 

# **Preparation of Nextera Adaptaters:**

**Created:** Aug 24, 2022

Nextera (NxT) adapters are prepared by hybridisation of the following primers:

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	A	В
Nxt-XTv2- B-N701-T CAAGCAGAAGACGGCATACGAGATTCG GAGATGTGTATAAGAGACAGT		CAAGCAGAAGACGGCATACGAGATTCGCCTTAGTCTCGTGGGCTCG GAGATGTGTATAAGAGACAGT
	Nxt-XTv2- B-3R-ac3- 5phos	/5Phos/CTGTCTCTATACACATCTCCGAGCCCACGAGAC/3InvdT/

- Preparation of the 5X annealing buffer (5X Tris NaCl buffer : 50 mM Tris, pH 7.5-8, 250 mM NaCl) :
- 500 µl Tris-HCl 1M pH 7.5
- 500 µl NaCl 5M
- 9 ml H20 mol.-grade
- Preparation of the adapters (40  $\mu$ M 50  $\mu$ L) :
- Resuspend both primers in water to obtain 100 μM stocks
- Mix 20 μl of each (Nxt-XTv2-B-N7XX-T and Nxt-XTv2-B-3R-ac3-phos5')
- Add 10 µl of 5X annealing buffer
- Annealing reaction in a thermocycler (decrease temperature from 98 to 4C (-0.1C/cycle(10s/cycle)))

#### Primers used for the first PCR:

A	В
Nxt_A	AATGATACGGCGACCACCGAGATCTACAC
Nxt_B CAAGCAGAAGACGGCATACGAGAT	

## Primers template for barcoding PCR:

	A	В
Nxt_i5_barco ding AATGATACGGCGACCACCGAGATCTACAC [8 Nu Index] TCGTCGGCAGCGTCAGATGTGTA  Nxt_i7_barco ding CAAGCAGAAGACGGCATACGAGAT [8 Nu Index] GTCTCGTGGGCTCGGAGATGTGTATAAG		AATGATACGGCGACCACCGAGATCTACAC [8 Nu Index] TCGTCGGCAGCGTCAGATGTGTA
		CAAGCAGAAGACGGCATACGAGAT [8 Nu Index] GTCTCGTGGGCTCGGAGATGTGTATAAG

### BEFORE START INSTRUCTIONS

All steps and master mixes need to be kept on ice as much as possible. Thermocyclers need to be cooled at 4C before inserting sample plate.

# Libraries

1h 34m

- 1 Transfer 2.5µl of DNA from the DNA extraction plate to a new PCR plate.
- 2 Prepare a fragmentation master mix with:

A	В
NEB Ultra II FS buffer	77 µl
NEB Ultra II FS enzyme	22 µl
Molecular grade water	11 µl

- 3 Add  $\underline{\mathsf{A}}$  1  $\mu L$  of the fragmentation master mix to each well.
- 4 Incubate in a thermocycler with the following protocol:

45m

- 00:15:00 at \$\ \$\ 37 \cdot \cdot
- 00:30:00 at \$\cdot 65 \cdot C
- Add  $\triangle$  1  $\mu$ L of 4 $\mu$ M Nextera (NxT) adaptors to each well.
- 6 Prepare a ligation master mix with :



A	В
NEB Ultra II ligation master mix	377.4 µl
NEB Ultra II ligation enhancer	12.1 µl

- 7 Add  $\underline{A}$  3.5  $\mu L$  of ligation master mix to each well.
- 8 Incubate in a thermocycler with the following protocol:

•	<b>©</b> 00:30:00	at	<b>₿</b> 20 °C
•	<b>(:)</b> 00:10:00	at	<b>\$</b> 65 °C

9 Prepare a PCR master mix with:

A	В
NxT_A primer 20 μM	880 µl
Nxt_B primer 20 µM	880 µl
Molecular grade water	8360 µl
PCR Mix 2X	11000 µl

- 10 Add  $\perp$  192  $\mu$ L of PCR master mix to each well.
- 11 Split the PCR reaction into 4 different plates (50µl per plate).
- 12 Incubate each plate in a thermocycler with the following cycles :

■ 👏 00:00:30 at 🖔 98 °C

■ 00:00:15 at \$\ 98 °C

■ 00:00:30 at \$ 72°C

■ Repeat from step 2 for 20~25 cycles\*

40m

3m 15s

■ 00:02:00 **\$** 72 °C

- Pool the 4 PCR replicates together in a.
- 14 Transfer  $\mathbb{Z}_{2\mu L}$  of DNA from the pool plate to a new PCR plate.
- 15 Add  $\underline{A}$  2  $\mu L$  of each barcoding primer to the DNA :
  - Nxt\_i5\_barcoding
  - Nxt\_i7\_barcoding
- 16 Prepare a PCR master mix with:

Α	В
Molecular grade water	2090 μΙ
PCR mix 2X	2750 μΙ

- 17 Add  $\underline{\mathbb{Z}}$  44  $\mu L$  of the PCR master mix to each well of the plate.
- 18 Incubate in a thermocycler with the following protocol:

3m 45s

- Ø 00:00:30 at \$ 98 °C
- 🕙 00:00:15 at ឺ 98 °C
- Repeat from step 2 for 5 cycles
- **③** 00:02:00 at **⑤** 72 °C

- 19 Pool together  $\angle 2 \mu L$  of each sample.
- 20 Purify with SPRI beads using a 0.8 ratio. Resuspend with  $\pm$  50  $\mu L$  of molecular grade water.
- 21 Proceed with QC and sequencing.