# Metagenomic library preparation

#### DNA Library Prep - Illumina DNA Prep Kit

Target insert size: ~150 bp

Final library size:  $\sim 600$  bp (for  $2 \times 150$  bp reads)

DNA input: 150 ng per sample

#### **DNA Quantification & Preparation**

• Quantify DNA using Qubit dsDNA HS Assay

• Check purity: A260/280 = 1.8-2.0

• Dilute 150 ng DNA in 30 μL 10 mM Tris-HCl (pH 7.5–8.5)

## **Tagmentation**

- Prepare Tagmentation Master Mix:
  - 11 μL BLT (bead-linked transposomes)
  - 11 µL TB1 (tagmentation buffer)
- Add 20 μL master mix to 30 μL DNA
- Mix well, seal plate
- $\bullet\,$  Thermocycle: 55°C for 15 min, hold at 10°C

#### **Post-Tagmentation Cleanup**

- Add 10 µL TSB (Stop Buffer), pipette to mix
- Thermocycle: 37°C for 15 min, hold at 10°C
- Place on magnetic stand, discard supernatant
- Wash beads twice with 100  $\mu L$  TWB
- Leave final wash in wells (do not dry)

### PCR Amplification (5 Cycles)

- Prepare PCR Master Mix per sample:
  - $-22 \mu L EPM$
  - 22 μL Nuclease-free water
- Add 40 µL master mix to beads
- Add 10 µL pre-paired index adapters (i5+i7)
- Thermocycle:
  - -68°C 3 min
  - 98°C 3 min
  - 5 cycles: 98°C 45s, 62°C 30s, 68°C 2 min
  - 68°C 1 min  $\rightarrow$  Hold at 10°C

## Library Cleanup (Ampure beads)

- Transfer 45  $\mu$ L PCR product to new plate
- Add 40  $\mu$ L water + 45  $\mu$ L SPB  $\rightarrow$  mix, incubate 5 min
- Transfer 125  $\mu L$  to new plate with 15  $\mu L$  SPB  $\rightarrow$  mix, incubate 5 min
- Wash twice with 200 µL 80% EtOH
- Air dry beads for 5 min
- Elute in 32 μL RSB, transfer 30 μL clean library

#### Library QC & Pooling

- Tapestation for analyzing the size of library
- Qubit for quantification of the final pooled library.