

## ***In Vitro* protocol in brief test LymphoTranscript**

### **1. Reverse transcription**

#### Reagents:

- Kit M-MLV RT
- dNTPs (10mM)
- Hexamers (100 $\mu$ M)
- ARN to be tested

#### Steps :

- Thaw the reagents
- Prepare reverse transcription mix
  - RT buffer (1,25  $\mu$ L)
  - DTT (0,5  $\mu$ L)
  - dNTPs (1  $\mu$ L)
  - Hexamers (1  $\mu$ L)
- Deliver 3,75  $\mu$ L mix per tube
- Add 2  $\mu$ L of RNA sample
- Vortex and centrifuge
- Put samples in thermocycler and start program 1 step 1a
- Add 0,5  $\mu$ L of M-MLV RT
- Centrifuge
- Put samples in thermocycler and start program 1 step 1b
- Once the program is over and the block cool down at 4°C, put out tubes
- Put tubes on ice or cooling block



Then proceed to step 2 or store cDNA products between -30°C et -15°C.

### **2. Hybridization of probes**

#### Reagents:

- SALSA MLPA Buffer
- GEP- LTPM** probe Mix

- Thaw reagents
- Prepare Hybridization mix
  - Salsa MLPA Buffer (1,5  $\mu$ L)
  - GEP-LTPM** Probe Mix (1,5  $\mu$ L)
- Vortex and centrifuge
- Deliver 3  $\mu$ L of mix per cDNA tubes
- Centrifuge
- Put samples in thermocycler and start program 1 step 2

### 3. Ligation

#### Reagents:

- SALSA Ligase Buffer A
- SALSA Ligase Buffer B
- SALSA Ligase 65
- Nuclease free water

- Thaw reagents
- Prepare ligation mix
  - Nuclease free water (25  $\mu$ L)
  - SALSA Ligase Buffer A (3  $\mu$ L)
  - SALSA Ligase Buffer B (3  $\mu$ L)
- Vortex and centrifuge
  - Salsa Ligase 65 (1  $\mu$ L)
- After 1h of hybridization, deliver 32  $\mu$ L of mix per cDNA tubes
- Start program 1 étape 3



Then proceed directly to step 4 or store the ligation products between -30°C and -15°C.



After this step, do not store the products at high temperature (e.g. 4°C or at room temperature) in order to avoid non-specific ligations which could result from a residual activity of the enzyme.

### 4. Amplification and incorporation of barcodes and adaptators

#### Reagents:

- GEP-BC-xxx**  
Barcodes
- GEP-BCC-xxx**  
GAPDH barcodes
- Q5 MasterMix
- Nuclease free water

- Thaw the reagents
- Prepare PCR mix
  - Q5 MasterMix (12,5  $\mu$ L)
  - Nuclease free water (5,5  $\mu$ L)
- Vortex and centrifuge
- Deliver 18  $\mu$ L of mix per wells of PCR plate
- Add 5  $\mu$ L of ligation product in each well
- Add 2  $\mu$ L of barcode (**GEP-BC-xxx** or **GEP-BCC-xxx** according to the test) in each well



Use different barcodes BEP-BC-xxx for each tested samples.

- Put samples in thermocycler and start program 2



Then proceed directly to step 5 or store the PCR products between -30°C and -15°C.



Do not store these products for prolonged periods at high temperature (e.g. 4°C in the thermocycler or at room temperature).

## 5. Purification and dosage sequencing librairies

### Reagents:

- 100% Ethanol
- Nuclease free water
- AMPure XP Magnetic beads
- TE Buffer
- Qubit dsDNA HS Assay

### Steps :



**Ensure beads are completely re-suspended before use.**

- Purify 25  $\mu$ L products with 45  $\mu$ L AMPure XP beads
- Elute PCR products in 50  $\mu$ L of TE buffer



**After purification, librairies can be stored between -30°C et -15°C before sequencing.**

- Dose 10  $\mu$ L of each library by fluorimetry



**This rapid protocol completes the notice. It does not dispense with complete reading of the notice.**