nCounter[®] miRNA Sample Prep



MicroRNA Sample Preparation for 12 Samples

n Program thermocycler protocols.

Annealing Protocol		Ligation Prote	Ligation Protocol	
Temperature	Time	Temperature	Time	Temperature
94°C	1 min	48°C	3 min	37°C
65°C	2 min	47°C	3 min	70°C
45°C	10 min	46°C	3 min	4°C
48°C	hold	45°C	5 min	Total Time
Total Time	13 min	65°C	10 min	
		4°C	hold	
		Total Time	24 min	

urification	Drotocol
urnication	Protocor

Temperature	Time
37°C	1 hr
70°C	10 min
4°C	hold
Total Time	1 hr 10 min

Prepare total RNA samples.

Using RNAse-free water, normalize total RNA samples to 33ng/uL in a total of 3uL to provide 100ng input (there is no need to enrich for small RNAs). Samples must be free of chaotropic salts and organic solvents.

3 Prepare controls.

Add 1µL of miRNA Assay Controls to 499µL of RNAse-free water in a sterile microfuge tube. Vortex and briefly spin down. Store on ice.

Anneal samples.

- Combine 13µL of Annealing Buffer, 26µL of nCounter miRNA Tag Reagent, and 6.5µL of the miRNA Assay Controls dilution prepared in Step 3 to create an annealing mastermix. Mix well by pipetting.
- Dispense 3.5µL of the annealing mastermix into provided 12 x 0.2mL strip tubes.
- Add 3µL of total RNA sample (100ng) into each tube with mastermix. Cap tube, flick to mix and spin down.
- Place strip in thermocycler and initiate Annealing Protocol.

6 Ligate samples.

- Combine 19.5µL of PEG and 13µL of Ligation Buffer in a microfuge tube and mix well by pipetting to prepare a ligation mastermix. PEG should be pipetted very slowly to ensure an accurate measurement.
- When the thermocycler has reached 48°C, remove tubes, add 2.5µL of the ligation mastermix to each tube in the strip. Cap tubes, flick to mix and spin down. Do not turn off thermocycler.
- Incubate tubes at 48°C in the thermocycler for 5-mins.
- While tubes remain in thermocycler, carefully uncap strips, add 1µL of ligase to each tube. Check the tip at each pipetting step to ensure all ligase was dispensed. There is no need to mix. To keep track of ligase addition, it can be helpful to line up 12 tips in front of the thermocycler discarding each tip after use.
- Immediately recap tubes in thermocycler, initiate thermocycler Ligation Protocol.

6 Clean up ligation.

- Remove tubes from thermocycler, carefully uncap strips, add 1µL of Ligation Clean-Up Enzyme to each reaction. Flick to mix and spin down.
- Place tubes in thermocycler and initiate Purification Protocol.
- Add 40µL of RNAse-free water. Samples may be stored at -20°C for up to several weeks before hybridization.

For more comprehensive information, sign in to our customer resources site (www.nanostring.com/sign/) and go to Support > Customer Resources* to view the manuals and other technical product literature. For technical support, please e-mail support@nanostring.com or in the U.S./Canada, call 1888 358 6266.

© 2013 NanoString Technologies, Inc. All rights reserved.

LBL-C0106-04