

nCounter[®] miRNA Sample Prep



MicroRNA Sample Preparation for 12 Samples

1 Program thermocycler protocols.

Annealing Protocol

Temperature	Time
94°C	1 min
65°C	2 min
45°C	10 min
48°C	hold
Total Time	13 min

Ligation Protocol

Temperature	Time
48°C	3 min
47°C	3 min
46°C	3 min
45°C	5 min
65°C	10 min
4°C	hold
Total Time	24 min

Purification Protocol

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

2 Prepare total RNA samples.

Using RNase-free water, normalize total RNA samples to 33ng/μL in a total of 3μL 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.

4 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**.

5 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 **1 888 358 6266**.