**CodeSet RNA:Protein Hybridization Setup**

Prepared samples are used as input into nCounter hybridization reactions containing NanoString Reporter and Capture probes. These overnight hybridization reactions enable specific hybridization of reporter of capture probes to their target. After hybridization, reactions are purified and imaged on nCounter® systems.

This manual provides instructions for producing hybridization Master Mix and setting up nCounter hybridizations using CodeSet chemistry.

**Materials and Reagents**

**Table 1. CodeSet RNA:Protein Hybridization Reagents**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Description</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reporter CodeSet</td>
<td>Barcoded Reporter Probes</td>
<td>At or below -80°C</td>
</tr>
<tr>
<td>Capture ProbeSet</td>
<td>Capture Probes</td>
<td>At or below -80°C</td>
</tr>
<tr>
<td>Hybridization Buffer</td>
<td>Supplied with nCounter Master Kits and SPRINT Reagent Packs</td>
<td>RT (15–25°C)</td>
</tr>
<tr>
<td>Protein TagSet (R)</td>
<td>Barcoded Capture and Reporter Probes for Protein Tags</td>
<td>At or below -80°C</td>
</tr>
</tbody>
</table>

**Important Probe Handling Instructions**

- During setup, do not vortex or pipette vigorously to mix.
- Mixing should be done by flicking or inverting the tubes.
- If using a microfuge to spin down tubes, do not spin any faster than 1,000 RCF for more than 30 seconds.
- Do not “pulse” to spin because that will cause the centrifuge to go to the maximum speed and may spin the probes out of solution.

**Nucleic Acid Sample Input Recommendation**

**Table 2. CodeSet RNA:Protein Hybridization Reagents**

<table>
<thead>
<tr>
<th>Analyte</th>
<th>MAX/FLEX</th>
<th>SPRINT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unamplified Total RNA</td>
<td>Fresh Frozen</td>
<td>100 ng (~33 ng/µL)</td>
</tr>
<tr>
<td>FFPE</td>
<td>300 ng (~100 ng/µL)</td>
<td>150 ng (~50 ng/µL)</td>
</tr>
</tbody>
</table>

For additional information related to nucleic acid sample preparation based on sample type, see Preparing Nucleic Acid from FFPE Samples (MAN-10050) or Preparing Nucleic Acid from Fresh Frozen Samples (MAN-10051).
CodeSet RNA:Protein Hybridization Setup

**IMPORTANT:** Check the reagent labels before you begin to ensure the correct reagents are being utilized.

1. Pre-heat a thermal cycler to **65°C** with a heated lid at **70°C**.
   
   **NOTE:** A thermal cycler with a heated lid is required for this protocol. NanoString recommends a thermal cycler with a programmable heated lid. Models without programmable lids may reach a high temperature that causes tubes to melt or deform during extended or overnight hybridization times, and if used, should be set to ensure that the heated lid does not exceed 110°C.

2. Remove Reporter CodeSet, Capture ProbeSet, and Protein Plus tubes from the freezer and thaw at room temperature. Invert or flick the tubes several times to mix well and briefly spin down reagents.

3. Create a hybridization master mix by adding the following reagents to the tube containing the appropriate Reporter CodeSet.
   
   **NOTE:** Do not add the Capture ProbeSet to the master mix.

<table>
<thead>
<tr>
<th>Table 3. CodeSet Master Mix for one nCounter assay</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Component</strong></td>
</tr>
<tr>
<td>Reporter CodeSet</td>
</tr>
<tr>
<td>Hybridization Buffer</td>
</tr>
<tr>
<td>Protein Plus</td>
</tr>
<tr>
<td><strong>Total Volume</strong></td>
</tr>
</tbody>
</table>

4. Flick or invert the tubes repeatedly to mix then briefly spin down the master mixes.

5. Label a 12-tube PCR hybridization strip. If necessary, ensure the strip will fit in a microfuge or picofuge by cutting both the strip tube and its lid in half prior to setting up the reactions, taking care not to crack the tubes.

6. Dilute (if necessary) and denature prepared protein lysate (from Protein Processing for Lysate Samples, MAN-10054):
   
   a. For protein samples from FFPE, dilute the lysate 1:5 in Buffer T.
   
   **NOTE:** This dilution is necessary to prevent saturation of the nCounter assay cartridge. The optimal dilution may vary depending on tissue type and size. Save the remaining undiluted sample at -80°C to re-run the assay if adjustments are to be made.

   b. Denature at 95°C for 5 min in a thermocycler with a heated lid at 100°C.

   c. Snap cool on ice, ice water, or quickly ramp the thermocycler to 4°C.

   d. If using nCounter SPRINT system, further dilute the denatured sample with an equal amount of nuclease-free water. A 1 µL aliquot of this sample is required per hybridization reaction. Additional dilution is not required with the MAX or FLEX analysis systems.

   e. Spin briefly, vortex, and spin briefly again to ensure a homogenous mixture.
7. Prepare hybridization reactions (16 µL total final volume per reaction):
   a. Add 10 µL of master mix to each well of a strip tube. Use a fresh tip for each pipetting step.
   b. Add 3 µL of the RNA sample to each tube (see Table 2).
   c. Add 1 µL of the **denatured diluted protein lysate** to each tube.
   d. Invert the Capture ProbeSet tube to mix and spin down the contents.
   e. Add 2 µL of Capture ProbeSet to each tube immediately before placing at 65°C.
   f. Cap the strip tubes tightly and mix the by inverting the tubes several times and flicking to ensure complete mixing.
   g. Spin briefly and immediately place the tubes in a pre-heated 65°C thermal cycler.

8. Incubate hybridization reactions for at least 16 hours. Maximum hybridization time should not exceed 48 hours.

9. Ramp reactions down to 4°C and process the following day. Do not leave the reactions at 4°C for more than 24 hours or increased background may result.
   
   **NOTE:** Selecting a fixed hybridization time followed by a ramp down to 4°C ensures equivalent hybridization time for all assays being directly compared in the same series of experiments. Counts continue to accumulate with time at 65°C, with total counts typically increasing 5% per hour between 16 and 24 hours. Although a 16-hour incubation is adequate for most purposes, a longer incubation increases sensitivity by increasing counts without significantly increasing background.

10. Once the hybridization reactions have been removed from the thermal cycler, proceed immediately to an nCounter Prep Station or SPRINT as described in the nCounter Analysis System User Manual (MAN-C0035) or nCounter SPRINT User Manual (MAN-10017). If using this protocol in conjunction with other 3D compatible products, see the next section for 3D post-hybridization processing.
3D Post-Hybridization Processing

**IMPORTANT:** Use only Master Kit strip tubes on the Prep Station. If you need additional strip tubes, contact NanoString.

1. After the parallel hybridization reactions are completed, remove the strip tubes from the thermal cycler and spin briefly.

2. Using a multichannel pipette, pool the full volume (16 μL) of the RNA:Protein Hybridization strip tube reactions with the (sample-matched) SNV strip tube reactions. Pooled sample will be processed and analyzed in a single nCounter cartridge lane. **Proceed immediately to the next step.**

3. Run the mixed hybridization reactions on the nCounter platform:
   - **MAX/FLEX:** Run the full volume of the mixed hybridization reactions immediately on an nCounter Prep Station as described in the nCounter Analysis System User Manual (MAN-C0035). For 3D applications, it is recommended to run the Prep Station in “Standard” mode.
   - **SPRINT:** Run 30 μL of the mixed hybridization reactions immediately on an nCounter SPRINT Profiler as described in the nCounter SPRINT User Manual (MAN-10017).