NucleoZOL Total RNA Isolation Protocol

(Xinyi Yu & TCH @ 08/21/2017)

NucleoZOL Reagent, Cat# 740404.200 (BD ClonTech-Takara)

- LYSE CELLS: For cells grown in monolayer, remove cell culture medium and lyse cells by addition of 1mL of NucleoZOL to the culture disk (e.g., 1ml per one confluent 100mm dish; or 500µl per 60mm dish). For cells grown in suspension, spin down cells and lyse directly by the addition of NuceoZOL at 1mL NucleoZOL/10⁷ cells. Lyse cells by pipetting up and down several times. Ensure complete lysis by repeated pipetting.
- PRECIPITATE PROTEINS/CELL DEBRIS: Add 200µL RNase-free water per 500uL NucleoZOL to the lysate. Shake the sample vigorously for 15 seconds. Centrifuge samples for 5~8min at 12,000 x g at 4°C or room temperature.
- 3) **PRECIPITATE TOTAL RNA:** Carefully pipette RNA containing supernatant from step 2 into a 1.7ml or 2.0ml RNase-free Eppendorf tube(s). Add **1mL isopropanol** per **1mL supernatant** (**i.e., equal volume**) in order to precipitate RNA. Make sure tubes are kept on ice. Centrifuge samples for **5~8min** at **12,000 x** *g* at 4°C or room temperature. (**NOTE**: You can stop here by storing the isopropanol-RNA mix at -80°C for days to years).
- 4) WASH RNA: Use 400–600μL 70 % ethanol when precipitating in 1.5mL or 2mL tubes. For larger tubes, add 500-700uL 70 % ethanol per 1mL supernatant. Centrifuge the pellets for 1~3 min at 12,000 x g.
- 5) Remove ethanol from the pellet by carefully pipetting. Repeat the ethanol washing step. Do not dry the RNA pellet between wash steps.
- 6) RECONSTITUTE RNA: Dissolve the RNA pellet in RNase-free water to obtain an RNA concentration of 1~2μg/μL (i.e., usually dissolve in 20~30μl for one 100mm dish, or 10~15ul for one 60mm dish). Vortex the sample for 2~5min at room temperature to dissolve total RNA.
- 7) Your RNA samples should be aliquoted and kept at -80°C (not your own -20°C freezer space).

NOTE: Similar to TRIzol reagent, NucleoZOL reagent contains a very low level of quinidine to inhibit RNase activity. Thus, you should take the following considerations in mind:

- 1). You should try to collect one sample at a time (keep the rest in the incubator). Once you lyse the cells, please collect the lysate and transfer to 2ml-Eppendorf tubes. Please keep the tubes on ice.
- 2). The RNA will severely degrade if the cell lysate is kept in -20°C or -80°C. You should process the samples all the way to isopropanol precipitation/ethanol washes (even though you may leave the isopropanol-filled tubes at -80°C for weeks/months).
- 3). If you plan to store your RNA samples for quite long periods, you can dissolve the RNA samples and perform PC-8 extraction, followed by ethanol precipitation/washes. PC-8 extraction will remove RNase completely.