PROTOCOL FOR TOTAL RNA ISOLATION USING RNAGENT KIT Quan Kang 7/10/2003, Commented by TCH

1. Aspirate the medium from the Flask (T-75).

2. Add 2~3ml ice-cold Denaturing Solution, and let it cover all surface of cells.

3. Shake the flask slowly.

4. Collect the lysate into 15ml- or 50ml-conical tubes; it should be thick and snotty [Note: Cell scrapers may be used to collect the lysate]. Keep the lysate on ice and follow the step 5 or @-80 °C for several days to months

5. Add 300ul 2M NaAc (supplied w/ kit), mix well and keep on ice.

6. Add 3ml phenol-chloroform (supplied w/ kit), vortex vigorously; then divide into three to four 2ml microfuge tubes (just use the bottom phase of the PC)
7. Spin down at top speed for 10 minutes in the cold room.

8. Transfer the supernatant to a new set of 1.5ml microfuge tubes, and avoid touching the cloudy interphase. [**Optional:** *the supernatant can be pooled and extracted with equal volume of phenol-chloroform. This should further improve the RNA quality*].

9. Add 3-5ul **glycogen** and 1ml (or at least equal volume of the supernatant) isopropanol to each 1.5ml tube; mix well. [**Note: seeDNA cannot be used as a carrier as it interferes with the OD readings at 260nm**].

10. Incubate @ -80°C freezer for 2-4 hours or overnight [the tubes can be stored for a couple of days if you don't use it immediately].

11. Spin down @ top speed for **10 minutes in the cold room**; aspirate the supernatant.

12. Wash pellets with 600ul 70% EtOH, spin down for 5 minutes in the cold room; Repeat the wash once. In this step, try to combine the pellets for one sample into one tube.

13. Dissolve pellets in 30~50ul RNase-free water [**Note**: if it is difficult to dissolve, you can incubate the tube briefly at room temperature].

14. Check RNA concentrations and quality (**typical yield: 100-200ug per T-75**).