A Simple and Swift Method for Isolating High Quality RNA from Jute (*Corchorus* spp.)

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High quality RNA extraction is a prerequisite for various types of genetic analyses. Many a time, the existing RNA isolation methods and commercial kits are either time consuming or fail to isolate high quality RNA from plants rich in polysaccharides, oil and other secondary metabolites since different plants contain different amounts of nucleic acids (Khan et al. 2004, Loomis 1974). This problem is particularly acute in case of jute (*Corchorus* spp.), which is rich in mucilage and other polysaccharides that tends to interfere with the downstream processes (Kundu et al. 2011, Pandey et al. 1996). Several guanidium salt based methods have been successful for RNA isolation from jute seedlings (Khan et al. 2004), but are often cumbersome and expensive; hence limit simultaneous processing of large number of samples. Here we report a simplified and swift protocol for isolating high quality RNA from jute by making key modifications in tissue denaturation and precipitation steps in the protocol described by Ghawana et al. (2011). The protocol allows consistent production of high quality RNA from different species, which makes it particularly suitable for comparative plant genome research. The extraction time has been reduced from two days (for standard guanidium-acid-phenol extraction protocols) to about one hour and the extracted RNA was suitable for downstream processes like cDNA synthesis and expression pattern analysis.

Seeds of *Corchorus capsularis* (white jute) and *C. olitorius* (Tossa) were germinated on moist blot papers in Petri dishes at 25°C, seedlings were collected after 3 days, frozen in liquid nitrogen and immediately stored at – 80°C until the time RNA was isolated (Mahmood et al. 2010).

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About 0.5 g of sample tissue was pulverized to a fine powder in liquid nitrogen using pre-chilled RNase-free mortar and pestle. Then 3 ml of
denaturing solution containing 88% (v/v) DEPC-phenol saturated with Tris, 0.2% (w/v) sodium dodecyl sulphate (SDS), 4% (v/v) of 1 M sodium acetate and 8% (v/v) of 0.5 M EDTA (pH 8.0) was added. The samples, which freeze after the addition of denaturing solution, are allowed to thaw at room temperature. Next 1 ml of DEPC-treated water was added, mixed by grinding and allowed to stand for 5 min. The sample was then divided into four aliquots and each of them was transferred to a 1.5 ml RNase-free microcentrifuge tube and allowed to stand for another 5 min. Then 0.3 ml of chloroform was added to each tube; mixed well by inversion, vortexed briefly and incubated at room temperature for 5-10 min. The tubes were then centrifuged at 13,000 rpm for 10 min at 4°C and the aqueous phase was transferred to a fresh microcentrifuge tube using a sterile transfer pipette. An equal volume of 10 M LiCl was added to each tube, mixed well by inversion, vortexed briefly and incubated at room temperature for 10 min to allow the nucleic acids to precipitate. After precipitation, the samples were centrifuged at 13,000 rpm for 10 min at 4°C and the supernatant was discarded. The pellet was then washed with 0.5 ml of 70% ethanol and centrifuged at 13,000 rpm for 10 min at 4°C. The supernatant was discarded and the tubes were kept inverted with their lids open on a stack of tissue papers in a laminar hood for 5-10 min. Next the pellet was air-dried for 2 - 3 min. The RNA pellet was then re-suspended in 20 μl of DEPC-treated water.

The quality and concentration of the RNA samples were checked using a spectrophotometer (NanoDrop, Technologies Inc.). The ratio of A$_{260}$/A$_{280}$ was within 1.8 to 2.0 signifying high purity of RNA without any protein contamination (Table 1). Further, the ratio of A$_{260}$/A$_{230}$ was greater than 1.8 indicating the absence of contamination proteins.

<table>
<thead>
<tr>
<th>Sample species</th>
<th>A$<em>{260}$/A$</em>{280}$</th>
<th>A$<em>{260}$/A$</em>{230}$</th>
<th>RNA yield (ng of RNA/g tissue)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C. capsularis</td>
<td>2.07</td>
<td>1.88</td>
<td>8507.50</td>
</tr>
<tr>
<td>C. olitorius</td>
<td>2.10</td>
<td>1.90</td>
<td>8246.75</td>
</tr>
<tr>
<td>A. thaliana</td>
<td>2.10</td>
<td>2.01</td>
<td>5273.75</td>
</tr>
</tbody>
</table>

To validate the effectiveness of this protocol in other plant species, high quality RNA has also been isolated from the seedlings of model plant Arabidopsis thaliana in the same way as described above.

The integrity of total RNA was determined by distinct 28 S and 18 S rRNA bands after running the samples on a 1.2% denaturing formaldehyde-agarose gel
To confirm that the isolated RNA was amenable to downstream applications, reverse transcription-polymerase chain reaction (RT-PCR) was done. For RT-PCR, 3 μg of poly (A+) RNA was reverse transcribed by SuperScript™ First-Strand Synthesis System of Invitrogen, USA, using a thermal cycler (Eppendorf Mastercycler Personal). Next normal PCR was carried out using the single stranded c-DNA with universal primers (β-actin) and the resultant double stranded DNA was checked by agarose gel electrophoresis. The presence of bands of desired sizes validated the success of the protocol for downstream processes (Fig. 1b).

In *Corchorus* spp., the nucleic acids are most often found to be contaminated with protein, possibly due to the highly proteinaceous nature of mucilages (Kundu et al. 2011). Therefore in order to remove these proteins strong denaturants like phenol and SDS were used during this experiment. Maintenance of acidic pH is the critical factor to ensure the separation of RNA from DNA and proteins. Under acidic conditions, total RNA remains in the upper aqueous phase, while most of DNA and proteins remain either in the interphase or in the lower organic phase (Chomczynski and Sacchi 2006). So, the denaturing solution was composed of such reagents that could ensure the overall acidic condition of
the reaction. During this study, RNA precipitation was done by LiCl; as it does not efficiently precipitate DNA, protein or carbohydrate (Barlow et al. 1963, Cathala et al. 1983). It was thus used to remove these inhibitors of protein translation or cDNA synthesis.

Previously, Khan et al. (2004) reported another protocol for RNA isolation from jute seedlings. The distinguishing feature of the protocol described here is its simplicity, time and cost-effectiveness; which makes it suitable for simultaneous isolation of good quality RNA from a large number of samples as required for expression analyses (Fig. 2).

![Graph](produced by Nanodrop software) showing the highest peak at a range close to 260 nm wavelength indicating efficient isolation of high quality RNA from 17 different samples within a very short period of time.

It can be concluded that this modified protocol is highly effective for extracting good quality RNA from jute vis-à-vis other plant species.

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**References**

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