RNA Extraction Method from Fruit Tissue High in Water and Sugar

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Abstract. RNA isolation from ripe fruit can be complicated by high concentrations of sugar and water. These sugars interfere with RNA extraction often resulting in low RNA quality and quantities, and high water concentrations dilute the RNA, making isolation difficult. We report a simple but novel method by which the majority of the excess sugar and water in mature fruit of tomato (Lycopersicon esculentum Mill.), watermelon (Citrullus lanatus (Thunb.) Matsum. & Nakai), and muskmelon (Cucumis melo L.) can be easily removed from tissue before RNA extraction. This method produced quality RNA in a shorter time than the currently accepted method for fruit tissue RNA isolation and does not require liquid nitrogen or a freeze dryer.

RNA is used in a multitude of laboratory experiments from Northern blots to reverse transcriptase–polymerase chain reactions (RT-PCR). Isolating high-quality RNA is important in the process of gene discovery and for studying temporal expression of genes during sequential developmental stages of plants. Unfortunately, the instability of RNA and the ubiquitous nature of RNases make RNA highly susceptible to degradation

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(Ausubel et al., 1997). Further complications arise from contaminants in the original tissue that render the isolated RNA unfit for downstream applications. For example, tissues from ripe fruit and seeds often have high concentrations of lipids, phenolic compounds, and polysaccharides and low concentrations of RNA molecules, causing low recovery or difficulty quantifying the sample (Asif et al., 2000; Callahan et al., 1989; Levi et al., 1992; Loomis, 1974; Taylor and Powell, 1982). Several groups have reported alternative protocols improving RNA yields and reducing contamination problems inherent in some fruit tissue (Asif et al., 2000; Callahan et al., 1989; Gebrig et al., 2000; Levi et al., 1992; Logemann et al., 1987; Mozer, 1980; Taylor and Powell, 1982). Callahan et al. (1989) reported a method to remove excess water from peach flesh to isolate high-quality RNA. However, this method requires the use of a freeze dryer to remove the excess moisture, which in turn concentrates the carbohydrates present in the dried sample. It may take up to 7 d to completely dry large or multiple samples, which increases the exposure time of RNA to oxidation and RNases, increases the potential for sample thawing resulting from mechanical failure, and slows down the isolation process. Additionally, the freeze-drying process makes the tissue difficult to grind by mortar and pestle or Polytron (Brinkmann Instruments Inc., Westbury, N.Y.). Therefore, these samples often do not fully homogenize in commercial RNA extraction solutions.

In this study, we report an efficient and economical procedure to reduce water and carbohydrate content in fruit tissue. The resultant RNA is of sufficient quality for downstream applications such as RT-PCR and 5' and 3' RACE.

Materials and Methods

'Fruit and sample preparation. Ripe fruit from watermelon ('Tri-x 313', 'Illini Red', and 'Summer Gold'), ripe red tomatoes (Roma type, 'Rutgers', and 'T5020'), and ripe and immature muskmelon ('Magnum 45' 15 d, and 2 d postpollination) were used for this study. They were grown and collected at the South Central Agricultural Research Laboratory in Lane, Okla. The fruit were rinsed with deionized water in the field, and small samples of flesh tissue (~5–10 g) were excised and processed using our juice extraction RNA isolation method and a conventional RNA freeze-dried method described subsequently. All glassware and utensils were purchased sterile or were cleaned with RNase Zap (Ambion Inc., Austin, Tex.), and rinsed with RNase-free water before each use. Utensils used in the field were washed between samples with 95% ethanol and allowed to air dry. Total soluble solids (Brix) were determined using a Reichert 10,430 refractometer (Cambridge Instruments Inc., Buffalo, N.Y.) on juice squeezed from fruit flesh tissue using a 7-inch sap extractor.

Juice extraction RNA isolation method. Flesh tissue was excised from ripe fruit; the juice was removed and discarded by squeezing the excised flesh with a sap extractor (such as a 7-inch Sap Extractor; Peaceful Valley Farms, Grass Valley, Calif.). The pulp was immediately scraped into a preweighed, prelabeled (etched, so liquid nitrogen would not remove the label) sterile 50-ml conical polypropylene tubes (Blue Max; Becton Dickinson Labware, Franklin Lakes, N.J.) that contained a predetermined amount of TRIzol (GibcoBRL In vitrogen; Life Technologies, Gaithersburg, Md.) reagent. For our purposes, 5 ml was used. The sample was then immediately placed on wet ice for transport to the laboratory. Liquid nitrogen should be used if available as an extra precaution against degradation. The tubes were reweighed and more TRIzol was added to bring the concentration to 1 ml of TRIzol per 100 mg of sample. Samples were homogenized at room temperature with a Brinkmann Polytron PT 10/35 with a 20-mm grinding blade for 60 to 90 s. The samples were not allowed to heat or froth. Polytron operation was performed in a fume hood with protective clothing. The homogenized samples were stored at ~80 °C until the RNA extraction procedure was performed. Total RNA was extracted using the TRIzol reagent following the company's protocol. Both
optional steps to reduce polysaccharide contamination were performed. This included a centrifugation step and a high salt precipitation step.

Testing RNA quality. Agarose and denaturing gel electrophoresis, Northern blotting, and PCR were performed using standard techniques (Ausubel et al., 1997). 5’ and 3’ RACE (GibcoBRL Invitrogen; Life Technologies) was performed according to company protocols. The denaturing RNA gel and Northern blot experiments were run using the NorthernMax, BrightStar PsoralenBiotin, and BrightStar BioDetect kits (Ambion) following the company’s protocol. Agarose gels were visualized on a Kodak Image Station 440CF (Eastman Kodak Co., Rochester, N.Y.) and were used to verify RNA quality and concentration determined by spectral analysis. All isolated RNA samples were diluted in 1 mM Na2HPO4 in nuclease-free water (pH 8.3) and readings were taken at 260 nm and 280 nm wavelengths using a Perkin Elmer Co. ultraviolet/VIS Spectrophotometer Lambda 3B (Wellesley, Mass.). A quartz 100-μl cuvette was used and the instrument was zeroed using the above diluent. A subset of samples was continuously scanned from 210 nm to 310 nm wavelengths using a NanoDrop ND-1000 Spectrophotometer (Wilmington, Del.).

Conventional RNA freeze-dried method. On excision, fruit tissue was immediately placed in a preweighed, prelabeled sterile 50-ml conical polypropylene tube. The tube was immediately placed in liquid nitrogen for transporting to a LabConco Co. Freeze Dryer 18 (Kansas City, Mo.). The samples were freeze-dried under shelf refrigeration at −35 °C until samples were fully dry (Callahan et al., 1989). This took from 24 h to 7 d depending on the size, number of samples, and the moisture content of the tissue samples. Once lyophilized, the preweighed tubes containing the samples were again weighed and TRIzol reagent was added to a final concentration of 1 ml TRIzol per 100 mg of sample. Samples were homogenized with a Polytron as described previously and stored at −80 °C until RNA extraction was performed as described.

Results and Discussion

Table 1. Yield and 260/280 ratio of RNA samples isolated using our juice extraction method and the conventional freeze-dried method.

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<tr>
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<th>Freeze-dried method</th>
<th>Juicy extraction method</th>
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<tbody>
<tr>
<td>Number of samples</td>
<td>Yield average (ug.g⁻¹)</td>
<td>Standard deviation</td>
</tr>
<tr>
<td>Watermelon</td>
<td>15</td>
<td>34</td>
</tr>
<tr>
<td>Tomato</td>
<td>6</td>
<td>42</td>
</tr>
<tr>
<td>Cantaloupe</td>
<td>2</td>
<td>15</td>
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*Total RNA yield from 1 g fresh sample.

Total RNA from 20 mature fruit samples processed using the juice extraction method yielded RNA in the range of 5 to 22 μg per 1 g fresh tissue. The average for the 12 mature watermelons was 11 μg.g⁻¹ fresh tissue and the average for six tomatoes was 13 μg.g⁻¹ fresh tissue. The average of two mature muskmelon fruits yielded 15 μg.g⁻¹ fresh tissue. This was a decrease over the freeze-dried method, which gave yields of 15 to 57 μg per 1 g fresh tissue from 21 fruit. The average of 15 freeze-dried mature watermelons was 34 μg.g⁻¹ fresh tissue and six tomatoes was 42 μg.g⁻¹ fresh tissue (Table 1). Cantaloupe was not analyzed using the freeze-dried method, but the juice extraction method on immature fruit samples yielded much higher RNA concentrations than on mature fruit. The juice extraction method yielded 236 μg.g⁻¹ fresh tissue from immature muskmelon and 146 μg.g⁻¹ fresh tissue from immature watermelon. RNA concentrations were verified at the University of Illinois on 17 of the samples using a NanoDrop spectrophotometer. This comparison determined that our analytical spectrophotometer was giving accurate concentration estimates. To determine if our RNA was of sufficient quality to perform accurate concentration estimates using ultraviolet spectrophotometry, we compared our estimated concentrations with samples run on denaturing agarose gels. The spectral data were used to estimate the amount of RNA needed to load each lane of the gels in Figures 1 and 2. These gels demonstrated the RNA was of high quality and can be accurately estimated using a spectrophotometer.

RNA quality was assessed using ultraviolet spectrophotometry (Tables 1 and 2), visualizing intact ribosomal RNA on a denaturing RNA gel (Figs. 1A and 2), detecting message on Northern blots (Fig. 1B), amplifying appropriately sized bands in RT-PCR (data not shown), and sequencing the 5’ and 3’ ends of watermelon genes using RACE (data not shown). Genomic DNA was not detected on denaturing gels (data not shown), Northern blots were performed on muskmelon total RNA purified with the juice extraction RNA isolation method using a tomato pgip probe and produced clean blots (Fig. 1). The results from these methods showed that RNA purified using the juice extraction RNA isolation method was free from genomic DNA and could be used to detect intact ribosomal RNA.
that the juice extraction method is not prefer-
entially isolating large RNA, which suggests that nuclear and cytoplasmic RNA are being retained equally. It is important to note that we may be preferentially selecting for RNA in certain cell types since we may be losing intact cells while squeezing out the liquid.

Removal of water and carbohydrates from juicy fruit is critical for obtaining high-quality and sufficient quantities of RNA. The procedure reported in this study can be useful for rapid removal of water and sugar for RNA extraction from a wide array of juicy fruits. It does not require use of a freeze-dryer and reduces the opportunity for RNA degradation resulting from mechanical failure or sample contamination from neighboring samples. It is simple enough to be used in the field, avoiding shipping-related changes in gene expression, and eliminates the need of liquid nitrogen, making it possible to collect RNA samples from almost anywhere.

By eliminating the water and water-soluble material, our method reduces the amount of sample weight and total soluble solids in watermelon by ∼90% (data not shown). However, when we tested the RNA concentration in the eliminated liquid, we found that we lose ∼50% of the RNA to the water-soluble material. This procedure results in a fivefold concentration of the RNA, whereas the freeze-dried method concentrates the RNA 10-fold. This is likely why the juice extraction method gives lower yields than the freeze-dried method.

Figure 2 shows RNA samples purified using the RNA juice extraction method and the freeze-dried method. This figure shows that the percent of small RNAs present is equivalent in both methods. This demonstrates that the juice extraction method is not prefer-
entially isolating large RNA, which suggests that nuclear and cytoplasmic RNA are being retained equally. It is important to note that we may be preferentially selecting for RNA in certain cell types since we may be losing intact cells while squeezing out the liquid.

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It is important to understand that certain cells within a given tissue may be more susceptible to removal with the juice or more susceptible membrane disruption and therefore may be prone to RNA losses. We feel this method will be useful for PCR sequenc-

### Literature Cited


