Total RNA Extraction from the Aromatic *Phalaenopsis bellina*,
Endemic Orchid in Sabah, Borneo

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Abstract

*Phalaenopsis bellina* is an attractive orchid due to its unique appearance and distinctive floral fragrance. Many past studies on this plant focused on the plant at the molecular level; however, this requires sufficient quantities of high-quality *P. bellina* RNA. RNA is more delicate to manipulate than DNA due to its structural instability and its vulnerability to various secondary metabolites, such as polyphenols and polysaccharides. Therefore, in this study, 4 RNA isolation methods, a modified phenol-chloroform method and 3 commercial kits (Vivantis, Novogene, and Analytik Jena) were used on the leaves and flowers of *P. bellina* for comparison. The yield and purity of the total RNA were determined using spectrophotometry. The results showed that the total RNA isolated using the modified phenol-chloroform method had the highest yield (1223.75±68.51 ng/µL) and purity compared to the 3 commercial kits, with an OD260/280 value of 2.07 and an OD260/230 value of 2.26, respectively. In particular, the isolated RNA did not show any detectable genomic DNA contamination or other impurities. The RNA isolated using the phenol-chloroform method was also evaluated by electrophoresis, reverse transcription, and PCR. The results indicated that the phenol-chloroform method appears to be superior for total RNA extraction. Thus, this developed method is proven to be suitable for the RNA extraction of plants rich in polysaccharides and polyphenols and is amenable for future molecular studies on *P. bellina*.

Keywords: *Phalaenopsis bellina*, Orchidaceae, RNA, Phenol-chloroform, Isolation, Flower

Introduction

*Phalaenopsis bellina*, commonly known as the “Lundu orchid” or “Norma orchid”, is an orchid endemic to Borneo [1]. The word *Bellina* is derived from the Latin word Bella, which means beauty or lovely [2]. The name befits its unique appearance, where the flower is brightly colored in white, greenish-white, yellow or, on rare occasions, orange tepals with an intense magenta blotch on the inner halves of the lateral sepals. Apart from its distinct floral appearance, the flower also emits a strong-sweet fragrance, with a hint of lemony and citrusy odors [3,4]. Due to its unique characteristics, *P. bellina* is often used as a parent for the breeding of scented hybrid cultivars [5].

Many of the molecular studies conducted on *P. bellina* revolved around the biosynthesis of the fragrance in the *P. bellina* flowers. To our knowledge, Hsiao et al. [6] was the first author who attempted to elucidate the fragrance in *P. bellina*. The researchers had conducted chemical profiling, along with gas chromatography mass spectrometry (GC-MS), the identification of floral expressed sequence tags (EST), and bioinformatics analysis. The authors managed to pinpoint 66 % of the biosynthetic steps from glyceraldehyde-3-phosphate (G3P) to the production of various fragrance derivatives, particularly that of...
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High-quality Ribonucleic Acid (RNA) must be extracted for studies on gene expression in plants. Reverse Transcription-Polymerase Chain Reaction (RT-PCR), Complementary Deoxyribonucleic Acid (cDNA) synthesis, and subsequent genetic analysis also require RNA with high purity and integrity [10]. However, obtaining high quality RNA from plant samples remains difficult. The presence of polysaccharides, polyphenols, lipids, and other secondary metabolites that may bind to the nucleic acids during extraction may interfere with subsequent molecular studies [11-14]. A large number of plant-specific RNA extraction protocols have been highlighted. All of these methods recommend a diversity of strategies to isolate sufficient RNA from plants, depending on a plant’s species condition [15]. Liu et al. [10] managed to isolate total RNA from *Dendrobium huoshansense*, an orchid that is known to have high levels of polysaccharides and polyphenols, using methods modified from those of Chan et al. [16]. Meanwhile, Abd Rahman et al. [17] managed to isolate the RNA from mangosteen (*Garcinia mangostana* L.) and screened it for RNA sequencing suitability. Compared to Liu et al.’s work [10], the method developed by Abd Rahman et al. [17] required 3 days of complete extraction, which is longer and time-consuming. Das et al. [18] optimized an RNA extraction method on tea roots (*Camellia sinensis*) based on sodium dodecyl sulphate (SDS) for the construction of the plant’s cDNA library. Ma and Yang [19] attempted to isolate RNA from the dry seeds of the sunflower (*Helianthus annuus*) using RNAiso, RNAiso Plus + RNAiso-mate, and Trizol kits. However, no total RNA of high quality was obtained.

To our knowledge, no protocol has been developed for the optimized RNA extraction of *P. bellina*. Therefore, the leaves and flowers of *P. bellina* were selected as the subjects of total RNA extraction by applying the modified phenol-chloroform method, as well as 3 other extraction kits (Vivantis, Analytik Jena, and Novogene). RNA extraction method suitability was chosen based on RNA purity ratio (A260/A280 and A260/230), gel electrophoresis, and PCR assessment.

Materials and methods

Orchid collection and maintenance

*Phalaenopsis bellina* orchid was brought from local vendors and maintained in the Institute for Tropical Biology and Conservation (ITBC) greenhouse, Universiti Malaysia Sabah. The orchid’s flowers and leaves were collected from April to October 2018. Harvested flowers and leaf samples were kept in liquid nitrogen before being stored at −80 °C for further use.

Total RNA extraction

RNA was extracted from the leaves and flowers of *P. bellina*. Three commercial kits were used [Analytik Jena (Germany), Vivantis (Malaysia), and Novogene (Singapore)] according to each respective manufacturer’s instruction. A brief overview of methods used by each kit is summarized in Table 1.
Table 1 Overview of methods used by the 3 kits for total RNA extraction.

<table>
<thead>
<tr>
<th>Steps/Brand</th>
<th>Analytik Jena</th>
<th>Vivantis</th>
<th>Novogene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell lysis</td>
<td>Uses readymade lysis buffer</td>
<td>Uses readymade lysis buffer and β-mercaptoethanol</td>
<td>Uses readymade lysis buffer and β-mercaptoethanol</td>
</tr>
<tr>
<td>Removal of genomic DNA</td>
<td>Included in the kit</td>
<td>Included in the kit</td>
<td>Not included in the kit</td>
</tr>
<tr>
<td>Selective binding of RNA</td>
<td>Utilizes different column (RNA binding column)</td>
<td>Utilizes different column (RNA binding column)</td>
<td>The same column used in the whole process</td>
</tr>
<tr>
<td>Washing of bound RNA</td>
<td>Ethanol-based washing buffer</td>
<td>Ethanol-based washing buffer</td>
<td>Ethanol-based washing buffer</td>
</tr>
<tr>
<td>Main component used</td>
<td>Guanidinium thiocyanate and guanidinium chloride</td>
<td>Guanidinium thiocyanate</td>
<td>Guanidinium thiocyanate</td>
</tr>
</tbody>
</table>

Manual RNA extraction was done by adapting methods loosely from Lewis et al. [20] with slight modifications. Concentration of LiCl was increased to 4M instead of 2M for total RNA precipitation, while total genomic DNA was completely removed using ethanol precipitation twice. Approximately 0.25 g of *P. bellina* petals or leaves were ground into powder in liquid nitrogen. The powdered sample was then quickly transferred into a falcon tube (Fischer, Malaysia) containing 5 mL of extraction buffer [250 mM (Tris-Hydrochloric acid) (Tris-HCl) (pH 7.5), 375 mM of Sodium Chloride (NaCl) (Sigma Aldrich, USA), 25 mM of Ethylenediaminetetraacetate disodium salt dihydrate (Na2EDTA) (pH 8.0) (Sigma Aldrich, USA), 1 % (w/v) of SDS (Sigma Aldrich, USA), and 1 % (v/v) of β-mercaptoethanol (Sigma Aldrich, USA)], and was vortexed immediately for 15 s. Then, 3 mL of phenol (Sigma Aldrich, USA) was added and it was vortexed again for 15 s. This was followed by the addition of 3 mL chloroform:isoamyl alcohol (24:1, v/v) (Sigma Aldrich, USA) and it was vortexed for 15 s. Then, the sample was centrifuged for 10 min at 5000 rotations per min (rpm). Aqueous or top phase obtained was transferred into a new tube and 3 mL of phenol:chloroform (1:1, v/v) (Sigma Aldrich, USA) was added. The sample was vortexed for 15 s and centrifuged for 10 min at 5,000 rpm using ultracentrifuge (Thermo Scientific, USA). The aqueous or top phase obtained was transferred into a new tube containing 8 mL of 4M LiCl (Sigma Aldrich, USA) to precipitate total nucleic acid. The sample was then incubated in −20 °C overnight to maximize nucleic acid precipitation.

After the overnight incubation, the sample was centrifuged at 7,000 rpm for 30 min at 4 °C to pellet out the total nucleic acid. The supernatant was then removed and the tube was dried on a paper towel for 10 min. Then, the nucleic acid pellet was resuspended in 500 μL of 0.1 % DEPC-treated deionized water. The sample was centrifuged again at 3,000 rpm for 5 min to pellet the insoluble material. The supernatant obtained was then transferred into a microcentrifuge tube containing 500 μL of 4M LiCl (Sigma Aldrich, USA). The mixture was mixed well and incubated at 4 °C overnight to precipitate the RNA.

After overnight incubation, the sample was centrifuged at 13,200 rpm for 15 min at room temperature to pellet total RNA. 500 μL of 70 % (v/v) ethanol (Sigma Aldrich, USA) was added, and then it was centrifuged again for 15 min at 13,200 rpm. The supernatant was removed and the ethanol step was repeated twice. Finally, the RNA pellet was air-dried for 10 min and dissolved in 50 μL of 0.1 % DEPC-treated deionized water. The RNA sample was then kept in −80 °C for future use.

**RNA analysis**

The total RNA extracted from the leaves and flowers of *P. bellina* was tested for integrity using 1 % Weight over Volume (w/v) agarose gel (Sigma Aldrich, USA) at 120V for 30 min. The A260/230 and A260/280 values for the RNA sample were measured using the Implen Nanophotometer® P-Class (Implen, Germany) to evaluate the RNA purity and yield.
First strand cDNA synthesis and PCR amplification

Approximately 2 μg of RNA was used for the first strand cDNA synthesis using the Viva cDNA synthesis kit (Vivantis, Malaysia) according to its manufacturer’s instruction. Two types of primers OligoDT (18) and a gene-specific primer were used. The candidate gene from the terpene synthase group was mined from Genebank’s database. The full-length nucleotide sequence of Phalaenopsis equestris’s terpene synthase gene (ascension number: EU124717.1) was used as a reference gene. The primer design was carried out using idtDNA tools and the Primer 3 software for specific PCR amplification. The forward primer of the terpene synthase used was 5’-TGT AAT GGA GAT TCA GCC ATC TTC - 3’, while the reverse primer was 5’TAC TAC CAC TTT GAC AAA ATA JGT TCC-3’.

PCR amplifications were carried out using degenerate primer pairs designed for the gene reference of P. equestris. A total reaction volume of 12.5 µL, containing 6.75 µL of GoTaq® Green Master Mix (Promega), 0.5µL of 10µM forward and reverse primer, and 2µL cDNA, was used. Amplifications were performed in a thermocycler (Applied Biosystem, USA) and programmed as follows: initial denaturation at 94 °C for 2 min, 30 cycles at 94 °C for 30 s, 56 °C for 30 s, then 72 °C for 2 min. A total of 5 µL of the PCR product was separated on 1.0 % agarose gel containing SYBR Safe (0.01 % v/v) (Thermo Fischer, USA) and TBE buffer (1 ×) at 100 V for 30 min. Gels were visualized under UV light and photographed with a gel image analyzer (Applegen, USA). 1Kb of DNA ladder (Invitrogen, USA) was used for base pair size estimation. The remaining products were stored at −20 °C.

Results and discussion

Although RNA isolation is considered a routine protocol, the extraction of RNA from aromatic plants could be critical due to the presence of large amounts of polysaccharides, polyphenols, and other secondary metabolites which limit the RNA extraction yield and purity [21]. There are many published RNA isolation protocols of which the single-step method using guanidine isothiocyanate [22] has proven to be the most useful in model plants, i.e., rice [23] and Arabidopsis [24]. The usage of guanidinium isothiocyanate could inhibit RNA activity, but it is not quite suitable for plant tissues rich in polysaccharides and phenolics [25].

Table 2 Quantitative analysis of total RNA extraction from flowers and leaves of P. bellina.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Method/kit</th>
<th>Column based?</th>
<th>Yield (ng/µL) for 50 µL sample</th>
<th>A_{260}/A_{280}</th>
<th>A_{260}/A_{230}</th>
<th>Total time required</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flowers</td>
<td>1AJ</td>
<td>Yes</td>
<td>32.33 ± 2.52</td>
<td>1.88 ± 0.25</td>
<td>0.091 ± 0.01</td>
<td>2 h</td>
</tr>
<tr>
<td></td>
<td>2V</td>
<td>Yes</td>
<td>264.67 ± 4.62</td>
<td>2.09 ± 0.01</td>
<td>2.39 ± 0.01</td>
<td>1 ½ h</td>
</tr>
<tr>
<td></td>
<td>3NV</td>
<td>Yes</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>Half day</td>
</tr>
<tr>
<td></td>
<td>Modified phenol-chloroform</td>
<td>No</td>
<td>1223.75 ± 68.51</td>
<td>2.07 ± 0.00</td>
<td>2.26 ± 0.05</td>
<td>3 days</td>
</tr>
<tr>
<td>Leaves</td>
<td>1AJ</td>
<td>Yes</td>
<td>15.0 ± 8.66</td>
<td>N/A</td>
<td>0.16 ± 0.09</td>
<td>2 h</td>
</tr>
<tr>
<td></td>
<td>2V</td>
<td>Yes</td>
<td>41.33 ± 0.58</td>
<td>2.54 ± 0.06</td>
<td>0.08 ± 0.00</td>
<td>1 ½ h</td>
</tr>
<tr>
<td></td>
<td>3NV</td>
<td>Yes</td>
<td>11.3 ± 0.58</td>
<td>6.67± 0.85</td>
<td>0.62 ± 0.02</td>
<td>Half day</td>
</tr>
<tr>
<td></td>
<td>Modified phenol-chloroform</td>
<td>No</td>
<td>249.75 ± 3.50</td>
<td>1.95 ± 0.10</td>
<td>2.13 ± 0.11</td>
<td>3 days</td>
</tr>
</tbody>
</table>

*1AJ = Analytik Jena, Germany, 2V = Vivantis, Malaysia, and 3NV = Novelgene, Singapore
*Data represent mean ± SD of 3 replicates
The RNA extraction from flowers was done using a modified phenol-chloroform method adapted from Lewis et al. [20] and 3 other commercial kits (AJ, V, and NV) (Table 2). Of all methods conducted, the AJ and V kits, along with the modified phenol-chloroform method, managed to isolate the total RNA from the *P. bellina* flowers at acceptable A260/A280 values. However, the AJ commercial kit did not manage to get full purity according to its A260/A230 reading, and the total RNA extracted appeared to be degraded (Figure 1). Although the V commercial kit had managed to obtain an acceptable total RNA yield with satisfactory A260/A280 and A260/230 values, no bands or smears were observed. The modified phenol-chloroform method managed to get the highest total RNA with suitable A260/A280 and A260/230 values. The NV kit did not manage to isolate the total RNA from the *P. bellina* flowers.

![Figure 1](image1.png)

**Figure 1** Qualitative results of total RNA extracted from flower tissue of *P. bellina* using different methods. Lane M: 1kb DNA ladder. Lane 1: using the commercial kit NV. Lane 2: using the commercial kit V. Lane 3: using the commercial kit AJ. Lane 4: using the modified phenol-chloroform method.

Apart from the flowers, total RNA was also extracted from the *P. bellina*’s leaves. All methods had managed to isolate the total RNA from the leaves. The AJ and NV kits yielded low total RNA values from the leaves; their A260/A280 and A260/230 values were too high or low, thus indicating contamination. The total RNA extracted using the AJ kit also appeared to be degraded when visualized under gel electrophoresis (Figure 2). While the V kit managed to isolate the total RNA from the leaves at a moderate yield, no bands were observed. The same goes for the NV kit. Compared to the other methods, the modified phenol-chloroform method managed to isolate the highest total RNA yield from the leaves (249.75 ± 3.50) with satisfactory A260/A280 and A260/230 values.
Figure 2 Qualitative results of total RNA extracted from leaf tissue of *P. bellina* using different methods. Lane M: 1kb DNA ladder. Lane 1: using the commercial kit NV. Lane 2: using the commercial kit V. Lane 3: using the commercial kit AJ. Lane 4: using the modified phenol-chloroform method.

All of the commercial kits (AJ, V, and NV) used column-based technologies that resulted in low RNA yield. The column-based technologies employed created other problems, which included the complicated manipulation of buffers while loading and washing, or sensitivity problems due to non-specific binding to the solid phase membrane provided in the column [17]. The manual method used was loosely modified from Lewis et al. [20] who had utilized phenol-chloroform extraction. The use of the extraction buffer containing EDTA acted as a chelating agent to sequester divalent cations, thereby inhibiting the deoxyribonuclease (DNase), ribonuclease (RNase), and enzyme activity to prevent the rapid degradation of the RNA. NaCl was used to control the lysate pH and create osmotic pressure [26]. Meanwhile, SDS used in the extraction acted as a detergent that effectively broke down the cell membrane [27]. β-mercaptoethanol was also used in the extraction buffer to counter oxidation [25] and eliminate RNases [28].

The use of the phenol and phenol-chloroform methods takes advantage of the nucleic acid structure. Nucleic acids have a negative charge due to their phosphate backbones, making them soluble in aqueous solutions. Proteins, lipids, and carbohydrates, on the other hand, have hydrophobic and hydrophilic domains, making them soluble in organic solutions or selective for the interface between the organic and the aqueous phases generated during the extraction process. In phenol extraction, phenol is added to an aqueous solution containing cellular constituents. The mixture and subsequent centrifugations separate the phases. At neutral or near-neutral pH, nucleic acids remain in the aqueous phase. RNA may be selectively extracted in an acidic environment because the phosphate groups of RNA are preferentially neutralized, and the DNA is thus trapped in the organic phase [29]. Elkins [30] also stated that the usage of phenol can further degrade DNA. Chloroform:isoamyl alcohol was used in between phenol and phenol:chloroform to separate the aqueous and organic phases and prevent foaming during centrifugation [31].

Nucleic acid (DNA and RNA) structures which contain phosphate groups cause the nucleic acids to act as negatively-charged polar molecules; therefore, they possess hydrophilic characteristics. This makes them completely insoluble in alcohols. To selectively precipitate RNA, lithium chloride has been used due to its ability to precipitate RNA and remove polysaccharides [17].
In terms of total time required, the modified phenol-chloroform method required 3 days to yield the RNA, compared to the commercial kits, which required less than a day. Tüzmen et al. [26] stated that the use of the phenol-chloroform method is indeed laborious, time-consuming, and generates toxic waste. However, the total RNA yield from both the leaves and flowers of the *P. bellina* was high compared to the commercial kits used. The total RNA extracted from the *P. bellina* flowers was higher than from its leaf and more noticeable when visualized under agarose gel electrophoresis than the latter. Liu et al. [10] suggested that the water content of each plant structure (leaf and flower) is one of the factors contributing to the quality of total RNA. Although this method was tedious, it is interesting to know that Liu et al. [10] have also developed a high-quality RNA extraction method for *Dendrobium huoshansense* which only takes 2 and a half h and utilizes reagents that are almost similar to the method developed in this research, such as the composition of extraction buffer, and the usage of phenol:chloroform (1:1, v/v), β-mercaptoethanol, and LiCl for RNA precipitation. Meanwhile, Abd Rahman et al. [17] took 3 days to isolate total RNA from *Garcinia mangostana*. It should be noted that different species require different RNA extraction protocols due to the diversity in plant structure and secondary metabolite content.

Since flowers contain the highest RNA yield, the total RNA from *P. bellina* flowers was used for reverse transcription to synthesize the cDNA and subsequent PCR (Figure 3). The cDNA can be amplified to obtain a target fragment of 650 bp when synthesizing cDNA using gene-specific primer or when synthesizing a cDNA using OligoDT (18). This shows that the total RNA extracted using the modified phenol-chloroform method can meet the requirements of related molecular biological researches.

**Figure 3** PCR amplification of cDNA. Lane M: 1kb DNA ladder. Lane 1: cDNA synthesized using gene-specific reverse primer (the same primer was also used for PCR). Lane 2: cDNA synthesized using OligoDT (18).
Conclusions

By using the modified phenol-chloroform method, the total RNA from the leaves and flowers of *Phalaenopsis bellina* can be extracted at high yield and quality, although the method is time-consuming and laborious. It is hoped that this study can be applied in further molecular studies on *Phalaenopsis bellina*.

Acknowledgements

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References


