Nucleic acid protocols: Extraction and optimization

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Abstract

Yield and quality are fundamental features for any researchers during nucleic acid extraction. Here, we describe a simplified, semi-unified, effective, and toxic material free protocol for extracting DNA and RNA from different prokaryotic and eukaryotic sources exploiting the physical and chemical properties of nucleic acids. Furthermore, this protocol showed that DNA and RNA are under triple protection (i.e., EDTA, SDS and NaCl) during lysis step, and this environment is improper for RNase to have DNA liberated of RNA and even for DNase to degrade the DNA. Therefore, the complete removal of RNA under RNase influence is achieved when RNase is added after DNA extraction, which gives optimal quality with any protocols. Similarly, DNA contamination in an isolated RNA is degraded by DNase to obtain high-quality RNA. Our protocol is the protocol of choice in terms of simplicity, recovery time, environmental safety, amount, purity, PCR and RT-PCR applicability.

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1. Introduction

Biomolecule extraction, such as deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) from a variety of starting biological materials to be used in downstream applications and other analytical or preparative purposes, is the most important first step in the molecular biology. The widely employed nucleic acid isolation methods can be divided into organic extraction method (phenol/chloroform), inorganic extraction method (salting out) and solid phase extraction method (solid matrix); moreover, four indispensable steps are generally required for successful nucleic acid purification:

1. Cell lysis through disruption of the cellular membranes, cyst wall or egg wall
2. Dehydration and precipitation of the cellular proteins (protein denaturation)
3. Separation of cellular proteins and other cellular components out of the nucleic acid

4. Precipitation and dissolving the nucleic acid

The routinely practised cell lysis step can be divided into three types to cope with different tissues, thereby achieving optimum nucleic acid yield:

1. Grinding in liquid nitrogen (mortar and pestle), such as different animal and plant tissues
2. Glass-bead grinding, for example, oocysts (e.g. Eimeria spp.), metacercariae (e.g. Fasciola spp.) and nematodes’ eggs (e.g. eggs of Haemonchus contortus)
3. Repetitive pipetting, notable examples of it are animal cells and zoites of apicomplexan parasites, such as sporozoites, merozoites, tachyzoites and bradyzoites, and trypansomal forms of Trypanosoma spp. and Leishmania spp., for example, trypanomastigote, promastigote, amastigote and epimastigote.

In recent years, the development of molecular techniques has created a need for establishing simple and efficient novel methods of DNA and RNA extraction for PCR amplification and other related techniques. Carbohydrates, tannins, polyphenols and proteins in addition to hazardous organic solvents, such as phenol and chloroform are the major enemies of the embattled researchers. No existence for DNA or RNA extraction method that is suitable for all prokaryotic and eukaryotic organisms.

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Furthermore, there is an urgent need to address the insufficiency of reasonable environment for RNase to have DNA free of RNA and even for DNase to degrade the DNA.

2. Materials and methods

2.1. Reagents

Proteinase K, 100% Ethanol, 70% Ethanol, Double distilled (DD) water, Ethylene diamine tetra acetic acid (EDTA), RNase, DNase, Pyrex beads, Agarose, Deoxyribonucleic acid (DNA) Marker, 2 × EasyPfu PCR SuperMix, 10% Sodium dodecyl sulfate (SDS), Glacial acetic acid (CH₃COOH), Hydrochloric acid (HCl) and Sodium hydroxide (NaOH).

2.2. Equipments

Mortar, Pestle, PCR machine, Microscope, Refrigerated Benchtop centrifuge (MIKRO200R, Germany), Weighing scale, Pipettes (20, 100, and 1000 µl), 15 and 50 ml falcon tubes, 50 ml centrifuge tubes and Disposable Polypropylene micro-centrifuge tubes

2.3. Reagent setup

Tris buffer, Tris-EDTA (TE), DEPC-treated water, Saturated salt solution (NaCl), Neutral saturated salt solution, Acidic saturated salt solution and Lysis buffer: 1X STE buffer (50 mM NaCl, 50 mM Tris-HCl and 100 mM EDTA; PH 8.0)

2.4. Procedure

2.4.1. Grinding in liquid nitrogen (Mortar and pestle)

2.4.1.1. DNA extraction protocol. Hepatic DNA extraction from mouse can be divided into six steps. These are:

2.4.1.1.1. Homogenization. 1 g of the liver was taken and cut into pieces then ground using a porcelain mortar and pestle in 3 ml of lysis buffer containing 900 µl of 10% SDS. The emulsion was transferred to micro-centrifuge tubes and 100 µg proteinase K was added per ml of emulsion solution, and incubated for 1 h at 50 °C.

2.4.1.2. Phase separation. 350 µl of neutral saturated salt solution (NaCl) per ml was added to the previous emulsion, the micro-centrifuge tube was capped and shaken gently by hand for 15 s, and then incubated at room temperature for 10 min. The micro-centrifuge tube was centrifuged at 590 × g for 15 min at room temperature with DNA remaining exclusively in the aqueous phase (see Fig. 1A for illustration).

2.4.1.3. DNA precipitation. The resulting aqueous phase was transferred into another micro-centrifuge tube, and mixed with two volumes of room temperature absolute ethyl alcohol. Then the micro-centrifuge tube was inverted several times for 10 s.

2.4.1.4. DNA wash. The supernatant was removed; the DNA pellet was washed once with 75% ethanol, and the DNA was precipitated out of the solution by centrifugation at 9500 × g for 5 min.

2.4.1.5. DNA dissolving. The DNA pellet was allowed to dry for 5 min, and dissolved in DD water. Then the DNA was quantified and aliquoted to be stored at −20 °C.

2.4.1.6. Removal of RNA from DNA preparation. 50 µg per ml RNase was added and the mixture was incubated for 1 h at 37 °C.

- Critical step: The treatment of DNA with RNase should be done in Tris buffer at the end of the extraction protocol. Salting out step can be repeated as before according to the protocol to obtain DNA with highest quality. The DNA can be precipitated and washed with 70% ethanol, and then the pellet can be dissolved in Tris-EDTA (TE) for DNA protection from degradation by metal dependent nucleases during storage.

2.4.1.2. RNA extraction protocol. Hepatic RNA extraction method from mouse can be listed as follows:

2.4.1.2.1. Homogenization. 1 g of the liver was taken and cut into pieces then ground using a porcelain mortar and pestle in 3 ml of lysis buffer containing 900 µl of 10% SDS. The emulsion was transferred to micro-centrifuge tubes.

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**Fig. 1.** Nucleic acid extraction and downstream application.

A. Neutral salting out (DNA extraction).
B. Acidic salting out (RNA extraction).
C. GoldView™ Nucleic Acid Stained 1.5% Agarose gel demonstrating the integrity of total DNA extracted from Eimeria tenella.
D. GoldView™ Nucleic Acid Stained 1.5% Agarose gel of total RNA of E. tenella.
E. Standard PCR amplification of MICII of E. tenella.
F. GoldView™ Nucleic Acid Stained 1.5% Agarose gel of EtMIC-2 RT-PCR amplified product of E. tenella using Finnzymes phusion™ High-Fidelity DNA Polymerase.
2.4.1.2.2. **Phase separation.** 350 μl of acidic saturated salt solution (NaCl) was added into each tube of the previous emulsion mixture, and the micro-centrifuge tube was capped and gently shaken by hand for 15 s and then incubated at room temperature for 10 min. The micro-centrifuge tube was centrifuged at 590 × g for 15 min at room temperature with RNA remaining exclusively in the aqueous phase (see Fig. 1B for elucidation).

2.4.1.2.3. **RNA precipitation.** The resulting aqueous phase was transferred into micro-centrifuge tubes and precipitated by mixing the aqueous phase with two volumes of cold absolute ethyl alcohol. Then the micro-centrifuge tube was inverted several times for 15 s.

2.4.1.2.4. **RNA wash.** The supernatant was removed; and the RNA pellet was washed once with cold 75% ethanol, and the RNA was precipitated out of the solution by centrifugation at 9500 × g for 5 min.

2.4.1.2.5. **RNA dissolving.** The RNA pellet was allowed to dry for 5 min and dissolved in DEPC-treated water. Then the RNA was quantified, and aliquoted to be stored at −80°C.

2.4.1.2.6. **DNA removal.** DNase I (RNase-free) kit was employed to remove any contaminating DNA from the sample as per manufacturer’s instructions.

- Critical step: The treatment of RNA with DNase should be done in Tris buffer at the end of the extraction protocol. Salting out step can be repeated as before according to the protocol to obtain RNA with highest quality. The RNA can be precipitated and washed with ethanol, and then the pellet can be dissolved in DEPC-treated water or Tris-EDTA (TE) for DNA protection from degradation by metal-dependent nucleases during storage.

2.4.2. **Repetitive pipetting: prokaryotes, for example E. coli**

2.4.2.1. **DNA extraction protocol.** Bacterial DNA extraction can be listed as follows:

2.4.2.1.1. **Homogenization.** 1 ml microbial culture was transferred into micro-centrifuge tubes and pelleted by centrifugation at 380 × g for 5 min at room temperature. Then the supernatant was discarded, and the pellet was resuspended with repetitive pipetting in 1 ml lysis buffer containing 100 μl of 10% SDS and 100 μg proteinase K. The mixture was incubated for 1 h at 50°C.

2.4.2.1.2. **Phase separation, DNA precipitation, DNA wash, DNA dissolving and. RNA removal.** They were conducted as previously mentioned in Section 2.4.1.1.

2.4.2.2. **RNA extraction protocol.** Bacterial RNA extraction can be divided into six sections:

2.4.2.2.1. **Homogenization.** 1 ml microbial culture in microcentrifuge tube was pelletted by centrifugation at 380 × g for 5 min at 37°C and the supernatant was discarded. The pellet was resuspended with repetitive pipetting in 1 ml lysis buffer containing 100 μl of 10% SDS.

2.4.2.2.2. **Phase separation, RNA precipitation, RNA wash, RNA dissolving and DNA removal.** They were performed as we previously mentioned in Section 2.4.1.2.

2.4.3. **Glass-bead grinding**

2.4.3.1. **DNA extraction protocol.** Eimerian DNA extraction from oocysts can be divided into six steps. These are:

2.4.3.1.1. **Homogenization.** 5 × 10⁶ sporulating/sporulated oocysts, 0.5 g of Pyrex beads and up to 3 ml of lysis buffer containing 900 μl of 10% SDS and 300 μg proteinase K were added in a 15 ml disposable polypropylene tube. The mixture was incubated for 1 h at 50°C.

2.4.3.1.2. **Phase separation.** 1 ml of neutral saturated salt solution was added to the previous mixture, and the 15 ml disposable polypropylene tube was capped and gently shaken by hand for 15 s and then incubated at room temperature for 10 min. The 15 ml disposable polypropylene tube was centrifuged at 590 × g for 15 min at room temperature.

2.4.3.1.3. **DNA precipitation, DNA wash, DNA dissolving and removing contaminating RNA from DNA.** These steps were then performed as previously reported in Section 2.4.1.1.

![Fig. 2. GoldView™ Nucleic Acid Stained 1.5% Agarose gel demonstrating. A. The integrity of total DNA extracted from the liver of the BALB/c mice. B. GAPDH amplified product of mouse genome. Lane 1: Trans DNA Marker III, Lane 2 and 3: GAPDH amplified product from the genomic DNA of mouse genome.](image-url)
2.4.3.2. RNA extraction protocol. Eimerian RNA extraction can be divided into six sections:

2.4.3.2.1. Homogenization. $5 \times 10^6$ sporulating/sporulated oocysts, 0.5 g of Pyrex beads and up to 3 ml of lysis buffer containing 900 μl of 10% SDS were added in a 15 ml disposable polypropylene tube, and the mixture was incubated for 1 h at 42 °C.

2.4.3.2.2. Phase separation. 1 ml of acidic saturated salt solution was added to the previous mixture, and the 15 ml disposable polypropylene tube was capped and gently shaken by hand for 15 s and then incubated at room temperature for 10 min. The 15 ml disposable polypropylene tube was centrifuged at 590 × g for 15 min at room temperature.

2.4.3.2.3. RNA precipitation, RNA wash, RNA dissolving and DNA removal. They were conducted as we previously reported in Section 2.4.1.2.

3. Results and discussion

3.1. Eukaryotes, such as

3.1.1. Eimeria spp.

3.1.1.1. DNA isolation. The average DNA purity $A_{260}/A_{280}$ ratio was 1.87 ± 0.025 of three repeated samples, which indicates low amounts of contaminants in the samples. The quantity of DNA extracted from $6 \times 10^6$ sporulated eimerian oocysts was 3.5985 ± 0.27 mg with integrity validated by 1.5% Agarose gel electrophoresis (Fig. 1C).

3.1.1.2. RNA isolation. The average RNA purity $A_{260}/A_{280}$ ratio was 1.42 ± 0.035 with yield average of 191.7 ± 1.7 μg/6 × 10⁶ unsporulated eimerian oocysts. As can be seen from Fig. 1D, the integrity of isolated total RNA was confirmed by 1.5% Agarose gel electrophoresis.

3.1.1.3. Assessing DNA and RNA for downstream applications. The ability to amplify a specific target from extracted DNA and RNA was proved using a pair of precise primers; MIC2-UP TATGCTCGACGGTTGTCCGGT and MIC2-D GTCAGGATGACTGTAAGTGC that were designed from the published Eimeria tenella microneme (MIC2) mRNA sequence (ACCESSION FJ807654) (Fig. 1E). The primers were synthesized by AuGt Biotechnology Synthesis Lab, Beijing, China. Fig. 1F presents RT-PCR amplification of MICII of E. tenella.

3.1.2. BALB/c mice

3.1.2.1. DNA isolation. The average $A_{260}/A_{280}$ ratio and yield of the total DNA extracted were 1.83 ± 0.025 and 20 ± 0.45 μg/g of liver tissue, respectively. The integrity of isolated total DNA was confirmed by 1.5% Agarose gel electrophoresis as shown in Fig. 2A.

3.1.2.2. Assessing DNA for downstream applications. The ability to amplify a specific target, such as 16S Ribosomal DNA from extracted DNA was proved using a pair of precise primers; forward GAPDH Primer, 5′-CAAGGTCTACCATGACAACCTT-3′ and the reverse GAPDH Primer, 5′-GTCACACCCCCTGTGCTAG-3′ that were provided with RevertAid™ First Strand cDNA Synthesis Kit to test the control sample (Fig. 2B).

### Fig. 3. GoldView™ Nucleic Acid Stained 1.5% Agarose gel illustrating.

A. The integrity of total DNA extracted from the Escherichia coli bacterial cultures, Lane 1: λ-EcoRI digest DNA Marker, Lane 2 Genomic DNA of the Escherichia coli. B. 16S ribosomal RNA sequence amplified product of E. coli genome. Lane 1: Trans DNA Marker III, Lane 2 16S ribosomal RNA amplified product from the genomic DNA of E. coli genome. C. Total RNA of E. coli. Lane 1 Trans DNA Marker 1Kb DNA ladder, Lane 2 total RNA of E. coli.

### Fig. 4. Electropherogram and gel image output (virtual gel) from Agilent Bioanalyzer’s Haemonchus contortus total RNA nano assay. The x-axis represents fluorescence unit (FU) and y-axis represents time (seconds).
Table 1
Test results of Agilant 2100.

| Parameters                  | Average ± standard deviation (SD) |
|-----------------------------|-----------------------------------|
| RNA concentration           | 164.7 ± 13.6 μg/g                 |
| RNA integrity number (RIN)  | 7.27 ± 0.32                       |

* Each value is a mean of three separate samples.

3.2. Prokaryotic examples; E. coli

3.2.1. DNA isolation

The average DNA purity ratio A_{260/A_{280}} was 1.87 ± 0.065 with DNA yield average of 48 ± 2.24 μg/1 ml (1 × 10^8 cells) of *Escherichia coli* bacterial cultures that were grown overnight in Luria Broth (LB) at 37 °C. The integrity of isolated total DNA was confirmed by 1.5% Agarose gel electrophoresis as presented in Fig. 3A.

3.2.2. Assessing DNA for downstream applications

The ability to amplify a specific target, such as the *E. coli* 16S ribosomal RNA sequence from extracted DNA was proved using a pair of precise primers that were designed from the published *E. coli* 16S ribosomal RNA sequence (ACCESSION NO: J01859/K02555/M24828/M24833/M24834/M24835/M24836/M24837/M24911/M24996) (Fig. 3B). The primers were synthesized by AuGCT Biotechnology Synthesis Lab, Beijing, China.

3.2.3. RNA isolation

The average purity of RNA samples was A_{260/A_{280}} ratio was 1.99 ± 0.01 and the quantity of RNA extracted from 1 × 10^8 *E. coli* was 22 ± 1.45 μg with integrity confirmed by 1.5% Agarose gel electrophoresis as can be seen from Fig. 3C.

3.3. Further assessment of RNA purity and integrity using *trichostongylid adult worm*

Total RNA from the barber’s pole worm, *Haemonchus contortus* was isolated employing salting out (acidic condition) followed by DNase digestion. For contaminant detection, the A_{260/A_{280}} and A_{260/A_{230}} values (i.e., for detection of protein contaminants and residual chemical contamination, such as EDTA and SDS) were 2 ± 0.02 and 2.076 ± 0.024, respectively [1–3]. Additionally, Agilent Bioanalyzer 2100 system (Agilent Technologies, CA, USA) was used in conjunction with the traditional 1.5% agarose gels for RNA integrity assessment (Fig. 4, Table 1).

These results present a simplified, semi-unified, effective, and toxic material free protocol for extracting DNA and RNA from different prokaryotic and eukaryotic sources exploiting the physical properties of the negatively charged molecules; DNA and RNA. The positively ions of saturated salt solution neutralize the negatively charged phosphate groups of the DNA and RNA backbone. Furthermore, in neutral saturated salt conditions, DNA will remain in the aqueous layer. However, RNA will partition into the aqueous layer by carrying out acidic saturated salt solution extraction.

Yield and quality are the ultimate goal for any researchers during DNA extraction procedure. Doubtless, the quality increases by getting RNA free of DNA contamination. Previous published studies failed to resolve this issue [4–12], and (http://www.natureprotocols.com/2009/05/27/a_protocol_for_high_molecular.php, http://www.nwsc.noaa.gov/publications/techmemos/tm14/dnaisol.html, http://csm.jmu.edu/biology/courses/bio480_580/mblab/genomic2.htm, www.promega.com).

The most common protocols used the chelating agent, ethylenediaminetetraacetic acid (EDTA), sodium dodecyl sulfate (SDS) as a detergent, and sodium chloride as a stabilizer in the lysis

Fig. 5. Diagrams illustrate the negatively charged phosphate (PO₄³⁻) of DNA and RNA molecules and the positively charged monovalent sodium cation shield around the helix axis.
A. Negatively charged phosphate group in the ribonucleic acid and deoxyribonucleic acid backbone.
B. Axial view of DNA with sodium cation shield modified from Berg et al., 2002.
buffer. The high affinity of EDTA to divalent cations, such as Ca\(^{2+}\), Mn\(^{2+}\) and Mg\(^{2+}\), which act as cofactors for nucleases could inhibit the degradation of DNA and RNA by DNases and RNases respectively [13–15]. SDS is an anionic detergent for cell and nucleus lysis to release ribonucleic and deoxyribonucleic acids. The nucleases; ribonuclease (RNase) and deoxyribonuclease (DNase) activities were inhibited by SDS [16,17]. The electrostatic repulsion between the two negatively charged helix strands destabilizing the helix was counteracted by positively charged sodium chloride [18]. The shielding effect of monovalent sodium cations leads to DNA and RNA stabilization through neutralization of the negative charge on the sugar phosphate backbone as is demonstrated in Fig. 5 [19].

Elevated salt concentration, SDS and EDTA were used to inhibit nuclease activity during extraction of DNA from tissues or organisms with high nucleic activity [20]. The use of sodium chloride in the lysis buffer decreases the susceptibility of DNA and RNA to be attacked by the action of nucleases possibly due to steric hindrance. Additionally, salting out denatures proteins and leaves nucleic acids intact. This is the most potent way of expeditiously inactivating nucleases.

We pointed out that DNA and RNA are under triple protection (i.e. EDTA, SDS and NaCl) and this environment is unsuitable for RNase to get DNA free of RNA and even for DNase to degrade DNA. Our conclusion is supported by results from treatments of different prokaryotic and eukaryotic sources as illustrated in Figs. 6A–C. The complete removal of RNA under the effect of RNase is achieved when RNase is eventually added (i.e. in Tris buffer without EDTA), which gives optimal quality with any DNA extraction protocols.

The polar phosphate groups of DNA and RNA can electrostatically interact with the polar environment allowing them to be easily dissolved in Tris buffer; therefore, the treatment of DNA and RNA with RNase and DNase respectively should be done in Tris buffer at the end of the extraction protocol as a standard measure. Salting out step can be repeated as before according to the protocol to obtain DNA with the highest quality without major changes in the nucleic acid yield. The DNA can be precipitated and washed with ethanol, and then the pellet can be dissolved in Tris-EDTA (TE) for DNA and RNA protection from degradation by divalent-metal-dependent nucleases during storage.

4. Conclusion

A simplified, semi-unified, effective, and toxic material free protocol for extracting DNA and RNA utilizing the physicochemical properties of nucleic acids has been described. Moreover, the unsuitable environment for endonucleases, such as RNase and DNase to have DNA liberated of RNA and even for DNase to degrade the DNA respectively has been addressed, and an appropriate alternative protocol has been presented that could be the top-first in the field of molecular biology.

Conflict of interest

The authors declare that there are not conflicts of interest.

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