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For Research Use Only



INSTRUCTION

MANUAL

DNA/RNA/Protein Extraction Kit

IB47700 (4 Preparation Sample Kit)

IB47701 (50 Preparation Kit)

IB47702 (100 Preparation Kit)

Advantages

Sample: cultured animal cells (up to 5×10^6), up to 25 mg of tissue, up to 500 μl of whole human blood, up to 200 μl of biological liquids (serum, plasma)

Yield: up to 9 μg of genomic DNA, 20 μg of total RNA, 120 μg of protein from 1.5×10^6 HeLa cells

Format: genomic DNA spin column and total RNA spin column

Operation Time: DNA/RNA purification within 25 minutes, protein precipitation within 50 minutes

Elution Volume: 50-200 μl (genomic DNA) / 25-50 μl (total RNA)

Kit Storage: dry at room temperature (15-25°C)

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Introduction

The DNA/RNA/Protein Extraction Kit provides an efficient method for purifying genomic DNA, total RNA and total protein simultaneously from a variety of samples (cultured cells, animal tissue, whole blood and biological liquids). Chaotropic salt is used to lyse cells and inactivate DNases and RNases, allowing DNA to bind to the genomic DNA spin column. The flow-through can then be transferred to the RNA spin column for RNA binding. The proteins in the flow-through can then be precipitated using acetone. Contaminants are effectively removed using wash buffers followed by pure genomic DNA elution in a low salt buffer and pure total RNA elution in RNase-free Water. DNA/RNA purification can be completed in 25 minutes without phenol/chloroform extraction or alcohol precipitation and protein purification can be completed in 50 minutes. The purified DNA, with approximately 20-30 Kb, is suitable for use in PCR or other enzymatic reactions and the purified RNA (including miRNA) is ready for use in RT-PCR, Real-time PCR, northern blotting, primer extension, mRNA selection and cDNA synthesis. The purified proteins can be directly analyzed on a SDS-PAGE and subsequent western blot.

Quality Control

The quality of the DNA/RNA/Protein Extraction Kit is tested on a lot-to-lot basis by isolating genomic DNA, total RNA and protein from cultured animal cells. The purified DNA and total RNA is quantified with a spectrophotometer and analyzed by electrophoresis on a 1% agarose gel. The purified protein is quantified by Bradford assay and analyzed on SDS-PAGE.

Kit Components

Content	IB47700	IB47701	IB47702
RBC Lysis Buffer	10 ml	100 ml	200 ml
DR Buffer	2 ml	30 ml	60 ml
RW1 Buffer	2 ml	30 ml	50 ml
RPE Buffer ¹ (Add Ethanol)	1 ml (4ml)	25 ml (100 ml)	50 ml (200 ml)
W1 Buffer	2 ml	45 ml	45 ml
Wash Buffer ¹ (Add Ethanol)	1 ml (4ml)	12.5 ml (50 ml)	25 ml (100 ml)
RNase-free Water	1 ml	6 ml	6 ml
Elution Buffer	1 ml	30 ml	30 ml
DV Buffer (8M Urea)	500 µl	6 ml	12 ml
Protein Loading Dye	60 µl	2 ml	2 ml
RB Column	4	50	100
GD Column	4	50	100
2 ml Collection Tube	16	200	400

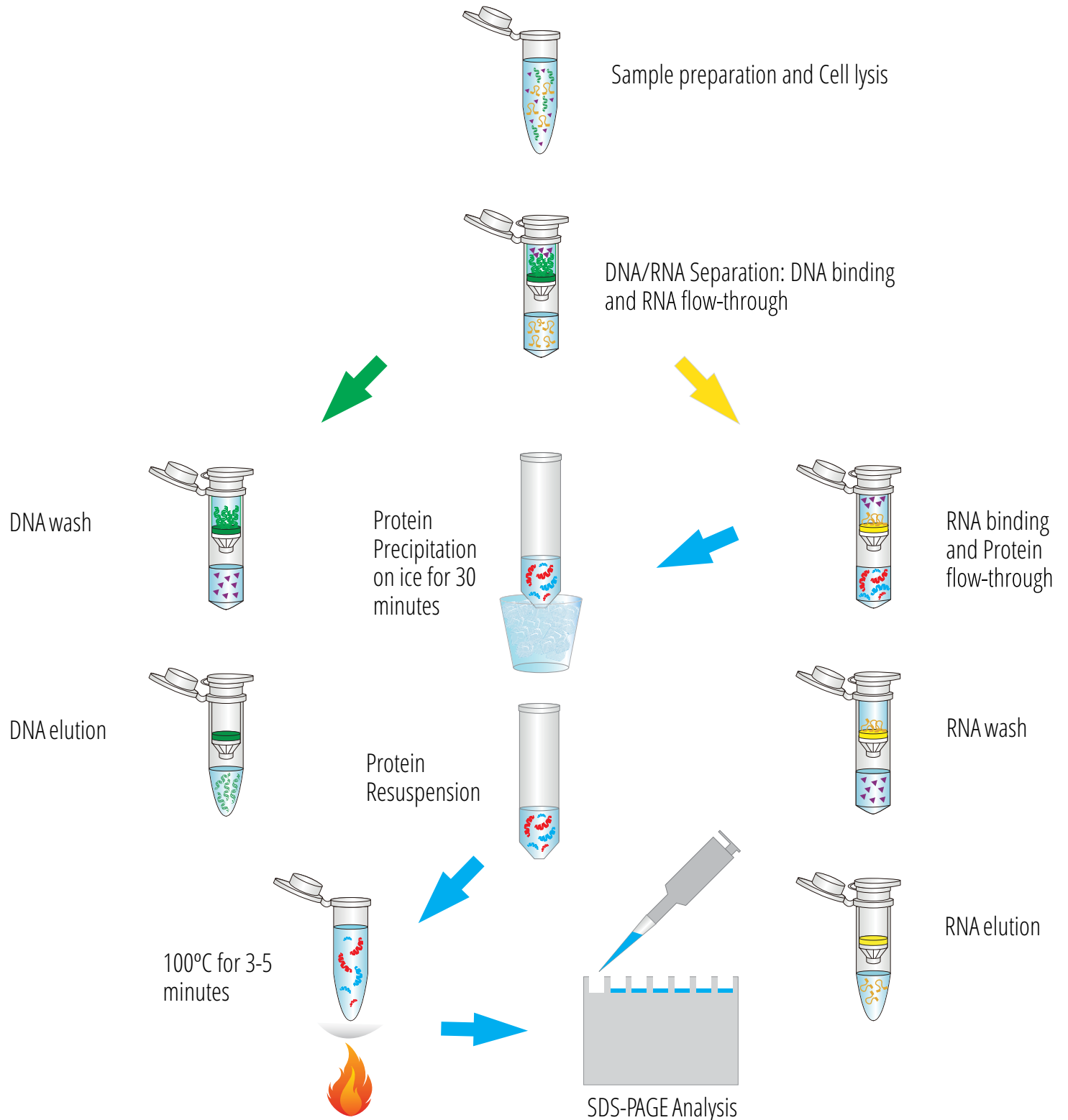
¹ Add absolute ethanol (see the bottle label for volume) to RPE Buffer and Wash Buffer then mix by shaking for a few seconds. Check the box on the bottle. Be sure and close the bottle tightly after each use to avoid ethanol evaporation.

Steps to prevent RNase contamination

1. Always wear a lab coat, disposable gloves, protective goggles and (anti-fog) procedure mask.
2. Plasticware and automatic pipettes should be sterile (RNase-free) and used only for RNA procedures.
3. Non-disposable glassware or plasticware should also be sterile (RNase-free).

 During the procedure, always wear a lab coat, disposable gloves, and protective goggles

Quick Protocol Diagram



DNA/RNA/Protein Extraction Kit Protocol

Please read the entire instruction manual prior to starting the Protocol Procedure.

IMPORTANT BEFORE USE!

1. Add absolute ethanol (see the bottle label for volume) to RPE Buffer and Wash Buffer then mix by shaking for a few seconds. Check the box on the bottle. Be sure and close the bottle tightly after each use to avoid ethanol evaporation.
2. Prepare Phosphate Buffered Saline (PBS, pH7.2) for adherent cultured cells.
3. Yield and quality of DNA/RNA will be higher when fresh samples or samples which have been flash frozen and stored at -70°C are used. DNA/RNA in samples which has been repeatedly frozen and thawed may be degraded.

Additional Requirements:

Absolute ethanol, β -mercaptoethanol, acetone; for cell samples: phosphate-buffered saline (PBS), 0.10-0.25% Trypsin; for tissue samples: TissueLyser or mortar and pestle, 20-G needle syringe.

Protocol Procedure

1. Sample Preparation.

Adherent Cultured Animal Cells (trypsinize cells prior to harvesting).

Remove the culture medium and wash cells in PBS. Aspirate PBS and add 0.10-0.25% Trypsin in PBS. Once cells detach add medium then transfer to a 15 ml centrifuge tube. Proceed with Suspension Cultured Animal cells.

Suspension Cultured Animal Cells.

Transfer cells (up to 5×10^6) to a 1.5 ml microcentrifuge tube then centrifuge for 5 minutes at 300 x g. Remove the supernatant completely then add 400 μ l of DR Buffer and 4 μ l of β -mercaptoethanol to resuspend cells by pipetting or vortex. Proceed with Step 2 DNA/RNA Separation.

Animal Tissue.

Excise 10-25 mg of tissue directly from the animal or remove the tissue sample from storage. Do not use more than 25 mg of tissue per reaction. Homogenize tissue samples using one of the following methods: A. Transfer the tissue to a 2 ml centrifuge tube containing ceramic beads or stainless steel beads, add 400 μ l of DR Buffer and 4 μ l of β -mercaptoethanol to the tissue sample and then homogenize the sample with a TissueLyser, Disruptor Genie or similar. B. Freeze the tissue in liquid nitrogen then grind the tissue thoroughly with a mortar and pestle. Transfer the tissue powder to a 1.5 ml microcentrifuge tube (do not allow the tissue to thaw) and add 400 μ l of DR Buffer and 4 μ l of β -mercaptoethanol. Shear the tissue by passing the lysate through a 20-G needle syringe 10 times. Proceed with Step 2 DNA/RNA Separation.

Biological Fluids.

Add 300 μ l of DR Buffer and 3 μ l of β -mercaptoethanol to 100 μ l of liquid sample and mix well by pipetting or vortex. Proceed with Step 2 DNA/RNA Separation.

Human Whole Blood.

Collect fresh human blood in anticoagulant-treated collection tubes. Transfer 500 μ l of blood to a sterile 15 ml centrifuge tube. Add 1.5 ml of RBC Lysis Buffer (3 volumes) and mix by inversion. Incubate the tube on ice for 10 minutes (vortex twice briefly during incubation). Centrifuge at 3,000 x g for 5 minutes then remove the supernatant completely. Add 400 μ l of DR Buffer and 4 μ l of β -mercaptoethanol to resuspend leukocyte pellet by pipetting or vortex. Proceed with Step 2 DNA/RNA Separation.

2. DNA/RNA Separation.

Incubate the sample lysate at room temperature for 5 minutes then centrifuge at 12-16,000 x g for 2 minutes. Place a GD Column in a 2 ml Collection Tube then transfer the supernatant to the GD Column. Centrifuge at 14-16,000 x g for 1 minute. Note: If the lysate mixture could not flow past the GD Column membrane following centrifugation, increase the centrifuge time until it passes completely. Save the flow-through in the 2 ml Collection Tube for RNA Purification. Place the GD Column in a new 2 ml Collection Tube and store at room temperature (15-25°C) or 4°C for DNA Purification. Do not store the GD Column for extended periods. Do not freeze the GD Column. At this time, preheat the required Elution Buffer (200 µl per sample) to 60°C (for DNA elution).

RNA Purification

3. RNA Binding

Add 0.8 volume of absolute ethanol to the flow-through in the 2 ml Collection Tube (e.g. add 320 µl of absolute ethanol to 400 µl of flow-through) and mix well by pipetting. Transfer the sample to the RB Column in a 2 ml Collection Tube. Centrifuge at 14-16,000 x g for 1 minute. Place the RB Column in a new 2 ml Collection Tube and save the flow-through for protein purification. Note: If DNA-free RNA is required, perform optional In Column DNase I Digestion below.

Optional In Column DNase I Digestion

Add 400 µl of RPE Buffer (make sure ethanol was added) to the RB Column then centrifuge at 14-16,000 x g for 30 seconds. Discard the flow-through and place the RB Column back in the 2 ml Collection Tube. Prepare DNase I solution in a 1.5 ml microcentrifuge tube (RNase-free) as follows:

D Nase I	5 µl (2 U/µl)
D Nase I Reaction Buffer	45 µl
Total Volume	50 µl

RNase-Free DNase I set can be purchased directly from IBI. Standard DNase buffers are incompatible with this in-column DNase digestion, which will effect RNA integrity and reduce yield.

Mix DNase I solution by pipetting gently (DO NOT vortex). Add DNase I solution (50 µl) into the CENTER of the RB column matrix. Incubate the column for 15 minutes at room temperature (20-30°C) then proceed with RNA Wash on page 6.

4. RNA Wash

Add 400 µl of RW1 Buffer into the RB Column and centrifuge at 14-16,000 x g for 30 seconds. Discard the flow-through and place the RB Column back in the 2 ml Collection Tube. Add 600 µl of RPE Buffer (make sure ethanol was added) into the RB Column. Centrifuge at 14-16,000 x g for 30 seconds. Discard the flow-through and place the RB Column back in the 2 ml Collection Tube. Add 600 µl of RPE Buffer (make sure ethanol was added) into the RB Column. Centrifuge at 14-16,000 x g for 30 seconds. Discard the flow-through and place the RB Column back in the 2 ml Collection Tube. Centrifuge at 14-16,000 x g for 3 minutes to dry the column matrix.

5. RNA Elution

Place the RB Column in a clean 1.5 ml microcentrifuge tube (RNase-free). Add 25-50 µl of RNase-free Water into the CENTER of the column matrix. Let stand for at least 1 minute to ensure the RNase-free Water is absorbed by the matrix. Centrifuge at 14-16,000 x g for 1 minute to elute the purified RNA.

Protein Purification

6. Protein Precipitation

Add 4 volumes of ice-cold acetone to the flow-through from step 3 in a 15 ml centrifuge tube (example: add 2.8 ml of ice-cold acetone to 700 µl flow-through). Incubate on ice or at -20°C for 30 minutes. At this time, perform DNA Purification on page 7. Centrifuge at 14-16,000 x g for 10 minutes then discard the supernatant.

7. Protein Resuspension

Add 100 μ l of ice-cold 70% ethanol to wash the protein pellet. Discard the supernatant then air-dry the protein pellet at room temperature.

NOTE! DO NOT overdry the protein pellet as it may result in difficult resuspension. Add up to 100 μ l of DV buffer (8M urea) or a buffer compatible with downstream application to dissolve the protein pellet.

8. SDS-PAGE Analysis

Add 2 μ l of Protein Loading Dye and 0.5 μ l of 2M DTT (optional) into a clean microcentrifuge tube. Add 8 μ l of the protein sample then mix well by pipetting. Incubate protein sample at 100°C for 3-5 minutes. Centrifuge the protein sample briefly then directly load to SDS-PAGE.

DNA Purification

9. DNA Wash

Add 400 μ l of W1 Buffer to the GD Column. Centrifuge at 14-16,000 x g for 30 seconds then discard the flow-through. Place the GD Column back in the 2 ml Collection Tube. Add 600 μ l of Wash Buffer (make sure absolute ethanol was added) to the GD Column. Centrifuge at 14-16,000 x g for 30 seconds then discard the flow-through. Place the GD Column back in the 2 ml Collection Tube. Centrifuge again for 3 minutes at 14-16,000 x g to dry the column matrix.

NOTE! Additional centrifugation at 14-16,000 x g for 5 minutes or incubation at 60°C for 5 minutes will completely dry the GD Column to avoid any residual ethanol carryover and ensure the most effective downstream applications.

10. DNA Elution

Standard elution volume is 100 μ l. If less sample is to be used, reduce the elution volume (30-50 μ l) to increase DNA concentration. If higher DNA yield is required, repeat the DNA Elution step to increase DNA recovery and the total elution volume to approximately 200 μ l.

Transfer the dried GD Column to a clean 1.5 ml microcentrifuge tube. Add 100 μ l of pre-heated Elution Buffer¹, TE Buffer² or water³ into the CENTER of the column matrix. Let stand for at least 3 minutes to allow Elution Buffer, TE Buffer or water to be completely absorbed. Centrifuge at 14-16,000 x g for 30 seconds to elute the purified DNA.

¹ Ensure that Elution Buffer (10 mM Tris-HCl, pH8.5 at 25°C) is added into the center of the GD Column matrix and is completely absorbed.

² Using TE (10 mM Tris-HCl, 1 mM EDTA, pH8.0) for elution is beneficial as EDTA preserves DNA for long term storage. However, EDTA will affect PCR and other sensitive downstream applications. Ensure that TE is added into the center of the GD Column matrix and is completely absorbed.

³ If using water for elution, ensure the water pH is between 7.0 and 8.5. ddH₂O should be fresh as ambient CO₂ can quickly cause acidification. Ensure that water is added into the center of the GD Column matrix and is completely absorbed. DNA Eluted in water should be stored at -20°C to avoid degradation.

Troubleshooting

Low Yield

Improper sample homogenization.

Yield and quality of DNA/RNA will be higher when fresh samples or samples which have been flash frozen and stored at -70°C are used. DNA/RNA in samples which has been repeatedly frozen and thawed may be degraded. Fresh blood is recommended. However, frozen or blood treated with anticoagulants can also be used. Increased storage length decreases DNA/RNA yield. Overloading the columns causes low nucleic acid yield.

Incomplete buffer preparation.

Add absolute ethanol (see the bottle label for volume) to RPE Buffer and Wash Buffer then mix by shaking for a few seconds. Check the box on the bottle. Be sure and close the bottle tightly after each use to avoid ethanol evaporation. Pre-heat Elution Buffer to 60°C prior to DNA elution.

Incorrect elution step.

Ensure that Elution Buffer, TE or water is added into the CENTER of the GD Column matrix and is completely absorbed. Make sure RNase-free Water is added to the CENTER of the RB Column and is absorbed completely. Use pre-heated Elution Buffer, TE, or water (60–70°C). If using water for elution, ensure the water pH is between 7.0 and 8.5. ddH₂O should be fresh as ambient CO₂ can quickly cause acidification. Elute twice to increase the DNA/RNA recovery.

Clogged column.

Use the recommended amount of starting material or separate into multiple tubes. After homogenization, centrifuge the lysate at 16,000 x g for 5 minutes to precipitate insoluble cell debris, make sure only supernatant was transferred to GD Column. All centrifugation steps should be at room temperature (20-25°C).

DNA Contaminated With RNA

Lysate applied to GD Column contains ethanol.

Only add the appropriate volume of ethanol to the lysate after passing lysate through the GD Column.

Low A260/280

Improper buffer preparation.

Add appropriate volume of absolute ethanol (see the bottle label) to the RPE Buffer and Wash Buffer prior to use. Incorrect buffer used for nucleic acid dilution. Use Elution Buffer (10 mM Tris, pH=8.0) instead of RNase-free Water for DNA/RNA dilution before measuring purity.

Degraded RNA

Incorrect sample preparation and/or storage.

Process or freeze samples at -70°C immediately after collection.

Incorrect storage temperature.

Extracted RNA should be stored at -70°C.

Eluted DNA/RNA Does Not Perform Well In Downstream Applications

Residual Ethanol Contamination.

Following the wash step, dry the GD/RB Column with additional centrifugation at 14-16,000 x g for 5 minutes to remove residual ethanol.

No Protein Detected On Western Blot Or Coomassie Stained Gel Protein pellet loss.

At the protein precipitation step, the protein pellet is loosely attached to the side of the 15 ml centrifuge tube. Carefully decant the supernatant.

DNA/RNA/Protein Extraction Kit

Functional Test Data

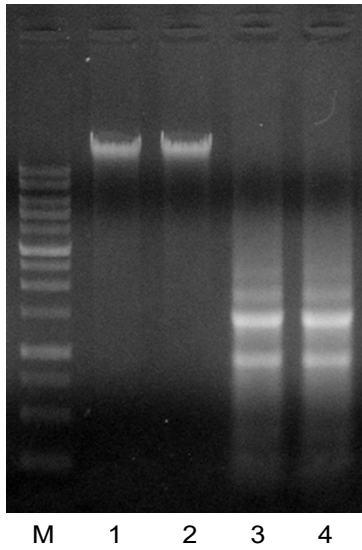


Figure 1. Genomic DNA and Total RNA from 1.5×10^6 HeLa cells was extracted using the DNA/RNA/Protein Extraction Kit. 10 μ l aliquots from a 200 μ l eluate of purified genomic DNA and 10 μ l aliquots from a 50 μ l eluate of purified total RNA were analyzed by electrophoresis on a 1% agarose gel.

1-2 = DNA from 1.5×10^6 HeLa cells

3-4 = RNA from 1.5×10^6 HeLa cells

M = 1 Kb DNA Ladder

Sample	μ g/ml	260/280	260/230	Yield
1. DNA	42.7	1.88	2.19	8.54 μ g
2. DNA	46.7	1.88	2.18	9.34 μ g
3. RNA	417.5	2.09	2.03	20.90 μ g
4. RNA	429.9	2.09	2.04	21.50 μ g

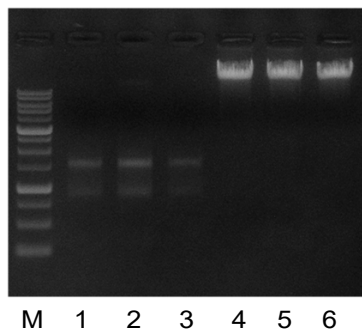


Figure 2. Genomic DNA and Total RNA from 3 human blood samples (500 μ l) was extracted using the DNA/RNA/Protein Extraction Kit. DNA yield: 8.0-10.0 μ g (100 μ l eluate, sample 4-6) RNA yield: 0.5-1.0 μ g (50 μ l eluate, sample 1-3). 10 μ l aliquots from a 100 μ l eluate of purified genomic DNA and 10 μ l aliquots from a 50 μ l eluate of purified total RNA were analyzed by electrophoresis on a 1% agarose gel.

M = 1 Kb DNA Ladder

Sample	ng/ μ l	260/280	260/230	Yield (μ g)
1. RNA	13.1	1.84	1.91	0.7
2. RNA	20.8	1.93	1.97	1.0
3. RNA	12.1	1.89	1.89	0.6
4. DNA	93.4	1.83	2.18	9.3
5. DNA	103.1	1.85	2.12	10.3
6. DNA	89.1	1.83	2.16	8.9

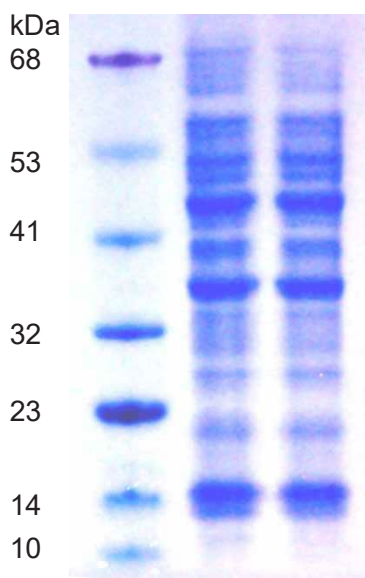


Figure 3. Protein was extracted from 1.5×10^6 HeLa Cells using the DNA/RNA/Protein Extraction Kit. 20 μ l of aliquots from a 200 μ l eluate of purified protein were analyzed on a NuPAGE4-12% Bis-Tris gel and stained with coomassie blue.

Related DNA/RNA Extraction Products

RNA Purification		
Product	Package Size	Catalogue Number
Total RNA Mini Kit (Blood/Cultured Cell)	50/100/300 preps	IB47321/322/323
Total RNA Mini Kit (Tissue)	50/100/300 preps	IB47301/302/303
Total RNA Mini Kit (Plant)	50/100/300 preps	IB47341/342/343
rBAC Mini RNA Bacteria Kit	100/300 preps	IB47421/422
rYeast Total RNA Mini Kit	100/300 preps	IB47411/412
miRNA Isolation Kit	100 preps	IB47371
IBI Isolate	100/200 rxns	IB47601/602
IBI Tri-Isolate	100/200 rxns	IB47631/632
RNA Pure Kit	50/100 rxns	IB47641/642
Virus DNA/RNA Purification		
Product	Package Size	Catalogue Number
Viral Nucleic Acid Extraction Kit II	50/100/300 preps	IB47401/402/403
Plasmid DNA Purification		
Product	Package Size	Catalogue Number
I-Blue Mini Plasmid Kit	100/300 preps	IB47171/172
I-Blue Midi Plasmid Kit	25 preps	IB47181
I-Blue Midi Plasmid Kit (Endotoxin Free)	25 preps	IB47191
Fast Ion Plasmid Midi Kit	25 preps	IB47111
Fast Ion Plasmid Midi Kit (Endotoxin Free)	25 preps	IB47113
Fast Ion Plasmid Maxi Kit	10/25 preps	IB47121/122
Fast Ion Plasmid Maxi Kit (Endotoxin Free)	10/25 preps	IB47124/125
96-Well Plasmid Kit	4/10 x 96 preps	IB47151/152
Post Reaction DNA Purification		
Product	Package Size	Catalogue Number
Gel/PCR DNA Fragments Extraction Kit	100/300 preps	IB47020/030
Small DNA Fragments Extraction Kit	100/300 preps	IB47061/062
96-Well Gel/PCR DNA Extraction Kit	4/10 x 96 preps	IB47040/050
Genomic DNA Purification		
Product	Package Size	Catalogue Number
Genomic DNA Mini Kit (Blood/Cultured Cell)	100/300 preps	IB47201/202
Genomic DNA Maxi Kit (Blood/Cultured Cell)	10 preps	IB47210
Genomic DNA Mini Kit (Tissue)	50/300 preps	IB47221/222
gMax Mini Kit (Blood/Tissue)	100/300 preps	IB47281/282
Genomic DNA Mini Kit (Plant)	100 preps	IB47230
Genomic DNA Maxi Kit (Plant)	10/25 preps	IB47240/241
gBAC Mini DNA Bacteria Kit	100/300 preps	IB47291/292
gYEAST Genomic DNA Kit	100/300 preps	IB47266/267
96-Well Genomic DNA Extraction Kit	4/10 x 96 preps	IB47251/252
96-Well Genomic DNA Extraction Kit (Plant)	4/10 x 96 preps	IB47271/272
96-Well Genomic DNA Bacteria Kit	4/10 x 96 preps	IB47296/297
IBI Plant Isolate	100/200 rxns	IB47611/612
DNA/RNA/Protein Purification		
Product	Package Size	Catalogue Number
DNA/RNA/Protein Extraction Kit	50/100 preps	IB47701/702

For additional product information please visit www.ibisci.com. Thank you



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