# gYEAST Genomic DNA Kit

For research use only

**Catalogue Numbers** 

IB47265 IB47266 IB47267



# Introduction

The gYEAST Genomic DNA Kit offers a simple and gentle reagent DNA precipitation method for isolating high molecular weight genomic, mitochondrial or viral DNA from *Saccharomyces cerevisiae* and a variety of other yeast and fungus species. This highly versatile solution based system offers a convenient procedure with minimal hands on time. The provided Sorbitol Buffer, when combined with zymolase or lyticase, will efficiently lyse yeast and other fungus species cell walls consisting of chitin and polysaccharides. The extracted DNA (A260/A280 = 1.8-2.0), is suitable for use in PCR or other enzymatic reactions.

# **Quality Control**

The gYEAST Genomic DNA Kit is tested on a lot-to-lot basis. Saccharomyces cerevisiae ( $2 \times 10^8$ ) is harvested by centrifugation at 5,000 x g for 10 minutes. A 15 µl aliquot of purified genomic DNA from a 100 µl eluate is analyzed by electrophoresis on a 1% agarose gel.

## Advantages

• High molecular weight genomic DNA extraction from a variety of yeast/fungus samples using a simple and gentle DNA precipitation method

- Sample: up to 2  $\times$  10<sup>8</sup> yeast and other fungus species
- Convenient: includes premixed Sorbitol Buffer
- Format: DNA precipitation reagent system
- Elution volume: 50-100 µl

# Applications

PCR, AFLP, RFLP/PADP, Southern Blotting, Real-time PCR

## Caution

The gYEAST Genomic DNA Kit contains irritants. During operation, always wear a lab coat, disposable gloves, protective goggles and (anti-fog) procedure mask.

## **Additional Requirements**

1.5 ml microcentrifuge tubes, zymolase or lyticase, RNase A (50 mg/ml), isopropanol, absolute ethanol for preparing 70% ethanol in  $ddH_2O$ 

## **Components and Storage**

Using DNA Hydration Buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0) is beneficial as EDTA preserves DNA for long term storage. However, EDTA will affect PCR and other sensitive downstream applications. If using water instead of DNA Hydration Buffer, ensure the water pH is between 7.0 and 8.5.  $ddH_2O$  should be fresh as ambient  $CO_2$  can quickly cause acidification. DNA in water should be stored at -20°C to avoid degradation.

| Item   | Volume | Product | Shipping         | Storage                           |
|--|--------|---------|------------------|-----------------------------------|
| Sorbitol Buffer  | 4.5 ml | IB47265 | room temperature | dry at room temperature (15-25°C) |
|  | 90 ml  | IB47266 |                  | for up to 1 year                  |
|  | 225 ml | IB47267 |                  |                                   |
| Cell Lysis Buffer  | 3 ml   | IB47265 | room temperature | dry at room temperature (15-25°C) |
|  | 40 ml  | IB47266 |                  | for up to 1 year                  |
|  | 100 ml | IB47267 |                  |                                   |
| Protein Removal Buffer                                     | 1 ml   | IB47265 | room temperature | dry at room temperature (15-25°C) |
|  | 15 ml  | IB47266 |                  | for up to 1 year                  |
|  | 40 ml  | IB47267 |                  |                                   |
| DNA Hydration Buffer<br>(10 mM Tris-HCl, 1 mM EDTA, pH8.0) | 1 ml   | IB47265 | room temperature | dry at room temperature (15-25°C) |
|  | 50 ml  | IB47266 |                  | for up to 1 year                  |
|  | 50 ml  | IB47267 |                  |                                   |

# Yeast and other Fungus Species Protocol Procedure

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

#### 1. Cell Harvesting

- 1. Transfer fungus cells (up to 2 x 10<sup>8</sup>) to a 1.5 ml microcentrifuge tube.
- 2. Harvest fungus cells by centrifugation for 10 minutes at 5,000 x g.
- 3. Discard the supernatant then resuspend the pellet in 600  $\mu l$  of Sorbitol Buffer.
- 4. Add 200 U of lyticase or zymolase then incubate at 30°C for 30 minutes.
- 5. Centrifuge the mixture for 10 minutes at 2,000 x g to harvest the spheroplast then remove the supernatant.

#### 2. Lysis

- 1. Add 300 µl of Cell Lysis Buffer then resuspend the cell pellet by pipette.
- 2. Incubate at 60°C for at least 10 minutes to ensure the sample lysate is clear.

NOTE: During incubation, invert the tube every 3 minutes

#### **Optional RNA Removal Step**

Following 60°C incubation, add 5 µl of RNase A (50 mg/ml) to the clear sample lysate then mix by vortex. Incubate at room temperature for 10 minutes.

#### 3. Protein Removal

1. Add 100 µl of Protein Removal Buffer to the sample lysate then vortex IMMEDIATELY for 10 seconds.

2. Centrifuge at 14-16,000 x g for 3 minutes to form a tight, white, protein pellet.

NOTE: Following centrifugation the protein should form a tight, white, pellet. If the pellet is not tight then incubate on ice for 5 minutes followed by centrifugation at 14-16,000 x g for another 3 minutes.

#### 4. DNA Precipitation

- 1. Being careful not to draw any of the protein pellet into the pipette, transfer the supernatant from Step 3 to a new 1.5 ml microcentrifuge tube.
- 2. Add 300 µl of isopropanol and mix well by gently inverting 20 times.
- 3. Centrifuge at 14-16,000 x g for 5 minutes.
- 4. Carefully remove the supernatant then add 300 µl of 70% ethanol to wash the pellet.
- 5. Centrifuge at 14-16,000 x g for 3 minutes.
- 6. Discard the supernatant then air-dry the pellet for 10 minutes.
- NOTE: DO NOT dry the DNA pellet with by vacuum centrifuge and avoid over drying the DNA pellet.

#### 5. DNA Rehydration

1. Add 50-100  $\mu$ I of DNA Hydration Buffer or ddH<sub>2</sub>O then incubate at 60°C for 10 minutes to dissolve the DNA pellet.

NOTE: Occasionaly tapping the bottom of the tube during incubation will promote DNA rehydration.

## gYEAST Genomic DNA Kit Functional Test Data



M 1 2 3

**Figure 1**. Genomic DNA (approx. 30 kb) was extracted using the gYEAST Genomic DNA Kit. *Saccharomyces cerevisiae*  $(2 \times 10^8)$  was harvested by centrifugation at 5,000 x g for 10 minutes. A 15 µl aliquot of extracted genomic DNA from a 100 µl eluate was analyzed by electrophoresis on a 1% agarose gel.

M = 1 Kb DNA Ladder

| Test | DNA Concentration | 260/280 | 260/230 | Yield   |
|------|-------------------|---------|---------|---------|
| 1    | 115.5 µg/ml       | 1.91    | 1.87    | 11.6 µg |
| 2    | 142.9 µg/ml       | 1.92    | 2.01    | 14.3 µg |
| 3    | 137.1 µg/ml       | 1.91    | 1.97    | 13.7 µg |

# Troubleshooting

| Problem                    | Cause   | Solution   |  |
|----------------------------|---|--|--|
|                            | A. Sample lysis or homogenization was         | A. Starting material should be reduced               |  |
| Low Yield                  | incomplete                                    | B. Following isopropanol addition, increase standing |  |
|                            | B. Incorrect DNA precipitation                | time to improve DNA precipitation. Following         |  |
|                            | C. Precipitate was formed during Step 4       | centrifugation, carefully remove the supernatant     |  |
|                            |   | without contacting the DNA pellet.                   |  |
|                            |   | C. Reduce starting material                          |  |
| Degraded DNA               | A. Incorrect sample preparation               | A. Process samples immediately after collection      |  |
| Degraded DNA               | B. Incorrect sample storage                   | B. Extracted DNA should be stored at -20°C           |  |
|                            | A. Did not perform optional RNase A treatment | A. If DNA is used for sensitive downstream           |  |
| <b>RNA</b> Contamination   |   | applications it might be necessary to extract        |  |
| RNA Contamination          |   | RNA-free DNA. Therefore, RNase A treatment           |  |
|                            |   | should be performed                                  |  |
| Eluted DNA does not        | A. Residual ethanol contamination             | A. Increase DNA pellet drying time to ensure         |  |
| perform well in downstream |   | residual ethanol is completely evaporated            |  |
| applications               |   |  |  |

# **Related DNA/RNA Purification and Extraction Products**

| 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 Extraction and 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      |

# **Related DNA/RNA Purification and Extraction Products**

| RNA Extraction and Purification          |                  |                  |
|--|------------------|------------------|
| Product                                  | Package Size     | Catalogue Number |
| Total RNA Mini Kit (Blood/Cultured Cell) | 50/100/300 preps | IB47321/322/323  |
| Total RNA Maxi Kit (Blood/Cultured Cell) | 10 preps         | IB47330          |
| Total RNA Mini Kit (Tissue)              | 50/100 preps     | IB47301/302      |
| Total RNA Maxi Kit (Tissue)              | 10 preps         | IB47310          |
| Total RNA Mini Kit (Plant)               | 50/100 preps     | IB47341/342      |
| Total RNA Maxi Kit (Plant)               | 10 preps         | IB47350          |
| rBAC Mini RNA Bacteria Kit               | 100/300 preps    | IB47421/412      |
| rYeast Total RNA Mini Kit                | 50/100/300 preps | IB47411/422      |
| 96-Well Total RNA Extraction Kit (Plant) | 4/10 x 96 preps  | IB47381/382      |
| 96-Well Total RNA Extraction Kit         | 4/10 x 96 preps  | IB47360/361      |
| miRNA Isolation Kit                      | 100 preps        | IB47371          |
| Virus DNA/RNA Purification               |                  |                  |
| Product                                  | Package Size     | Catalogue Number |
| Viral Nucleic Acid Extraction Kit II     | 50/100/300 preps | IB47401/402/403  |

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