

# 96 Well Genomic DNA Bacteria Kit

**IB47295** (2 x 96 well plates/kit)

**IB47296** (4 x 96 well plates/kit)

**IB47297** (10 x 96 well plates/kit)

## Advantages

**Sample:** up to  $5 \times 10^8$  Gram-positive and Gram-negative bacteria cells

**Yield:** up to 5  $\mu$ g of genomic DNA

**Format:** 96 Well gDNA Binding Plate

**Operation Time:** within 90 minutes

**Elution Volume:** 100~200  $\mu$ l

**Kit Storage:** dry at room temperature (15-25°C), Lysozyme is shipped at room temperature and should be stored at -20°C for extended periods

## Table of Contents

Introduction.....	2
Quality Control.....	2
Kit Components.....	2
Safety Measures.....	3
Quick Protocol Diagram.....	3
Centrifuge Protocol Procedure.....	4
Test Data.....	7
Troubleshooting.....	8
Related Products.....	9

## Introduction

The 96 Well Genomic DNA Bacteria Kit is optimized for high-throughput genomic and viral DNA purification from Gram (-) negative and Gram (+) positive bacterial cells. Gram+ Buffer, when combined with Lysozyme, will efficiently lyse bacterial cell walls consisting of the peptidoglycan layer. Proteinase K and chaotropic salt are used to further lyse cells and degrade protein, allowing DNA to easily bind to the glass fiber matrix of the binding plate. Contaminants are removed using a Wash Buffer (containing ethanol) and the purified genomic DNA is eluted by a low salt Elution Buffer, TE or water. Phenol/chloroform extraction or alcohol precipitation is not required and the purified genomic DNA is ready for use in a variety of downstream applications.

## Quality Control

The quality of the 96 Well Genomic DNA Bacteria Kit is tested on a lot-to-lot basis by isolating DNA from *Escherichia coli* ( $5 \times 10^8$ ) culture. 5  $\mu$ l from a 100  $\mu$ l eluate of purified DNA is analyzed by electrophoresis on a 1% agarose gel.

## Kit Components

Component	IB47295	IB47296	IB47297
Gram+ Buffer	50 ml	100 ml	220 ml
GT Buffer	60 ml	100 ml	200 ml
GB Buffer	60 ml	100 ml	155 ml x 1 60 ml x 1
W1 Buffer	80 ml	200 ml	200 ml x 2
Wash Buffer <sup>1</sup> (Add Ethanol)	50 ml (200 ml)	50 ml x 2 (200 ml x 2)	50 ml x 4 (200 ml x 4)
Proteinase K <sup>2</sup> (Add ddH <sub>2</sub> O)	11 mg x 4 (1.1 ml x 4)	65 mg x 1 (6.5 ml) 11 mg x 2 (1.1 ml x 2)	55 mg x 4 (5.5 ml x 4)
Lysozyme <sup>3</sup>	110 mg x 2	110 mg x 1 250 mg x 1	610 mg x 1 250 mg x 1
Elution Buffer	60 ml	100 ml	120 ml x 2
Presto™ gDNA 96 Well Binding Plates	2	4	10
Microtubes (Racked)	2	2	2
Microtubes (8-strip)	N/A	12 x 2	12 x 8
Caps for Microtubes (8-strip)	24	48	60 x 2
96 Deep Well Plates <sup>4</sup>	2	2	2
Adhesive Film	10	20	50

<sup>1</sup>Add absolute ethanol (see the bottle label for volume) to Wash Buffer then mix by shaking for a few seconds. Check the box on the bottle. Close the bottle tightly after each use to avoid ethanol evaporation.

<sup>2</sup>Add ddH<sub>2</sub>O to Proteinase K (see the bottle label for volume) then vortex to ensure it is completely dissolved. Check the box on the bottle. Once it is dissolved completely, centrifuge for a few seconds to spin the mixture down. For extended periods, the mixture should be stored at 4°C. Use only fresh ddH<sub>2</sub>O as ambient CO<sub>2</sub> can quickly cause acidification.

<sup>3</sup>Lysozyme should be stored at -20°C for extended periods.

<sup>4</sup>96 Deep Well Plates are reusable. After use, rinse with water then incubate in 0.4M HCl for 1 minute at room temperature. Wash the plate thoroughly with ddH<sub>2</sub>O then autoclave.



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

## Quick Protocol Diagram



Transfer 1.5 ml of cultured bacteria broth (up to  $5 \times 10^8$  bacterial cells) into each tube of the 96 Deep Well Plate.



Lyse bacteria cells with Proteinase K and Lysozyme



Incubate at 60°C for 10 min. (Proteinase K) and/or 37°C for 30 min. (Lysozyme).



DNA binding



Wash



Elution of pure DNA into Microtubes (Racked)

# 96 Well Genomic DNA Bacteria 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 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. Add ddH<sub>2</sub>O to Proteinase K (see the bottle label for volume) then vortex to ensure it is completely dissolved. Check the box on the bottle. Once it is dissolved completely, centrifuge for a few seconds to spin the mixture down. For extended periods, the mixture should be stored at 4°C. Use only fresh ddH<sub>2</sub>O as ambient CO<sub>2</sub> can quickly cause acidification.
3. 96 Deep Well Plates are reusable. After use, rinse with water then incubate in 0.4M HCl for 1 minute at room temperature. Wash the plate thoroughly with ddH<sub>2</sub>O then autoclave.

### Additional Requirements

Centrifuge with microplate buckets, 37°C and 60°C incubator, additional 96 Deep Well Plates, and absolute ethanol.

## Centrifuge Protocol Procedure

### 1. Sample Collection

Transfer up to **1.5 ml of cultured bacteria broth or 10-20 mg (wet weight) of bacteria pellet (up to 5 x 10<sup>8</sup> bacterial cells)** into each well of a **96 Deep Well Plate**. Dry the top of the plate with paper towel then seal the plate with **Adhesive Film**. Centrifuge the **96 Deep Well Plate** for 5 minutes at 3,000 x g to pellet the bacterial culture. Carefully remove the **Adhesive Film** from the **96 Deep Well Plate** and remove the supernatant in each well by quickly inverting the plate.

### 2A. Gram (-) Negative Bacteria Preparation

#### Proteinase K Working Solution

- A. Mix **20 µl of Proteinase K and 180 µl of GT Buffer** per sample by vortex.
- B. For 96 samples, mix **2 ml of Proteinase K and 18 ml of GT Buffer** by vortex.

Add **200 µl of Proteinase K working solution** into each well of the 96 Deep Well Plate. Re-suspend the cell pellet by pipette until all traces of the pellet have been dissolved. Dry the top of the plate with paper towel then seal with new Adhesive Film. Incubate at 60°C for 10 minutes. For optimal lysis, mix occasionally or place the plate on a rocking platform during incubation.

NOTE: Pre-warm the required volume of **Elution Buffer (200 µl/ sample)** to 60°C for step 6 DNA Elution.

Proceed to Optional RNA Removal Step or Lysis on Page 5.

## 2B. Gram (+) Positive Bacteria Preparation

### Lysozyme Working Solution

A. Mix **0.8 mg of Lysozyme and 200 µl of Gram+ Buffer** per sample by vortex.

B. For 96 samples, mix **80 mg of Lysozyme and 20 ml of Gram+ Buffer** by vortex.

Add **200 µl of Lysozyme Working Solution** into each well of the **96 Deep Well Plate** containing the bacterial pellet. Resuspend the cell pellet by pipette until all traces of the cell pellet have been dissolved. Dry the top of the plate with paper towel. Seal the plate with new **Adhesive Film** then incubate at 37°C for 30 minutes. For optimal lysis, mix occasionally or place the plate on a rocking platform during incubation. Briefly centrifuge the **96 Deep Well Plate** at 3,000 x g to collect any solution from the **Adhesive Film**. Allow the centrifuge to reach 3,000 x g prior to stopping. Carefully remove the **Adhesive Film** then add **20 µl of Proteinase K** into each well of the **96 Deep Well Plate**. Dry the top of the plate with paper towel then seal with new **Adhesive Film**. Mix the samples by vortex then incubate the **96 Deep Well Plate** at 60°C for 10 minutes. For optimal lysis, mix occasionally or place the plate on a rocking platform during incubation.

### Optional RNA Removal Step

Briefly centrifuge the 96 Deep Well Plate at 3,000 x g to collect any solution from the Adhesive Film. Allow the centrifuge to reach 3,000 x g prior to stopping. Carefully remove the Adhesive Film. Add 10 µl of RNase A (50 mg/ml) into each well of the 96 Deep Well Plate and mix well by pipette. Incubate the 96 Deep Well Plate for 10 minutes at room temperature.

NOTE: Pre-warm the required volume of **Elution Buffer (200 µl/ sample)** to 60°C for step 6 DNA Elution.

## 3. Lysis

Briefly centrifuge the **96 Deep Well Plate** at 3,000 x g to collect any solution from the **Adhesive Film**. Allow the centrifuge to reach 3,000 x g prior to stopping. Carefully remove the **Adhesive Film** then add **200 µl of GB Buffer** into each well. Dry the top of the plate with paper towel then seal with new **Adhesive Film**. Mix the sample by inverting the plate 10 times. Incubate the **96 Deep Well Plate** at 70°C for 10 minutes. For optimal lysis, mix occasionally or place the plate on a rocking platform during incubation.

#### 4. Binding

Briefly centrifuge the **96 Deep Well Plate** at 3,000 x g to collect any solution from the **Adhesive Film**. Allow the centrifuge to reach 3,000 x g prior to stopping. Carefully remove the **Adhesive Film** then add **200 µl of absolute ethanol** into each well. Dry the top of the plate with paper towel then seal with new **Adhesive Film**. Mix the sample by shaking the plate vigorously for 15-30 seconds. Briefly centrifuge the **96 Deep Well Plate** at 3,000 x g to collect any solution from the **Adhesive Film**. Allow the centrifuge to reach 3,000 x g prior to stopping. Place a **96 Well gDNA Binding Plate** on a new **96 Deep Well Plate**. Carefully remove the **Adhesive Film** from the **96 Deep Well Plate** and transfer all of the sample lysate into each well of the **96 Well gDNA Binding Plate**, being careful not to get any lysate on the rims of the wells. Centrifuge the **96 Well gDNA Binding Plate** and **96 Deep Well Plate** together at 3,000 x g for 5 minutes. Discard the flow-through then place the **96 Well gDNA Binding Plate** back on the **96 Deep Well Plate**.

#### 5. Wash

Add **400 µl of W1 Buffer** to each well of the **96 Well gDNA Binding Plate** then centrifuge the **96 Well gDNA Binding Plate** and **96 Deep Well Plate** together at 3,000 x g for 5 minutes. Discard the flow-through then place the **96 Well gDNA Binding Plate** back on the **96 Deep Well Plate**. Add **500 µl of Wash Buffer (make sure ethanol was added)** to each well of the **96 Well gDNA Binding Plate** then centrifuge the **96 Well gDNA Binding Plate** and **96 Deep Well Plate** together at 3,000 x g for 5 minutes. Add another **500 µl of Wash Buffer (make sure ethanol was added)** to each well of the **96 Well gDNA Binding Plate** then centrifuge the **96 Well gDNA Binding Plate** and **96 Deep Well Plate** together at 3,000 x g for 5 minutes. Discard the flow-through then place the **96 Well gDNA Binding Plate** back on the **96 Deep Well Plate**. Centrifuge the **96 Well gDNA Binding Plate** and **96 Deep Well Plate** together at 3,000 x g for 10 minutes to dry the membrane.

## 6. Elution

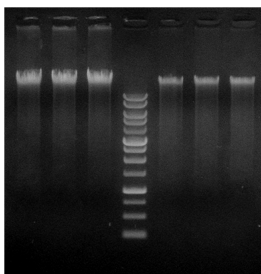
Remove the **96 Well gDNA Binding Plate** from the **96 Deep Well Plate** then blot the nozzles on a clean absorbent paper towel to remove residual ethanol. Place the **96 Well gDNA Binding Plate** on **Microtubes (Racked)**. Add **100 µl of pre-heated Elution Buffer<sup>1</sup>**, TE<sup>2</sup> or water<sup>3</sup> to the center of each well of the **96 Well gDNA Binding Plate**. Let stand for at least 2 minutes to ensure the Elution Buffer, TE or water is absorbed by the membrane. Centrifuge the **96 Well gDNA Binding Plate** and **Microtubes (Racked)** together at 3,000 x g for 5 minutes to elute the purified DNA. Seal the **Microtubes (Racked)** with new caps and store the purified DNA at -20°C. NOTE: For maximum DNA yield, repeat the elution step by adding 100 µl of pre-heated Elution Buffer, TE or water to each well of the 96 Well gDNA Binding Plate then centrifuge again. If a higher DNA concentration is required, use 50 µl of pre-heated Elution Buffer then repeat the Elution step by adding the same 50 µl of Elution Buffer (which now contains the eluted DNA) to each well of the 96 Well gDNA Binding Plate then centrifuge again.

<sup>1</sup>Ensure that Elution Buffer (10 mM Tris-HCl, pH8.5 at 25°C) is added into the CENTER of the well matrix and is completely absorbed.

<sup>2</sup>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 well matrix and is completely absorbed.

<sup>3</sup>If using water for elution, ensure the water pH is ≥8.0. ddH<sub>2</sub>O should be fresh as ambient CO<sub>2</sub> can quickly cause acidification. Ensure that water is added into the CENTER of the well matrix and is completely absorbed. DNA eluted in water should be stored at -20°C to avoid degradation.

## 96 Well gDNA Kit Functional Test Data



**Figure 1.** Genomic DNA was extracted from bacteria cells using the 96 Well Genomic DNA Bacteria Kit. The purified genomic DNA was eluted in 100 µl of Elution Buffer and 10 µl aliquots of the final sample (chosen from 6 random wells) were analyzed by electrophoresis on a 1% agarose gel.

M = 1 Kb DNA Ladder

1-3: 5 x 10<sup>8</sup> *Escherichia coli* cells, 4-6: 5 x 10<sup>8</sup> *Bacillus subtilis* cells

Sample	ng/µl	260/280	Yield (µg)
1	49.2	1.80	3.9
2	56.2	1.79	4.5
3	55.8	1.80	4.5
4	22.4	1.87	1.8
5	19.9	1.84	1.6
6	20.4	1.80	1.6

1 2 3 M 4 5 6

# Troubleshooting



## Low Yield

### **Incorrect Sample Lysis**

When extracting genomic DNA from Gram (+) positive bacteria, prepare and use Lysozyme working solution to lyse bacteria cells. When extracting genomic DNA from Gram (-) negative bacteria, prepare and use Proteinase K working solution to lyse bacteria cells.

### **Too much starting material**

Extract genomic DNA from up to  $5 \times 10^8$  bacteria cells per reaction.

### **Incorrect DNA elution step**

Use pre-heated Elution Buffer, TE or water (60 °C) to elute DNA. Ensure Elution Buffer, TE or water is added into the center of the matrix and is completely absorbed. If using water for elution, ensure the water pH is between 7.5 and 8.5. Elute twice to increase the DNA recovery.

## DNA contaminated with RNA

### **RNA carry-over**

Perform RNA removal step.

## Eluted DNA does not perform well in downstream applications

### **Residual ethanol contamination**

Following the wash step, dry the Binding plate with additional centrifugation at 3,000 x g for 10 minutes to ensure the membrane is completely dry.



## 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](http://www.ibisci.com). Thank you!





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