

Instruction Manual Ver. 02.10.17 For Research Use Only

Presto[™] 96 Well PCR Cleanup Kit

96DFH02 (2 x 96 well plates/kit) **96DFH04** (4 x 96 well plates/kit) **96DFH10** (10 x 96 well plates/kit)

Advantages

Sample: up to 100 µl of PCR products per well Fragment Size: 70 bp-20 kb Recovery: up to 95% Format: Presto[™] PCR Cleanup 96 Well Binding Plates for efficient vacuum filtration and centrifugation Operation Time: 20 minutes Elution Volume: 60 µl from 80 µl elution buffer volume, and 40 µl from 60 µl elution buffer volume Kit Storage: dry at room temperature (15-25°C)

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Introduction

Presto[™] 96 Well PCR Cleanup Kits were designed to recover or concentrate DNA fragments from PCR or other enzymatic reactions using an efficient 96 well binding plate system. Chaotropic salt is used to denature enzymes. DNA fragments in the chaotropic salt are bound by the glass fiber matrix in each well of the binding plate. Contaminants are removed with a Wash Buffer (containing ethanol) and the purified DNA fragments are eluted by a low salt Elution Buffer, TE or water. Salts, enzymes and unincorporated nucleotides can be effectively removed from the reaction mixture without phenol extraction or alcohol precipitation and the purified DNA is ready for use in subsequent reactions.

Quality Control

The quality of the Presto[™] 96 Well PCR Cleanup Kit is tested on a lot-to-lot basis by purifying DNA fragments of various sizes from PCR products. The purified DNA is analyzed by electrophoresis.

Kit Components

Component	96DFH02	96DFH04	96DFH10
Binding Buffer	40 ml	80 ml	240 ml x 1
Wash Buffer ¹ (Add Ethanol)	50 ml (200 ml)	50 ml x 2 (200 ml x 2)	50 ml x 5 (200 ml x 5)
Elution Buffer	30 ml	60 ml	100 ml
Presto™ PCR Cleanup 96 Well Binding Plates	2	4	10
0.35 ml Collection Plates	2	4	10
Adhesive Film	4	8	20

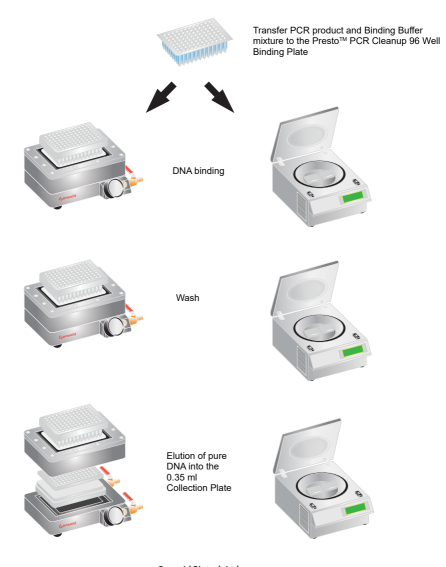
¹Add absolute ethanol (see the bottle label for volume) to Wash Buffer then mix by shaking for a few seconds. Check the box on the bottle. Be sure and close the bottle tightly after each use to avoid ethanol evaporation.

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Binding Buffer contains guanidine thiocyanate. During the procedure, always wear a lab coat, disposable gloves, and protective goggles.

Quick Vacuum and Centrifuge Protocol Diagram







Presto[™] 96 Well PCR Cleanup 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. It is not necessary to remove mineral oil or kerosene from the PCR sample prior to cleanup.

Additional Requirements absolute ethanol, 96 Deep Well Plate (optional), Square-well Block (optional)

PCR Cleanup Vacuum Protocol Procedure

1. Vacuum Manifold Preparation

Place the waste tray on the manifold base then place the binding top plate on the manifold base. Place the **Presto[™] PCR Cleanup 96 Well Binding Plate** in the binding top plate aperture. Seal unused wells of the **Presto[™] PCR Cleanup 96 Well Binding Plate** with **Adhesive Film** then attach the vacuum manifold to a vacuum source.

2. DNA Binding

Add **3 volumes of Binding Buffer to 1 volume of PCR sample** then mix by pipetting. Transfer the sample mixture to each well of the **Presto[™] PCR Cleanup 96 Well Binding Plate** (E.g. Add 150 µl of Binding Buffer to 50 µl of PCR sample). Apply vacuum at 15 inches Hg until samples pass through completely (approximately 10 seconds) then turn off the vacuum.

NOTE: If the PCR sample is less than 50 µl, adjust the volume to 50 µl with ddH₂O.

3. Wash

Add **500 µl of Wash Buffer (make sure ethanol was added)** to each well of the **Presto[™] PCR Cleanup 96 Well Binding Plate**. Let stand for 1 minute. Apply vacuum at 15 inches Hg until Wash Buffer passes through completely (approximately 10 seconds) then turn off the vacuum. **Add 500 µl of Wash Buffer (make sure ethanol was added)** to each well. Apply vacuum at 15 inches Hg until Wash Buffer passes through completely. Continue to apply vacuum for an additional 10 minutes to dry the membrane then turn off the vacuum.

4. Elution

Remove the **Presto[™] PCR Cleanup 96 Well Binding Plate** from the binding top plate aperture and blot the nozzles on a clean absorbent paper towel to remove residual ethanol. Remove the waste tray from the manifold base then place the collection plate spacer on the manifold base. Place a **0.35 ml collection plate** on top of the collection plate spacer. Place the binding top plate back on the manifold base then place the **Presto[™] PCR Cleanup 96 Well Binding Plate** back in the binding top plate aperture. Add **60-80 µl of Elution Buffer**¹, TE² or water³ into the **CENTER** of each well of the **Presto[™] PCR Cleanup 96 Well Binding Plate**. Let stand for at least 3 minutes to ensure the Elution Buffer, TE or water is absorbed by the membrane. Apply vacuum at 15 inches Hg for 5 minutes then turn off the vacuum. Seal the **0.35 ml Collection Plate** with **Adhesive Film** and store the purified DNA at -20°C.

NOTE: The average eluate volume is 60 μl from 80 μl elution buffer volume, and 40 μl from 60 μl elution buffer volume.

 1 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.

²Using 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.

³If using water for elution, ensure the water pH is ≥ 8.0 . ddH₂O should be fresh as ambient CO₂ 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.

PCR Cleanup Centrifuge Protocol Procedure

1. DNA Binding

Place the **Presto[™] PCR Cleanup 96 Well Binding Plate** on a 96 Deep Well Plate or a standard Square-Well Block. Add **3 volumes of Binding Buffer to 1 volume of the PCR sample** then mix by pipetting. Transfer the sample mixture to each well of the **Presto[™] PCR Cleanup 96 Well Binding Plate**. (E.g. add 150 µl of Binding Buffer to 50 µl PCR sample).

NOTE: If the PCR sample is less than 50 μ l, adjust the volume to 50 μ l with ddH₂O.

Centrifuge the **Presto[™] PCR Cleanup 96 Well Binding Plate** and 96 Deep Well Plate together at 3,000 x g for 5 minutes. Discard the flow-through then place the **Presto[™] PCR Cleanup 96 Well Binding Plate** back on the 96 Deep Well Plate.



2. Wash

Add 500 µl of Wash Buffer (make sure ethanol was added) to each well of the Presto[™] PCR Cleanup 96 Well Binding Plate. Let stand for 1 minute. Centrifuge the Presto[™] PCR Cleanup 96 Well Binding Plate and 96 Deep Well Plate together at 3,000 x g for 5 minutes. Discard the flow-through then place the Presto[™] PCR Cleanup 96 Well 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 Presto[™] PCR Cleanup 96 Well Binding Plate and 96 Deep Well Plate. Centrifuge the Presto[™] PCR Cleanup 96 Well Binding Plate and 96 Deep Well Plate together at 3,000 x g for 5 minutes. Discard the flow-through then place the Presto[™] PCR Cleanup 96 Well Binding Plate and 96 Deep Well Plate together at 3,000 x g for 5 minutes. Discard the flow-through then place the Presto[™] PCR Cleanup 96 Well Binding Plate back on the 96 Deep Well Plate. Centrifuge the Presto[™] PCR Cleanup 96 Well Binding Plate and 96 Deep Well Plate together at 3,000 x g for 5 minutes.

3. Elution

Remove the **Presto[™] PCR Cleanup 96 Well Binding Plate** from the 96 Deep Well Plate then blot the nozzles on a clean absorbent paper towel to remove residual ethanol. Place the **Presto[™] PCR Cleanup 96 Well Binding Plate** on a **0.35 ml Collection Plate**. Add **60-80 µl of Elution Buffer**¹, TE² or water³ to the center of each well of the **Presto[™] PCR Cleanup 96 Well Binding Plate**. Let stand for at least 3 minutes to ensure the Elution Buffer, TE or water is absorbed by the membrane. Centrifuge the **Presto[™] PCR Cleanup 96 Well Binding Plate** and **0.35 ml Collection Plate** together at 3,000 x g for 5 minutes to elute the purified DNA. Seal the **0.35 ml Collection Plate** with **Adhesive Film** and store the purified DNA at -20°C. NOTE: The average eluate volume is 60 µl from 80 µl elution buffer volume, and 40 µl from 60 µl elution buffer volume.

¹Ensure 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.

²Using 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.

³If using water for elution, ensure the water pH is \geq 8.0. ddH₂O should be fresh as ambient CO₂ 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.

Troubleshooting



Incomplete Wash Buffer preparation.

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.

Incorrect DNA Elution step.

Low Yield

Ensure that Elution Buffer, TE or water is added into the **CENTER** of the well matrix and is completely absorbed. If DNA fragments are larger than 5 kb, use pre-heated Elution Buffer, TE, or water ($60 \sim 70^{\circ}$ C). If using water for elution, ensure the water pH is $\geq 8.0.$ ddH₂O should be fresh as ambient CO₂ can quickly cause acidification.

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Presto[™] 96 Well PCR Cleanup Kit Functional Test Data

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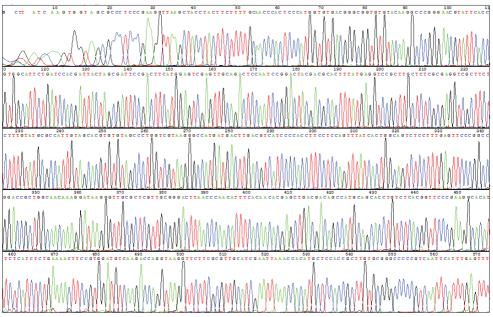


Figure 1. Sequencing data of PCR product (*E. coli* 16S ribosomal DNA fragment) purified using the Presto™ 96 Well PCR Cleanup Kit.

Related Post Reaction DNA Extraction Products

Product	Package Size	Catalogue Number
GenepHlow™ Gel Extraction Kit	100/300 preps	DFG100/300
GenepHlow™ PCR Cleanup Kit	100/300 preps	DFC100/300
GenepHlow™ Gel/PCR Kit	100/300 preps	DFH100/300
GenepHlow™ DNA Cleanup Midi Kit	100/300 preps	DFI100/300
GenepHlow™ DNA Cleanup Maxi Kit	10/25 preps	DFM010/025
Small DNA Fragments Extraction Kit	100/300 preps	DF101/301
Presto [™] Max Gel/PCR Kit (Large DNA Fragments)	100/300 preps	DFL100/300
Presto [™] 96 Well PCR Cleanup Kit	4/10 x 96 preps	96DFH04/10
Presto [™] 96 Well Gel Extraction Kit	4/10 x 96 preps	96DFG04/10
G-25 Gel Filtration Desalting Column	50 rxns	CG025
G-50 Gel Filtration Dye Terminator Removal Column	50 rxns	CG050
96-Well G-50 Gel Filtration Plate	4/10 x 96 rxns	CGP04/10







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