

E-Z 96 Plant DNA Kit

D1086-01	1 x 96 preps
D1086-02	4 x 96 preps
D1086-03	20 x 96 preps

Manual Date: May 2019
Revision Number: v7.0

For Research Use Only

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E-Z 96 Plant DNA Kit

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Introduction

E-Z 96 Plant DNA Kit allows for the rapid and reliable isolation of high-quality total cellular DNA from a wide variety of plant species and tissues in a 96-well plate format. E-Z 96 Plant DNA Kit adapted the buffer system from Omega Bio-tek's SP Plant DNA system which is used to process a variety of plant types, particularly those with unusually high levels of phenolic compounds or polysaccharides such as cotton, pine, and peanut samples. Up to 50 mg wet tissue (or 12 mg dry tissue) can be processed in each well in less than one hour. The system combines the reversible nucleic acid-binding properties of the HiBind® matrix with the speed and versatility of the E-Z 96 DNA Plate to eliminate polysaccharides, phenolic compounds, and enzyme inhibitors from plant tissue lysates. Purified DNA is suitable for PCR, restriction digestion, and hybridization applications.

If using the E-Z 96 Plant DNA Kit for the first time, please read this booklet to become familiar with the procedures. Dry or fresh plant tissue is disrupted and lysed in a specially formulated buffer containing a proprietary detergent mixture. Proteins, polysaccharides, and cellular debris are precipitated. Binding conditions are adjusted and the sample is transferred to the E-Z 96 DNA Plate. Two rapid wash steps remove trace contaminants such as residual polysaccharides and pure DNA is eluted in water or low ionic strength buffer. Purified DNA can be used directly in downstream applications without the need for further purification. Optional E-Z 96 Lysate Clearance Plates (Product No. FL96-01) are available for use with this kit.

New In this Edition

May 2019

- SPW Wash Buffer has been renamed SPW Buffer. This is a name change only. The formulation has not changed.

April 2018

- 1.2 mL HTS Plate has been replaced with Lysis Plate.
- The number of Caps for Racked Microtubes has been reduced and Silicone Mats are now included for use with the Lysis Plate.

January 2016

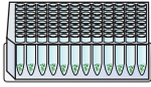
- 96-well Round-well Plates (1.2 mL) have been replaced with 1.2 mL HTS Plates.
- Caps for Round-well Plates have been replaced with Caps for Racked Microtubes.

April 2013

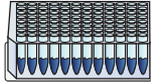
- Equilibration Buffer is no longer included with this kit. An optional Column Equilibration Protocol has been added to the protocol for your convenience.
- Equilibration Buffer is replaced with 3M NaOH provided by the user.

Centrifugation Protocol

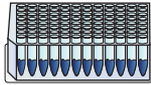
Vacuum Protocol



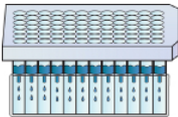
Collect Plant Tissue and Homogenize



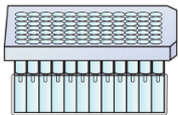
Lyse and Precipitate Polysaccharides



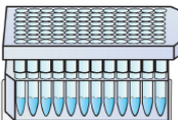
Transfer Cleared Lysate and Adjust Binding Conditions



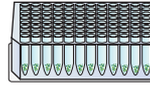
Bind and Wash 2X



Dry Membrane



Elute



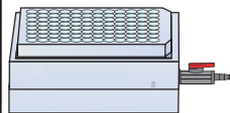
Collect Plant Tissue and Homogenize



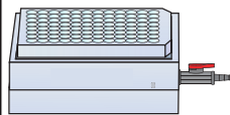
Lyse and Precipitate Polysaccharides



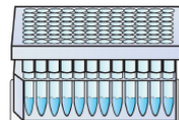
Transfer Cleared Lysate and Adjust Binding Conditions



Bind and Wash 2X



Dry Membrane



Elute

Kit Contents

Product Number	D1086-01	D1086-02	D1086-03
E-Z 96 DNA Plate	1	4	20
Lysis Plate	1	4	20
Silicone Mat	2	8	4 x 10
96-well Racked Microtubes (1.2 mL)	2	8	40
Caps for Racked Microtubes	25 x 8	100 x 8	500 x 8
AeraSeal Film	4	16	80
96-well Square-well Plate (2.2 mL)*	1	2	10
SP1 Buffer	60 mL	240 mL	2 x 600 mL
SP2 Buffer	20 mL	80 mL	440 mL
SP3 Buffer	30 mL	100 mL	2 x 250 mL
SPW Buffer	50 mL	200 mL	4 x 200 mL
Elution Buffer	60 mL	250 mL	1000 mL
RNase A	450 µL	1.35 mL	6 mL
User Manual	✓	✓	✓

* 96-well Square-well Plates (2.2 mL) are reusable; see Page 5 for cleaning instructions.

Storage and Stability

All components of the E-Z 96 Plant DNA Kit are guaranteed for at least 12 months from date of purchase when stored as follows. Store RNase A at 2-8° C. All other components should be stored at room temperature. During shipment or storage in cool ambient conditions, precipitates may form in SP1 Buffer and SP3 Buffer. Dissolve such deposits by warming the solution at 37°C and gently shaking.

Preparing Reagents

- Dilute SP3 Buffer with 100% ethanol as follows and store at room temperature.

Kit	100% Ethanol to be Added
D1086-01	60 mL
D1086-02	200 mL
D1086-03	500 mL per bottle

- Dilute SPW Buffer with 100% ethanol as follows and store at room temperature.

Kit	100% Ethanol to be Added
D1086-01	200 mL
D1086-02	800 mL
D1086-03	800 mL per bottle

- Prepare SP1 Buffer: Add 2 μ L RNase A per 400 μ L SP1 Buffer.

Cleaning Instructions for 96-well Square-well Plates

96-well Square-well Plates can be used to collect filtrate from the E-Z 96 DNA Plate. They are designed for repeated use. Wash the plates thoroughly in tap water after each use. Let sit for 5 minutes at room temperature in 0.5M HCl. Rinse with distilled water. Used plates can also be autoclaved after washing.

Guidelines for Vacuum Manifold

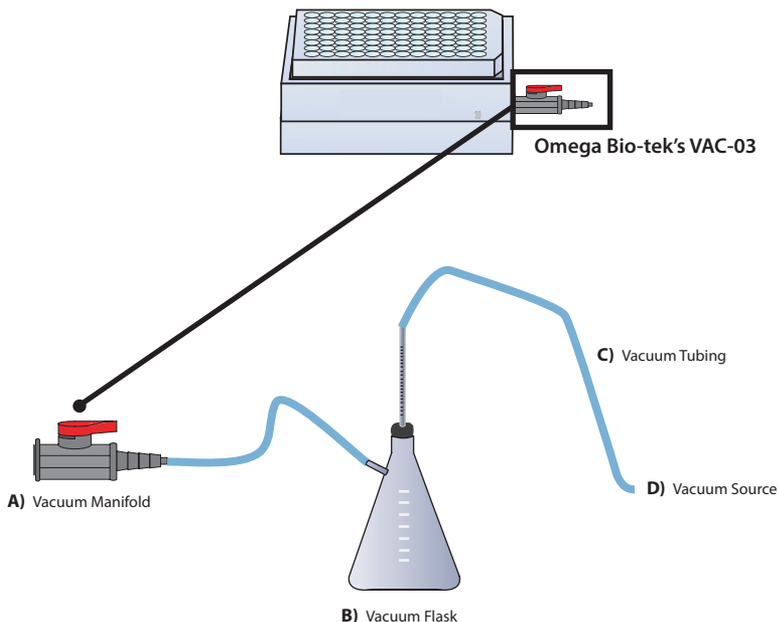
The following is required for use with the Vacuum Protocol:

- A) Vacuum Manifold (We recommend Omega Bio-tek's VAC-03)
Other Compatible Vacuum Manifolds: Qiagen QIAvac24, Sigma Aldrich VM20, Promega Vacman®, or manifold with standard Luer connector
- B) Vacuum Flask
- C) Vacuum Tubing
- D) Vacuum Source (review tables below for pressure settings)

Manifold	Recommended Pressure (mbar)
VAC-03	-200 to -400

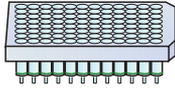
Conversion from millibars:	Multiply by:
Millimeters of Mercury (mmHg)	0.75
Kilopascals (kPa)	0.1
Inches of Mercury (inchHg)	0.0295
Torrs (Torr)	0.75
Atmospheres (atmos)	0.000987
Pounds per Square Inch (psi)	0.0145

Illustrated Vacuum Setup



Guidelines for Vacuum Manifold

DNA Bind and Wash Setup



E-Z 96 DNA Plate



Vacuum Manifold Top

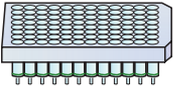


Waste Collection



Vacuum Manifold Base

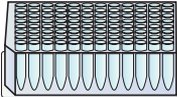
Standard Elution Setup



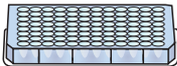
E-Z 96 DNA Plate



Vacuum Manifold Top



Racked Microtubes

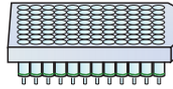


Microplate (300 µL)



Vacuum Manifold Base

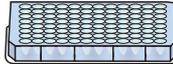
Optional Elution Setup



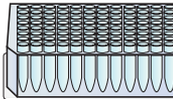
E-Z 96 DNA Plate



Vacuum Manifold Top



Microplate (300 µL)



Racked Microtubes



Vacuum Manifold Base

Disruption of Plant Tissues

1. Grind samples with pestle

A) Dry Specimens

Drying allows storage of field specimens for prolonged periods of time prior to processing. Samples can be dried overnight in a 45°C oven, powdered, and stored dry at room temperature. To prepare dried samples, place ~15 mg of dried tissues into a microcentrifuge tube (1.5 mL tubes are recommended) and grind using a pellet pestle. Disposable Kontes pestles work well and are available from Omega Bio-tek (Cat# SSI-1014-39 & SSI-1015-39). For critical work such as PCR and cloning, pestles are best used a single time then soaked in a dilute bleach solution immediately after use until clean. Disposable pestles may be autoclaved several times. A fine powder will ensure optimal DNA extraction and yield.

B) Fresh/Frozen Specimens

Due to the tremendous variation in water and polysaccharide content of plants, sample size should be limited to ~50 mg for first time users. It is very important to not overload the E-Z 96 DNA Plate. Too much starting material will decrease the yield and purity due to inefficient lysis. However, for some plant species, increasing the starting material can increase DNA yield. We recommend starting with 50 mg tissue. If results obtained are satisfactory, then increase amount of starting material. Best results are obtained with young leaves or needles.

Although various means of sample disruption can be used for this kit, such as beads or pestles, we recommend grinding the sample in liquid nitrogen. To prepare samples, collect tissue in a 1.5 mL or 2 mL microcentrifuge tube and dip the tube in liquid nitrogen with a pair of tweezers to fill the tube. Grind the tissue using disposable Kontes pellet pestles, which are available from OBI (Cat# SSI-1015-39). Alternatively, allow the liquid nitrogen to evaporate and store the samples at -70°C for later use. For critical work such as PCR and cloning, pestles are best used a single time then soaked in a dilute bleach solution immediately after use until clean. Disposable pestles may be autoclaved several times. For standard Southern analysis, the same pestle can be reused several times to grind multiple tissue samples by rinsing with ethanol and carefully wiping the surfaces clean between samples. Transfer the ground sample into a 96-well racked microtubes or 96-well deep-well plate.

Note: Do not allow the sample to thaw during handling and weighing. To prevent the sample from thawing, keep the rack or plate on a bed of dry ice.

Disruption of Plant Tissues

2. Disrupt Samples With Commercial Homogenizers

Fresh, frozen, and dried plant tissue can be effectively disrupted and homogenized by rapid agitation in the presence of beads.

For Fresh, Frozen and Lyophilized/Dried Tissue

1. Add one 3-4 mm stainless steel bead to each well of a 96-well round-well plate.
2. Close the individual tubes with Cap Strips.
3. Freeze the sample in liquid nitrogen.

Note: Lyophilized/Dried samples do not require freezing with liquid nitrogen.

4. Place the racks or plates into the clamps of the homogenizer.
5. Homogenize for 60-90 seconds at 30 Hz. Tissue samples are disrupted and simultaneously homogenized with the shearing and crushing action of the beads. Refer to manufacturer's protocol regarding use of liquid nitrogen with the homogenizer.

E-Z 96 Plant DNA Kit Protocols

E-Z 96 Plant DNA Kit Protocol - Centrifugation Protocol

Materials and Equipment to be Supplied by User:

- Centrifuge equipped with swing-bucket rotor capable of at least 3,000 x *g*
- Water baths, ovens, or incubators capable of 80°C
- Vortexer
- 100% ethanol
- Liquid nitrogen for freezing/disrupting samples (for fresh/frozen specimens)
- Equipment for disrupting plant tissue
- Ice, freezer, or 96-well cryorack at -20°C
- Optional: 3M NaOH

Before Starting:

- Prepare SP1 Buffer, SP3 Buffer, and SPW Buffer according to Preparing Reagents section on Page 5
- Set a water bath, oven, or incubator to 65°C
- Heat SP1 Buffer and Elution Buffer to 65°C (SP1 Buffer should be equilibrated to 80°C if liquid nitrogen is used for sample disruption)

1. Homogenize plant tissue following one of the methods described in the Disruption of Plant Tissue section on Pages 8-9.
2. Transfer up to 15 mg dry powdered tissue or 50 mg fresh (or frozen) tissue to a Lysis Plate (provided).

Note: No more than 50 mg (wet weight) or 15 mg (dry weight) starting material is recommended. More or less can be used depending on results. Water content (and buffer absorption) of samples affect optimal starting amounts.

3. Add 400 μ L SP1 Buffer to each sample. Seal the wells with Silicone Mat (provided). Vortex to mix thoroughly.

Note: SP1 Buffer must be mixed with RNase A before use. Please see the Preparing Reagents section on Page 5 for instructions. Ensure that all the samples are completely suspended and that there are no clumps in the solution. Clumps will result in low yields.

E-Z 96 Plant DNA Kit Protocols

4. Incubate at 65°C for 10 minutes. Mix samples twice during incubation by briefly shaking the plate side to side.
5. Centrifuge briefly to collect any drops of liquid from the mat. Do not prolong this step.
6. Remove and discard the mat.
7. Add 140 µL SP2 Buffer to each lysate.
8. Seal the wells with a new Silicone Mat. Vortex at maximum speed or shake the plate side to side vigorously for 20 seconds.
9. Centrifuge briefly to collect any drops of liquid from the mat. Do not prolong this step.

Note: Do not prolong this step. The brief centrifugation prevents the precipitates from freezing to the mat making it difficult to open the mat after incubation at -20°C in next step.

10. Incubate at -20°C for 10 minutes.
11. Centrifuge at 3,000-6,000 x *g* for 10 minutes.

Note: Optional E-Z 96 Lysate Clearance Plates (Cat# FL9601) are available for use with this kit at this step. Briefly, place an E-Z 96 Lysate Clearance Plate on top of the 96-well Racked Microtubes (provided). Transfer the desired volume of supernatant following the -20°C incubation in Step 10 to the E-Z 96 Lysate Clearance Plate. Centrifuge at 3,000-5,000 x *g* for 5 minutes. Proceed to Step 13.

12. Remove and discard the mat.
13. Transfer 400 µL supernatant to the 96-well Racked Microtubes. If less than 400 µL supernatant is recovered, adjust the volume of SP3 Buffer in Step 14.

E-Z 96 Plant DNA Kit Protocols

14. Add 1.5 volumes SP3 Buffer. A precipitate may form at this point; it will not interfere with DNA isolation.

Note: For example, if 400 μL lysate was transferred, add 600 μL SP3 Buffer.

15. Seal the microtubes with Caps for Racked Microtubes (provided). Vortex at maximum speed or shake the plate side to side vigorously for 20 seconds.
16. Centrifuge briefly to collect any drops of liquid from the caps. Do not prolong this step.
17. Place the E-Z 96 DNA Plate on to a 96-well Square-well Plate (provided).

Optional Protocol for Plate Equilibration:

1. Add 150 μL 3M NaOH to each well.
 2. Let sit at room temperature for 4 minutes.
 3. Centrifuge at 3,000-5,000 $\times g$ for 2 minutes.
 4. Discard the filtrate and reuse the 96-well Square-well Plate.
18. Carefully transfer 1 mL sample to the E-Z 96 DNA Plate. Be careful not to spill sample liquid onto the rims of the wells during the transfer.
 19. Seal the E-Z 96 DNA Plate with AeraSeal Film (provided).
 20. Centrifuge at 3,000-5,000 $\times g$ for 5 minutes or until all the sample has passed through the HiBind® membrane.
 21. Discard the filtrate and reuse the 96-well Square-well Plate.
 22. Remove the AeraSeal Film.
 23. Add 800 μL SPW Buffer to each well of the E-Z 96 DNA Plate. Seal the plate with new AeraSeal Film.

Note: SPW Buffer must be diluted with 100% ethanol prior to use. Please see Page 5 for instructions.

E-Z 96 Plant DNA Kit Protocols

24. Centrifuge at 3,000-5,000 x *g* for 5 minutes.
25. Discard the filtrate and reuse the 96-well Square-well Plate.
26. Repeat Steps 23-25 for a second SPW Buffer wash step.
27. Centrifuge at 3,000-5,000 x *g* for 15 minutes to dry the plate.

Note: It is important to dry the plate membrane before elution. Residual ethanol may interfere with downstream applications.

28. Remove the AeraSeal Film.
29. Transfer the E-Z 96 DNA Plate to new 96-well Racked Microtubes (provided) or a 96-well microplate (not provided).
30. Add 100 μ L Elution Buffer heated at 65°C to each well. Seal the plate with new AeraSeal Film.
31. Incubate at 65°C for 5 minutes.
32. Centrifuge at 5,000 x *g* for 5 minutes.
33. Repeat Steps 30-32 for a second elution step.

Note: To maintain higher DNA concentration, second elution may be performed with first eluate.
34. Seal the 96-well Racked Microtubes with Caps for Racked Microtubes.
35. Store DNA at -20°C.

E-Z 96 Plant DNA Kit Protocols

E-Z 96 Plant DNA Kit - Vacuum Protocol

The following protocol is based on using Omega Bio-tek's vacuum manifold (Cat# VAC-03).

Materials and Equipment to be Supplied by User:

- Vacuum manifold and vacuum source
- Centrifuge equipped with swing-bucket rotor capable of at least 3,000 x *g*
- Water bath, oven, or incubator capable of 80°C
- Vortexer
- 100% ethanol
- Liquid nitrogen for freezing/disrupting samples (For Fresh/Frozen Specimens)
- Equipment for disrupting plant tissue
- Ice, freezer, or 96-well cryorack at -20°C
- Sealing film
- Optional: 3M NaOH

Before Starting:

- Prepare SP1 Buffer, SP3 Buffer, and SPW Buffer according to Preparing Reagents section on Page 5
- Set a water bath, oven, or incubator to 65°C
- Heat SP1 Buffer and Elution Buffer to 65°C (SP1 Buffer should be equilibrated to 80°C if liquid nitrogen is used for sample disruption)

1. Homogenize plant tissue following one of the methods described in the Disruption of Plant Tissue section on Pages 8-9.
2. Transfer up to 15 mg dry powdered tissue or 50 mg fresh (or frozen) tissue to a Lysis Plate (provided).

Note: No more than 50 mg (wet weight) or 15 mg (dry weight) starting material is recommended. More or less can be used depending on results. Water content (and buffer absorption) of samples affect optimal starting amounts.

E-Z 96 Plant DNA Kit Protocols

3. Add 400 μ L SP1 Buffer to each sample. Seal the wells with a Silicone Mat (provided). Vortex to mix thoroughly.

Note: SP1 Buffer must be mixed with RNase A before use. Please see the Preparing Reagents section on Page 5 for instructions. Ensure that all the samples are completely suspended and that there are no clumps in the solution. Clumps will result in low yields.

4. Incubate at 65°C for 10 minutes. Mix samples twice during incubation by briefly shaking the plate side to side.
5. Centrifuge briefly to collect any drops of liquid from the mat. Do not prolong this step.
6. Remove and discard the mat.
7. Add 140 μ L SP2 Buffer to each lysate.
8. Seal the wells with a new Silicone Mat. Vortex at maximum speed or shake the plate side to side vigorously for 20 seconds.
9. Centrifuge briefly to collect any drops of liquid from the mat. Do not prolong this step.

Note: Do not prolong this step. The brief centrifugation prevents the precipitates from freezing to the mat making it difficult to open the caps after incubation at -20°C in next step.

10. Incubate at -20°C for 10 minutes.
11. Centrifuge at 3,000-6,000 $\times g$ for 10 minutes.

Note: Optional E-Z 96 Lysate Clearance Plates (Cat# FL9601) are available for use with this kit at this step. Briefly, place an E-Z 96 Lysate Clearance Plate on top of the 96-well Racked Microtubes (provided). Transfer the desired volume of supernatant following the -20°C incubation in Step 10 to the E-Z 96 Lysate Clearance Plate. Centrifuge at 3,000-5,000 $\times g$ for 5 minutes. Proceed to Step 13.

E-Z 96 Plant DNA Kit Protocols

12. Remove and discard the mat.
13. Transfer 400 μL supernatant to the 96-well Racked Microtubes. If less than 400 μL supernatant is recovered, adjust the volume of SP3 Buffer in Step 14.
14. Add 1.5 volumes SP3 Buffer. A precipitate may form at this point; it will not interfere with DNA isolation.

Note: For example, if 400 μL lysate was transferred, add 600 μL SP3 Buffer.
15. Prepare the vacuum manifold according to manufacturer's instructions.
16. Place an E-Z 96 DNA Plate on the top part of the vacuum manifold. Place the waste collection tray inside the base of the manifold. Seal any unused wells with sealing film (not provided).

Optional Protocol for Plate Equilibration:

1. Add 150 μL 3M NaOH to each well.
 2. Let sit at room temperature for 4 minutes.
 3. Turn on the vacuum source to draw the NaOH through the column.
 4. Turn off the vacuum.
-
17. Transfer 1 mL sample (including any precipitate that may have formed) from Step 14 to the E-Z 96 DNA Plate.
 18. Turn on the vacuum source to draw the sample through the plate.
 19. Turn off the vacuum.
 20. Repeat Steps 17-19 until all the sample has been transferred to the E-Z 96 DNA Plate.

E-Z 96 Plant DNA Kit Protocols

21. Add 800 μ L SPW Buffer to each well.

Note: SPW Buffer must be diluted with 100% ethanol prior to use. Please see Page 5 for instructions.

22. Turn on the vacuum source to draw the SPW Buffer through the plate.

23. Turn off the vacuum.

24. Repeat Steps 21-23 for a second SPW Buffer wash step.

25. Add 400 μ L 100% ethanol to each well.

26. Turn on the vacuum source to draw the ethanol through the plate.

27. Turn off the vacuum.

28. Continue to apply the vacuum for 10 minutes after all liquid has passed through the E-Z 96 DNA Plate.

29. Turn off the vacuum.

30. Discard the filtrate and collection plate.

31. Place the E-Z 96 DNA Plate upside down on a stack of paper towels and tap several times to remove any residual ethanol.

Note: It is very important to completely dry the E-Z 96 DNA Plate before elution. If a swing bucket centrifuge with a 96-well plate adaptor is available, centrifuge at 5,000 $\times g$ for 5 minutes to dry the plate. Or if an oven/incubator is available, dry the plate at 70°C for 10 minutes.

32. Place the 96-well Racked Microtubes inside the base of the manifold.

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33. Place the E-Z 96 DNA Plate on top of the manifold.
34. Add 100 μ L Elution Buffer heated to 65°C to each well.
35. Let sit at room temperature for 5 minutes.
36. Turn on the vacuum source to draw the Elution Buffer through the plate.
37. Turn off the vacuum.

Optional: Repeat Steps 34-37 for a second elution step.

Note: 100 μ L Elution Buffer is sufficient to elute up to 85% of the DNA from each well of the E-Z 96 DNA Plate. A second elution step with same 100 μ L elute containing DNA, reheated to 65°C, will increase yield by up to 10-15%. Total DNA yields vary depending on type and quantity of sample.

38. Seal the 96-well Racked Microtubes with Caps for Racked Microtubes (provided).
39. Store DNA at -20°C.

Troubleshooting Guide

Please use this guide to troubleshoot any problems that may arise. For further assistance, please contact the technical support staff, toll free, at **1-800-832-8896**.

Possible Problems and Suggestions

Problem	Cause	Solution
Clogged well	Debris carryover	Following precipitation with SP2 Buffer, make sure no particulate material is transferred.
	Sample too viscous	Do not exceed suggested amount of starting material.
	Incomplete precipitation following addition of SP2 Buffer	Increase RCF or time of centrifugation after addition of SP2 Buffer.
Problem	Cause	Solution
Low DNA yield	Incomplete disruption of starting material	Completely homogenize sample.
	Poor lysis of tissue	Decrease the amount of starting material or increase the amount of SP1 Buffer and SP2 Buffer.
	DNA remains bound to column	Increase elution volume to 200 μ L and incubate the plate at 65°C for 5 minutes before centrifugation.
	DNA washed off	Dilute SPW Buffer by adding appropriate volume of 100% ethanol prior to use (Page 5).
Problem	Cause	Solution
Problems in downstream applications	Salt carryover	Repeat wash step with SPW Buffer.
	Ethanol carryover	Following the second wash spin, ensure that the plate is completely dried before elution.

Ordering Information

The following components are available for purchase separately.
(Call Toll free at 1-800-832-8896)

Product	Part Number
SP1 Buffer (250 mL)	PD072
SP2 Buffer (60 mL)	PD073
SP3 Buffer (100 mL)	PD074
Elution Buffer (100 mL)	PDR048
SPW Buffer (25 mL)	PDR045
RNase A (400 µL)	AC117
Sealing Film	AC1200
AeraSeal Film	AC1201
96-well Square-well Plate (2.2 mL)	EZ9602
RNase A (400 µL)	AC117
E-Z 96 Homogenizer Plates (4 x 96)	HCR9601-02
E-Z 96 Lysate Clearance Plates (10 x 96)	FL9601
Vacuum Manifold	VAC-03

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Qiagen®, QIAvac® and Vacman® are all trademarks of their respected companies.
PCR is a patented process of Hoffman-La Roche. Use of the PCR process requires a license.

For more purification solutions, visit www.omegabiotek.com

AVAILABLE FORMATS



Spin Columns



96-Well Silica Plates



Mag Beads

SAMPLE TYPES



Blood / Plasma



Plasmid



Cultured Cells



Plant & Soil



NGS Clean Up



Tissue



FFPE



Fecal Matter



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