

AccuPrep® Plant Genomic DNA Extraction Kit

Cat. No. K-3031







AccuPrep® Plant Genomic DNA Extraction Kit

Kit for the extraction of total DNA from plants

User Guide

K-3031

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Please read all the information in booklet before using the unit



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Intended Use

AccuPrep[®] Plant Genomic DNA Extraction Kit is developed and supplied for research purposes only. Certain applications possible with this kit may require special approval by appropriate local and/or national regulatory authorities in the country of use.

Safety Warning and Precaution

Wear appropriate protection when handling any irritant or harmful reagents. The use of a laboratory coat, protective gloves and safety goggles are highly recommended. For more information, please consult the appropriate Material Safety Data Sheet (MSDS).

Warranty and Liability

All BIONEER products undergo extensive Quality Control testing and validation. BIONEER guarantees quality during the warranty period as specified, when following the appropriate protocol as supplied with the product. It is the responsibility of the purchaser to determine the suitability of the product for its particular use. Liability is conditional upon the customer providing full details of the problem to BIONEER within 30 days.

Quality Management System ISO 9001 Certified

Every aspect of our quality management system from product development, production to quality assurance and supplier qualification meets the world-class standards.

Trademark

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Contents

Product Information	1
Components	1
Storage	1
Specifications	2
Precautions	2
Introduction	3
Product Description	3
Features & Benefits	4
Experimental Procedures	5
Before You Begin	5
Plant Genomic DNA Extraction	6
Troubleshooting	8
Ordering Information	10
Related Products	10
Explanation of Symbols	11



Product Information

Components

This kit contains adequate reagents for 100 reactions.

Components	Amount	Storage		
Proteinase K powder, lyophilized	25 mg x 2 ea	Refer to the "Storage" below.		
RNase A powder, lyophilized	24 mg x 2 ea	Relei to the Storage below.		
PL Buffer (Plant Lysis)	40 ml			
PC Buffer (Precipitation)	25 ml			
WA1 Buffer (Binding)	40 ml			
W2 Buffer (Washing)	120 ml x 2 ea			
EA Buffer (Elution)	25 ml	Store at room temperature (15-25°C).		
AccuPrep® Binding Column-I	100 ea	,		
Collection Tube (Filtration)	100 ea			
1.5 ml Tube (Elution)	100 ea			
One Page Protocol	1 ea			

Storage

The kit will maintain performance for at least two years under standard storage conditions.

All components of the kit should be stored dry at room temperature (15-25°C). Before use, lyophilized Proteinase K and RNase A should be completely dissolved in 1,250 µl and 600 µl of nuclease-free water, respectively. For short term storage, dissolved Proteinase K and RNase A should be stored at 4°C. For long term storage, it is recommended to aliquot these enzymes into separate tubes and store at -20°C.

* Note: Please note that repeated freezing and thawing may reduce its activity.

Specifications

Sample Type	Amount of Starting Sample	Typical DNA Yield	DNA Purity	
Soybean	100 mg	2-5 µg		
Maize	100 mg	1-4 µg	$A_{260}/A_{280} > 1.7$	
Potato	100 mg	1-4 µg		

^{*} Note: There may be differences in measured values depending on the type of samples.

Precautions

- Take appropriate laboratory safety precautions and wear gloves when handling because WA1 Buffer contains chaotropic salts which are irritants.
- Thoroughly mix PL Buffer and PC Buffer by shaking before use.



Introduction

Product Description

AccuPrep® Plant Genomic DNA Extraction Kit provides a rapid and convenient genomic DNA extraction method. AccuPrep® Plant Genomic DNA Extraction Kit is designed for extraction of total DNA from 100 mg of a variety of botanical sources such as beans and corns. The kit employs spin-column with glass fiber filter for nucleic acid binding in the presence of chaotropic salts. The DNA binds to the silica of glass fiber, while proteins and other contaminants are eliminated through a series of short wash-and-spin steps using ethanol. Finally genomic DNA is eluted by low salt solutions. The process does not require phenol/chloroform extraction and ethanol precipitation.

DNA extracted through this kit can be used for a variety of applications, including: gene cloning, PCR, Real-time PCR, southern blotting, and SNP genotyping.

Features & Benefits

- Comprehensive: High quality and yield of genomic DNA extraction from various plant samples such as leaves, stems, roots, or seeds.
- Convenient & Rapid: The process does not require phenol/chloroform extraction, ethanol precipitation, or other time-consuming steps.
- Efficient: Contaminants such as proteins and nucleases which may interfere with PCR reactions are completely removed.
- Ready-to-use: Extracted DNA is ready-to-use for various application.
- Minimized DNA damage: DNA damage is minimized by avoiding precipitation and use of organic solvents.
- High binding capacity: Uses silica-based DNA binding column with a high binding capacity of up to 30 µg.



Experimental Procedures

Before You Begin

Before proceeding, please check the following:

- 1. Completely dissolve Proteinase K powder in 1,250 μl of nuclease-free water before use. Dissolved Proteinase K should be stored at 4°C.
 - * Note: Do not add Proteinase K directly to PL Buffer.
- 2. Completely dissolve RNase A powder in 600 μl of nuclease-free water before use. Dissolved RNase A should be stored at 4°C.
- 3. Pre-heat EA Buffer at 60°C before use. This process is essential for maximal recovery in filter-based extraction.
- 4. Add indicated volume of absolute ethanol (not provided) to WA1 Buffer before use (see bottle label).

Plant Genomic DNA Extraction

- 1. Grind (or homogenize) ≤ 100 mg of plant sample with a mortar and pestle (or homogenizer) and place it into a clean 1.5 ml tube.
 - * **Note:** If the sample is not ground completely, it will result in significantly reduced DNA yields and clogging of the binding column.
- 2. Add 300 μ l of PL Buffer, 20 μ l of Proteinase K, and 10 μ l of RNase A to the sample from step 1 and mix well by vortexing.
 - * Note: The sample should be completely immersed in the buffer.
- 3. Incubate at 60°C for 10 minutes.
- 4. Add 100 µl of PC Buffer to the lysate and mix well by vortexing.
 - * Note: You should completely resuspend the sample to achieve maximum lysis efficiency.
- 5. Incubate for 5 minutes on ice.
- 6. Centrifuge at 13,000 rpm for 5 minutes.
- 7. Transfer the cleared lysate to a new 1.5 ml tube and add WA1 Buffer. The added buffer should be 1.5 times the volume of the lysate. Mix well by vortexing or pipetting and briefly spin down to collect lysate clinging under the lid.
- 8. Carefully apply the cleared lysate to the *AccuPrep*[®] Binding Column-I fit in a Collection Tube.
- 9. Centrifuge at 8,000 rpm for 1 minute and discard the flow through. Reuse the Collection Tube in step 10.
 - * **Note:** You should put a cap on the column to avoid aerosol formation during centrifugation. If the lysate has not completely passed through the column after centrifugation, centrifuge again at a higher than 10,000 rpm until the column is empty.
- 10. Repeat steps 8-9 with remaining sample.
- 11. Wash the *AccuPrep*® Binding Column-I by adding 500 µI of W2 Buffer.



- 12. Centrifuge at 8,000 rpm for 1 minute and discard the flow through. Reuse the Collection Tube in step 13.
- 13. Repeat steps 11-12.
- 14. Centrifuge once more at 13,000 rpm for 1 minute to remove residual ethanol completely and check whether there is no droplet clinging to the bottom of the binding column.
 - * Note: Residual W2 Buffer in the binding column may cause problems in later applications.
- 15. Place the *AccuPrep*[®] Binding Column-I in a clean 1.5 ml Tube (supplied with a kit). Add 50-200 μl of EA Buffer or nuclease-free water to elute DNA.
 - * **Note:** A smaller volume will result in a more concentrated solution, but total yield may be reduced. For long-term DNA storage, you should elute with EA Buffer and store at -20°C because DNA stored in water is subject to acid hydrolysis. Furthermore, pre-warmed EA Buffer will improve efficiency of elution (see "Before You Begin" on page 5).
- 16. Incubate at room temperature for 1 minute to be absorbed the EA Buffer completely into the glass fiber of the binding column.
 - * Note: To increase DNA yield, you should incubate for 5 minutes after adding EA Buffer.
- 17. Centrifuge at 8,000 rpm for 1 minute.
- 18. To recover more DNA, repeat once more elution step using the eluate from step 17.
- 19. Centrifuge at 8,000 rpm for 1 minute.
 - * **Note:** The eluted genomic DNA is stable and can be used directly or stored at 4°C for future experiment.

Troubleshooting

Problem	Comments			
Low DNA yield or purity	Buffers or other reagents may have been exposed to conditions that reduce their effectiveness. Store the reagents at room temperature (15-25°C) at all times upon arrival and keep all reagent bottles tightly closed after use to preserve pH and stability and avoid contamination. After reconstitution of the lyophilized reagents, divide it into aliquots and store at -20°C.			
	Ethanol may not have been added to WA1 Buffer. Add indicated volume of absolute ethanol (not provided) to the WA1 Buffer (see "Before You Begin" on page 5) and mix well. Mark WA1 Buffer bottle label to indicate whether ethanol has been added or not.			
	Reagents and samples may not have been completely mixed. Always mix the sample tube well after adding each reagent.			
	You may not have used optimal reagents for eluting nucleic acid. An alkaline pH is required for optimal elution. Use EA Buffer included in the kit.			
	The lysis may have been incomplete. Mix the sample immediately after adding Proteinase K. Always mix the sample tube well with ethanol before loading the lysate into the binding column.			
	Glass fiber from binding column may have been eluted with nucleic acid. These fibers can scatter light, resulting in a high absorbance value. In the last stage of elution, too much centrifugation could result in mixing the debris of glass fiber in the binding column into eluate. Please refer to the "Comments" below for instructions on removing the glass fibers.			



Incomplete or no restriction enzyme cleavage of isolated DNA	Glass fiber from binding column may have been eluted with nucleic acid. These fibers may inhibit enzyme reactions. After final elution step, centrifuge the tube at maximum speed for 1 minute. Glass fibers may be visible at the bottom of the tube. Transfer the supernatant to a new tube.
Degraded DNA from plant samples	 There may have been nuclease activity in the unlysed sample. Sample should be frozen at -20°C immediately after harvesting and remained so until lysis step begins. Grind (or homogenize) sample to a fine powder with mortar and pestle (or homogenizer) in liquid nitrogen.
White precipitates in PL Buffer	Buffers may have been stored for prolonged periods of time at lower temperature. Warm the buffers at 60°C and mix well to dissolve the precipitates.

9 BQ-042-101-01 **www.bioneer.com** Revision : 4 (2022-02-14)

Ordering Information

Description	Cat. No		
AccuPrep® Plant Genomic DNA Extraction Kit	K-3031		

Related Products

Description	Cat. No
Proteinase K Powder	KB-0111
RNase A Powder	KB-0101

10 BQ-042-101-01 www.bioneer.com Revision : 4 (2022-02-14)



Explanation of Symbols

LOT	Batch Code	Ţ <u>i</u>	Consult Instructions For Use	RUO	Research Use Only	\triangle	Caution
&	Biological Risks	Σ	Contains Sufficient for <n> tests</n>	1	Temperature Limitation		Manufacturer
REF	Catalog Number	2	Do not Re-use	\subseteq	Use-by Date		

11 BQ-042-101-01 www.bioneer.com Revision : 4 (2022-02-14)

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