



NZY Plant/Fungi gDNA Isolation kit

Catalogue numbers:

MB17701, 50 columns

MB17702, 4 x 50 columns

Description

NZY Plant/Fungi gDNA Isolation kits are designed for the simple and rapid small-scale preparation of highly pure genomic DNA from plant tissues and fungi samples. The method is spin column silica-based and requires no phenol or chloroform extraction. This kit uses two optimized lysis buffers systems based on the established CTAB and SDS methods. NZY Homogenization columns are included in this kit for conveniently clearing the crude lysates. The clear flow-through is mixed with binding buffer PN for optimal binding of DNA to the silica membrane. Then, the DNA is selectively absorbed into the NZYSpin Plant Column and others impurities such as proteins and salts are removed during the washing steps. The eluted genomic DNA has a $A_{260/280}$ ratio between 1.7 and 1.9 what makes it ready to use in applications like sequencing, PCR, multiplex-PCR, genotyping and a wide range of other enzymatic manipulations.

The NZY Plant gDNA Isolation kit is optimized to isolate up to 30 μg of DNA from up to 100 mg (wet weight) or 20 mg (dry weight) of plant/fungi samples, depending of the plant/fungi species. We suggest not using more than the recommended starting material to prevent reductions in yield and purity.

Storage conditions and reagents preparation

All kit components can be stored at room temperature (15-25 °C) and are stable till the expiry date. Before use, add 100 mL of ethanol (96-100%) to Buffer PNW2 bottle. Add 0.6 mL of water to the RNase A vial. Store the RNase A solution at 4 °C for up to 3 months. For longer storage (up to 1 year), the RNase A solution should be divided into small aliquots and stored at -20 °C. Buffers PNL1, PNL2 and PN may form a precipitate when stored at low temperatures, if necessary, dissolve the precipitate by warming the solutions at 37 °C. Buffers PNL1, PNL2, PN and PNW1 contain guanidine hydrochloride and/or detergents like CTAB or SDS. Wear gloves and goggles when using this kit. DO NOT add bleach or acidic solutions directly to the sample-preparation waste.

System Components

Component	50 columns
Buffer PNL1	22 mL
Buffer PNL2	20 mL
Buffer PNL3	5 mL
Buffer PN	25 mL
Buffer PNW1	22 mL
Buffer PNW2 (concentrate)	25 mL
Buffer PNE	7 mL
RNase A (lyophilized)	6 mg
NZYSpin Homogenization columns (purple rings)	50
NZYSpin Plant columns (green rings)	50
Collection tubes (2 mL)	100

Standard protocol for isolating genomic DNA

NZY Plant/Fungi gDNA Isolation kits include two different lysis buffers for optimal results with most common plant species. Please, see Table 1 (in appendix) for choosing the optimal lysis buffer system for your individual plant/fungi sample.

A. Genomic DNA from plant

1. Sample preparation

Homogenize up to 100 mg wet weight or up to 20 mg dry weight (lyophilized) plant material. Proceed with cell lysis using Buffer PNL1 (step 2a) or Buffer PNL2 (step 2b).

2a. Cell lysis (Buffer PNL1)

Transfer the resulting powder to a new tube and add 400 μ L Buffer PNL1. Mix thoroughly by vortex.

Note: If the samples are difficult to lyse, additional Buffer PNL1 can be added. Note that the volumes of RNase A (step 2a) and buffer PN (step 4) must be increased proportionally.

Add 10 μ L of RNase A solution and mix sample thoroughly. Incubate the suspension for 10 min at 65 $^{\circ}$ C.

Note: For some plant samples, it might be advantageous to increase the incubation time to 30-60 min.

Proceed with step 3.

2b. Cell lysis (Buffer PNL2)

Transfer the resulting powder to a new tube and add 300 μL Buffer PNL2. Mix thoroughly by vortex.

Note: *If the samples are difficult to lyse, additional Buffer PNL2 can be added. Note that the volumes of RNase A, Buffer PNL3 (step 2b) and Buffer PN (step 4) must be increased proportionally.*

Add 10 μL of RNase A solution and mix sample thoroughly. Incubate the suspension for 10 min at 65 $^{\circ}\text{C}$.

Note: *For some plant samples, it might be advantageous to increase the incubation time to 30-60 min.*

Add 75 μL of Buffer PNL3 and mix sample thoroughly. Incubate for 5 min on ice to precipitate SDS completely.

Proceed with step 3.

3. Clarification of crude lysate

Transfer the mixture from step 2 into a NZYSpin Homogenization column (purple ring) placed in a new 2 mL collection tube. Centrifuge for 2 min at $>11,000 \times g$, collect the clear flow-through and discard the NZYSpin Homogenization column.

Notes:

If not all liquid has passed the filter, repeat the centrifugation step.

If insoluble particles are visible in the flow-through, transfer the clear supernatant to a new 1.5 mL microcentrifuge tube. Alternatively, centrifuge the crude lysate for 5 min at full speed and transfer the supernatant to a new microcentrifuge tube or pass the precleared supernatant through the NZYSpin Homogenization column to remove the insoluble particles completely.

4. Adjust DNA binding conditions

Add 450 μL of Buffer PN and mix thoroughly by pipetting up and down for 5 times.

5. DNA binding

Place a NZYSpin Plant column (green ring) into a new collection tube and load 700 μL of the sample (maximum loading capacity of NZYSpin Plant column).

Centrifuge for 1 min at $> 11,000 \times g$ and discard flow-through. For higher sample volumes repeat the loading step.

6. Wash silica membrane

Add 400 μL of Buffer PNW1 to the NZYSpin Plant column. Centrifuge for 1 min at $> 11,000 \times g$. Discard flow-through and place the column back into the collection tube.

Add 700 μL of Buffer PNW2 (make sure ethanol was previously added) to the NZYSpin Plant column and centrifuge for 1 min at $> 11,000 \times g$. Discard flow-through.

Add another 200 μL of Buffer PNW2 to the NZYSpin Plant column and centrifuge for 2 min at $> 11,000 \times g$ to remove wash buffer and dry the silica membrane completely.

7. Elute DNA

Place the NZYSpin Plant column into a clean microcentrifuge tube and add 50 μL of Buffer PNE directly in the membrane column (preheating of elution buffer to 65 $^{\circ}\text{C}$ may improve yield). Incubate 1 min at room temperature and centrifuge at $> 11,000 \times g$ for 2 min to elute DNA.

Repeat this step with another 50 μL of Buffer PNE (65 $^{\circ}\text{C}$) and elute into the same tube.

The genomic DNA can be stored at 4 $^{\circ}\text{C}$ or -20 $^{\circ}\text{C}$.

B. Genomic DNA from fungi

1. Sample preparation

Wash 50-200 mg mycelium (fresh weight) or material from fruiting body of macro fungi in ethanol. Mycelium can be obtained from a liquid culture or scraped off (with or without agar) from the surface of a solid medium. Cover sample completely with ethanol and mix carefully. Short washing in ethanol is sufficient in most cases, although incubation overnight sometimes increases DNA yield.

Remove the ethanol by pipetting and squeezing the mycelium.

2. Cell lysis

Place the sample into a 1.5 mL microcentrifuge tube. Add 150 mg siliconized glass beads or sea sand and 200 μL of Buffer PNL1. Homogenize sample using a micro pestil and vortex regularly. Add additional 100 μL of Buffer PNL1 and continue to homogenize the sample.

Note:

If the samples cannot be handled easily, additional buffer PNL1 can be added. Note that the volume of Buffer PN (step 4) must be increased proportionally. If the samples are rich in RNA or protein, we recommend adding 10 μL of RNase A and/or Proteinase K (5-10 mg/mL stock solution) to the PNL1 lysis solution to minimized contaminants.

Incubate for 10 min at 65 $^{\circ}\text{C}$. For some fungi, it might be advantageous to increase the incubation time to 30-60 min.

Add 100 μL chloroform, vortex for 10 seconds and separate phases by centrifugation for 15 min at 20,000 $\times g$. Pipette the top aqueous layer into a new 1.5 mL microcentrifuge tube.

Proceed with section A, step 3.

Quality control assay

Functional assay

All components of NZY Plant/Fungi gDNA Isolation kit are tested following the isolation protocol described above. The purification system must isolate 1-30 µg of gDNA/column, depending of the tested species.

V2401

Certificate of Analysis

Test	Result
Functional assay	Pass

Approved by:



Patrícia Ponte
Senior Manager, Quality Systems

For research use only

Appendix

Table 1. Plant species tested with NZY Plant and Fungi isolation kit.

Plant Species	Plant tissue/organ	Lysis buffer successfully tested	
		PNL1	PNL2
<i>Abies alba</i> (fir)	Needle	Ok	Ok
<i>Amorphorphallus titanium</i>	Leaf	Ok	Not tested
<i>Apium graveolens</i> (celery)	Corm	Ok	Ok
<i>Arabidopsis thaliana</i>	Leaf	Ok	Ok
<i>Boreava orientalis</i>	Leaf, herbarium sample	Ok	Ok
<i>Cleisostoma racemiferum</i>	Inflorescence rachis, silica-gel dried	Ok	Not tested
<i>Doritis pulcherrima</i>	Leaf, silica-gel dried	Ok	Not tested
<i>Eichornia azurea</i>	Leaf	Ok	Not tested
<i>Encephalartos natalensis</i>	Leaf	Ok	Not tested
<i>Galium aparine</i>	Leaf	Ok	Ok
<i>Hordeum</i> sp. (barley)	Leaf	Ok	Ok
<i>Isatis kotschyana</i>	Leaf, herbarium sample	Ok	Ok
<i>Laurus azorica</i> (laurel)	Leaf	Ok	Not tested
<i>Lupinus</i> sp. (lupin)	Leaf	Ok	Ok
<i>Lycopersicon esculentum</i> (tomato)	Stem	Ok	Ok
<i>Myagrum perfoliatum</i>	Leaf, herbarium sample	Ok	Ok
<i>Oryza sativa</i> (rice)	Leaf	Ok	Ok
<i>Persea feru./caerulea</i>	Leaf	Ok	Not tested
<i>Pteridium</i> sp.	Leaf	Ok	Not tested
<i>Pterocarya fraxinifolia</i>	Leaf	Ok	Not tested
<i>Rosa</i> sp. (rose)	Leaf	Ok	Ok
<i>Rubus fruticosus</i> (blackberry)	Leaf	Ok	Ok
<i>Sameraria nummularia</i>	Leaf, herbarium sample	Ok	Ok
<i>Secale</i> sp. (rye)	Leaf	Ok	Ok
<i>Stereochilus</i> sp.	Leaf, silica-gel dried	Ok	Not tested
<i>Tauscheria lasiocarpum</i>	Leaf, herbarium sample	Ok	Ok
<i>Trachycarpus takil</i>	Leaf	Ok	Not tested
<i>Trichoglottis</i> sp.	Leaf, silica-gel dried	Ok	Not tested
<i>Triticum aestivum</i> (wheat)	Leaf	Ok	Ok
<i>Vigna radiate</i> (mung bean)	Root	Ok	Ok
<i>Zea mays</i> (maize)	Leaf	Ok	Ok
<i>Zea mays</i> (maize)	Grain, dried, ground coarsely	Ok	Ok
Fungal mycel (not specified)		Ok	Not tested
Green algae (not specified)		Ok	Not tested



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