

Manual

Genomic Micro AX Plant Gravity

Gravity flow kit for genomic DNA purification from plant tissue.

catalog#	size
103-100	100 isolations

For research use only.

Guarantee

 $A\&A\ Biotechnology\ provides\ a\ guarantee\ on\ this\ product.$

The company does not guarantee the correct performance of this kit in the event of:

- not adhering to the supplied protocol
- use of not recommended equipment or materials
- use of other reagents than recommended or which are not a component of the product
- use of expired or improperly stored product or its components

Table of Contents

Contents	3
Additional equipment and reagents	3
Necessary	3
Optional	3
Important information	4
Isolation protocol	4
Lysate flow rate troubleshooting	ć
DNA neutralization	ć
E buffer functionality test	ć
Gravity flow technology	7
Safety information	7

Contents

component	100 isolations	storage
Micro AXD columns	100 pcs	2-8 °C
Gravity tubes	100 pcs	15-25 ℃
LS lysis suspension	100 ml	15-25 ℃
K1G equilibrating solution	60 ml	15-25 ℃
W1G first wash solution	70 ml	15-25 ℃
W2 second wash solution	60 ml	15-25 ℃
E elution buffer (without EDTA)	20 ml	2-8°C
N neutralizing buffer	1 ml	15-25 ℃
Tsolution	400 μΙ	2-8 °C
Proteinase K	2 x 1.1 ml	4-8°C

The binding capacity of the column is $20 \,\mu g$.

Additional equipment and reagents

Necessary

- 1.5 ml, 2 ml sterile Eppendorf tubes
- Incubator or thermoblock set to 50 °C
- Centrifuge, vortex

Optional

- RNAse (cat. # 1006-10, 1006-50)
- Gravity flow rack (cat. # 008-1)

Important information

 E elution buffer loses activity upon prolonged contact with air. Always close the E elution buffer vial tightly directly after use. Store E elution buffer at 2-8 °C.

Isolation protocol

- Transfer up to 20 mg of dried, powdered plant material or up to 100 mg fresh / frozen cut plant tissue to a 2 ml Eppendorf tube (not included).
- 2. LS lysis suspension should be mixed by inverting the tubes before use.

Add $900 \,\mu l$ of LS lysis suspension and $20 \,\mu l$ of proteinase K.

3. Vortex and incubate for 10 min at 50 °C. Vortex the sample a few times.

The incubation step can be performed in Eppendorf Thermomixer or analogous equipment at 1400 RPM and 50 °C.

RNA digestion (optional): add 5 µl of RNAse (10 mg/ml solution) (not included). Mix and incubate for 5 min at room temp.

4. During incubation prepare the Micro AXD columns by securely attaching them to the top of Gravity tubes and placing them upright in the rack.



Reference photo:

placing the columns and receiving tubes in the Gravity flow rack.

5. Apply 500 µl of K1G equilibrating solution onto the Micro AXD column. Wait for the solution to flow through the column.

It is a good practice to apply the K1G solution to the column wall to avoid accidental blockage of the column flow by an air bubble between the membrane and the K1G solution.

The column is ready for use when the solution stops dripping from the capillary.

6. After incubation, vortex the sample for 2 min at 1000-1500 RPM.

This is a key step for DNA efficiency.

Centrifuge for 5 min at 10 000-14 000 RPM to remove residual material from the tube caps.

Apply the sample onto the equilibrated Micro AXD column.
 Wait 10 min for the lysate to flow through the column by gravity.

The flow rate strongly depends on DNA concentration in the sample. The more DNA, the slower the flow rate. As soon as the lysate stops dripping, proceed to the next step.

Lysate flow rate troubleshooting - page 6.

Note: applies to points 7-9 of the isolation protocol

- In the case of DNA isolation from a smaller number of samples (up to 10), it should be observed whether the
 lysate has completely passed through the column. When the solution stops dripping from the capillary,
 proceed to the next step in the isolation protocol.
- In the case of DNA isolation from a larger number of samples (over 10), we recommend waiting up to 10 min, instead of observing the process in individual columns.
- 8. Apply 600 µl of W1G first wash solution onto the Micro AXD column. Wait for the solution to flow through the column.
- Apply 500 µI of W2 second wash solution onto the Micro AXD column.
 Wait for the solution to flow through the column.
- 10. Before using E buffer, it is recommended to do a functionality test page 6.

Apply $60 \mu l$ of E elution buffer onto the Micro AXD column. Keep for $5 \min$ at room temp.

E elution buffer loses activity upon prolonged contact with air. Always close the E elution buffer vial tightly directly after use. Store E elution buffer at 2-8 °C.

The purpose of this step is to decrease the total volume of eluate, since the column void volume is about 60 µl.

11. Prepare the 1.5 ml elution tubes (not included).

Apply $5 \mu I$ of N neutralizing buffer onto the bottom of each tube.

DNA neutralization - page 6.

- 12. Transfer the Micro AXD columns to the prepared elution tubes.
- 13. Before using E buffer, it is recommended to do a functionality test page 6.

Apply 120 µl of E elution buffer onto the Micro AXD column. Wait 10 min for the buffer to flow through the column by gravity.

E elution buffer loses activity upon prolonged contact with air. Always close the E elution buffer vial tightly directly after use. Store E elution buffer at 2-8 °C.

After 10 min check that the buffer has passed through the column. If not, this indicates a very large amount of DNA in the sample. In this case, it is recommended to centrifuge the sample (column within tube) for 30-60 s at 5000 RPM.

14. Remove the Micro AXD column. Close the tube with purified DNA and store until later use.

Lysate flow rate troubleshooting

problem	reason	solution
very slow rate of lysate through the Micro AXD column	highly concentrated DNA in the sample	- place the Micro AXD column into an Eppendorf tube and centrifuge. - for the next isolation, reduce the amount of sample by half.
air bubbles in the receiving tube capillary	the Micro AXD column is not not securely attached to the receiving tube	- "tightening" the Micro AXD column reattach the column in luer-like fitting simultaneously by pressing the column down and twisting.

DNA neutralization

E elution buffer is strongly alkaline and may cause DNA degradation upon freezing. Thus it is necessary to use a N neutralizing buffer. We recommend adding the N neutralizing buffer to the elution tube before the elution step. If the N neutralizing buffer was not added before the elution step it can be added directly before freezing DNA samples. The use of N neutralizing buffer enables secure DNA storage conditions at 10 mM TrisHCl, pH 8.5.

E buffer functionality test

E buffer has a critical influence on DNA elution efficiency and thus overall DNA purification yield. The kit contains T solution which enables testing of the E buffer correct functionality.

Typically it is suggested to perform such a test in the following cases:

- E buffer was not used for a long period of time (at least 2 months).
- E buffer was stored at room temp. for a long period of time (at least 2 weeks).
- E buffer vial was not closed tightly.

Procedure:

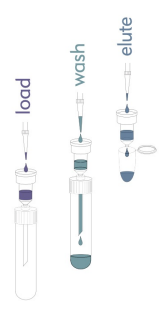
Transfer 20 µl of E buffer to PCR tubes; add 2 µl of T solution; mix the sample, wait 2 min. Compare the mixture color with the reference color guide.



Buffer E

Buffer E

Gravity flow technology



Safety information



Proteinase K

H315 Causes skin irritation.

H319 Causes serious eye irritation.

H334 May cause allergy or asthma symptoms or breathing difficulties if inhaled.

H335 May cause respiratory irritation.

P261 Avoid breathing dust.

P305+P351+P338 If in eyes: rinse cautiously with water for several minutes. Remove contact lenses,

if present and easy to do. Continue rinsing.

P342+P311 If experiencing respiratory symptoms call a Poison Center or doctor/physician.



LS lysis suspension

H302 Harmful if swallowed.

H315 Causes skin irritation. H319 Causes serious eye irritation.

P305+P351+P338 If in eyes: rinse cautiously with water for several minutes. Remove contact lenses,

if present and easy to do. Continue rinsing.



E elution buffer

H314 Causes severe skin burns and eye damage.

P280 Wear protective gloves/ protective clothing/ eye protection/ face protection.

 $P305 + P351 + P338 \ If in eyes: rinse cautiously with water for several minutes. Remove contact lenses, and the property of the property of$

if present and easy to do. Continue rinsing.

P310 Immediately call a Poison Center or doctor/physician.



A&A Biotechnology, ul. Strzelca 40, 80-299 Gdańsk, Poland phone +48 883 323 761, +48 600 776 268 info@aabiot.com, www.aabiot.com