

SUPPLEMENTARY PROTOCOL

Ver 2.0

KIT-9230

PrimeMag Plant Genomic DNA Extraction Kit

A supplementary protocol is an additional guide provided alongside the main protocol to assist users in applying the product to special or less common sample types, workflows, or conditions. It expands the usability of the main kit by offering validated methods for alternative applications while maintaining performance and quality standards. The support protocol does not replace the main protocol, but serves as supplementary guidance.





List of Supplementary Protocol for KIT-9230

1. **New sample type:** Mushroom
2. **Optional column:** To filter lysate



1. New sample type: Mushroom

Materials Supplied by Users

- ✓ Centrifuge at speed of 10,200 rpm – 14,800 rpm
- ✓ Thermomixer with temperature settings of 65 °C, 56 °C and room temperature (DLAB HM100-Pro Thermo Mix or equivalent)
- ✓ Magnetic Rack (1st BASE, KIT-MAG16A)
- ✓ Vortex mixer
- ✓ 2 mL microcentrifuge tubes
- ✓ 1.5 mL microcentrifuge tubes
- ✓ Isopropanol (IPA)
- ✓ Absolute ethanol (≥ 99.5%)



Protocol

Preparation

- I. Prepare 50 mg (up to 150 mg) of pulverized mushroom using liquid nitrogen.

Tips:

- ✓ *To obtain the best result from plant nucleic acid extraction, bring liquid nitrogen to plant field to snap-freeze the plant tissue immediately during sample collection before process in laboratory.*
- ✓ *Pulverize plant tissue in liquid nitrogen using a set of sterilized mortar and pestle.*
- ✓ *Replenish the liquid nitrogen in the mortar 2 to 3 times and continue to grind sample until a fine, homogenous powder is obtained.*

- II. Set thermomixer to 65 °C.

- III. Freshly prepare 80% ethanol.

Example: For 1 prep, to prepare 600 μ L 80% ethanol, add 480 μ L into 120 μ L of dH₂O

- IV. Add absolute ethanol ($\geq 99.5\%$) to WB1 Buffer as following:

Kit Preps	WB1 Buffer	Ethanol to be added
10 preps	4 mL	6 mL
48 preps	16 mL	24 mL
96 preps	30 mL	45 mL



Lysis

- Transfer 50 mg of pulverized mushroom sample into 2 mL tube. Add **700 µl Plant Lysis Buffer**, **50 µl PG Buffer** and **10 µl RNase A into the sample tube**. Mix the sample by gently inverting the tube 10 times.
Note: Always add Plant Lysis Buffer and PG Buffer separately and never premix the solution.
- Incubate the mixture at 65 °C for 30 minutes in Thermo Mix (e.g. DLAB HM100-Pro) while shaking at 1,000 rpm.

Binding

- Vortex the **PlantMag Beads** for 30 seconds to ensure they remain in suspension before use.
- Prepare the **binding mixture (BindMix)** in a 2 mL microcentrifuge tube as following:

Components	BindMix Volume (per prep)
Isopropanol (IPA)	360 µL
BD2 Buffer	120 µL
PlantMag Beads	30 µL

Note: Mix isopropanol and Binding Buffer just before use. The freshly prepared mixture must be used within 1 hour.

- After 30 minutes of incubation from step 2, centrifuge the lysate at 16,160 x g at room temperature for 5 minutes. Change the Thermo Mix to room temperature.
- Leave the tube (with centrifuged lysate) at room temperature for 10 - 15 minutes or until the top layer appears.
- Transfer **480 µL of supernatant** (middle layer) to the **BindMix** tube.



Binding

- Mix the mixture in the **BindMix** tube by pipetting up and down for 10 times and shake at 1,000 rpm in room temperature, for 5 minutes in Thermo Mix.
- Place the tube on a magnetic rack (1st BASE, KIT-MAG16A) for 2 minutes, or until the **PlantMag Beads** separation has been completed.
- Remove and discard the supernatant.

Washing

- Add **600 μ L WB1 Buffer** (ethanol added) into the **PlantMag Beads** tube.
- Shake at 1, 000 rpm in room temperature, for 5 minutes in Thermo Mix.
- Place the tube on a magnetic rack for 1 minutes or until the **PlantMag Beads** separation has been completed.
- Remove and discard the supernatant.
- Add **600 μ L 80% ethanol** into the **PlantMag Beads** tube.
- Shake at 1, 000 rpm in room temperature for 5 minutes in Thermo Mix.
- Place the tube on a magnetic rack for 1 minute or until the **PlantMag Beads** separation has been completed.
- Remove and discard the supernatant.



Washing	<p>19. Air-dry the PlantMag Beads at room temperature for 5 to 10 minutes.</p> <p>Note: <i>DO NOT over dry the beads. Over drying could result in lower DNA yield.</i></p>
Elution	<p>20. Remove PlantMag Beads tube from the magnetic rack. Add 100 μL of Elution Buffer into the PlantMag Beads tube.</p> <p>21. Shake at 1,000 rpm at 56 °C for 5 minutes.</p> <p>22. Place the tube on a magnetic rack for 1 minute or until the PlantMag Beads separation has been completed.</p> <p>23. Transfer the supernatant with purified DNA to a new 1.5 mL tube.</p> <p>Optional: <i>If sediment is observed in the DNA eluent, centrifuge the sample at a maximum speed of 16,160 \times g for 1 minute. Carefully transfer the clear supernatant to a new 1.5 mL microcentrifuge tube.</i></p> <p>Note: <i>Execution of this optional step may result in reduced DNA yield.</i></p>



2. Optional column: To filter lysate

Materials Supplied by Users

- ✓ Centrifuge at speed of 10,200 rpm – 14,800 rpm
- ✓ Thermomixer with temperature settings of 65 °C, 56 °C and room temperature (DLAB HM100-Pro Thermo Mix or equivalent)
- ✓ Magnetic Rack (1st BASE, KIT-MAG16A)
- ✓ biocomma® Filtration column (biocomma®, FC0015-PES-45W)
- ✓ Vortex mixer
- ✓ 2 mL microcentrifuge tubes
- ✓ 1.5 mL microcentrifuge tubes
- ✓ Wide bore tips
- ✓ Isopropanol (IPA)
- ✓ Absolute ethanol (≥ 99.5%)



Protocol

Preparation

- I. Prepare 50 mg of plant tissue using liquid nitrogen.

Tips:

- ✓ *To obtain the best result from plant nucleic acid extraction, bring liquid nitrogen to plant field to snap-freeze the plant tissue immediately during sample collection before process in laboratory.*
- ✓ *Pulverize plant tissue in liquid nitrogen using a set of sterilized mortar and pestle.*
- ✓ *Replenish the liquid nitrogen in the mortar 2 to 3 times and continue to grind sample until a fine, homogenous powder is obtained.*

- II. Set thermomixer to 65 °C.

- III. Freshly prepare 80% ethanol.

Example: For 1 prep, to prepare 600 μ L 80% ethanol, add 480 μ L into 120 μ L of dH₂O

- IV. Add absolute ethanol ($\geq 99.5\%$) to WB1 Buffer as following:

Kit Preps	WB1 Buffer	Ethanol to be added
10 preps	4 mL	6 mL
48 preps	16 mL	24 mL
96 preps	30 mL	45 mL



Lysis

- Transfer 50 mg of pulverized plant sample into 2 mL tube. Add **700 μ L Plant Lysis Buffer**, **50 μ L PG Buffer** and **10 μ L RNase A** into the sample tube. Mix the sample by gently inverting the tube 10 times.
Note: Always add Plant Lysis Buffer and PG Buffer separately and never premix the solution.
- Incubate the mixture at 65 °C for 30 minutes in Thermo Mix (e.g. DLAB HM100-Pro) while shaking at 1,000 rpm.

Binding

- Vortex the **PlantMag Beads** for 30 seconds to ensure they remain in suspension before use.
- Prepare the **binding mixture (BindMix)** in a 2 mL microcentrifuge tube as following:

Components	BindMix Volume (per prep)
Isopropanol (IPA)	360 μ L
BD2 Buffer	120 μ L
PlantMag Beads	30 μ L

- Note:** Mix isopropanol and Binding Buffer just before use. The freshly prepared mixture must be used within 1 hour.
- After 30 minutes of incubation from step 2, transfer the **entire lysate** with wide bore tips into the **filter tube** (e.g. biocomma® Filtration Columns, FC0015-PES-45W) and centrifuge at 16,160 x g at room temperature for 1 minute to 5 minutes. Change the Thermo Mix to room temperature.
 - Transfer **480 μ L** of filtrate to the **BindMix** tube.



Binding	<ol style="list-style-type: none">7. Mix the mixture in the BindMix tube by pipetting up and down for 10 times and shake at 1,000 rpm in room temperature, for 5 minutes in Thermo Mix.8. Place the tube on a magnetic rack (e.g. 1st BASE, KIT-MAG16A) for 2 minutes, or until the PlantMag Beads separation has been completed.9. Remove and discard the supernatant.
Washing	<ol style="list-style-type: none">10. Add 600 μL WB1 Buffer (ethanol added) into the PlantMag Beads tube.11. Shake at 1, 000 rpm in room temperature, for 5 minutes in Thermo Mix.12. Place the tube on a magnetic rack for 1 minute or until the PlantMag Beads separation has been completed.13. Remove and discard the supernatant.14. Add 600 μL 80% ethanol into the PlantMag Beads tube.15. Shake at 1, 000 rpm in room temperature for 5 minutes in Thermo Mix.16. Place the tube on a magnetic rack for 1 minute or until the PlantMag Beads separation has been completed.17. Remove and discard the supernatant.18. Air-dry the PlantMag Beads at room temperature for 5 to 10 minutes. Note: <i>DO NOT over dry the beads. Over drying could result in lower DNA yield.</i>



Elution

19. Remove **PlantMag Beads** tube from the magnetic rack. Add **100 μ L** of Elution Buffer into the **PlantMag Beads** tube.
20. Shake at 1,000 rpm at 56 °C for 5 minutes.
21. Place the tube on a magnetic rack for 1 minute or until the **PlantMag Beads** separation has been completed.
22. Transfer the supernatant with purified DNA to a new 1.5 mL tube.



Product Ordering Information

Product Number	Product Description	Remarks
KIT-9230-10	PrimeMag Plant Genomic DNA Kit	Sufficient for 10 preps.
KIT-9230-48	PrimeMag Plant Genomic DNA Kit	Sufficient for 48 preps.
KIT-9230-96	PrimeMag Plant Genomic DNA Kit	Sufficient for 96 preps.
KIT-MAG16A	1st BASE Magnetic Rack	For 16 tubes processing.



Document Revision History

Revision	Date of Revision	Description of Change
2.0	22-May-26	<ol style="list-style-type: none">1. Revision of “Supplementary protocol” main description.2. Addition of supplementary protocol “Optional column: To filter lysate ”.