

magnetiC Plant DNA Extraction Kit

Quick Start Guide



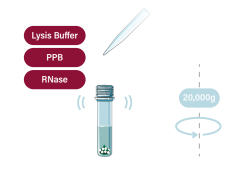
Add up to 50mg of ground fresh or dried plant leaves to the lysis bead tube provided.

Add 600µl of Lysis Buffer, 60µl PPB, and 5µl RNase.

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Mix for 3 mins using TissueLyser at max speed or vortex for 10 mins; then centrifuge at 20,000g for 2 mins.



Avoiding pellet, transfer up to 300-400µl of supernatant to clean centrifuge tube.

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Add 1000µl of Binding Buffer, vortex for 10-20s, and wait 5 mins.





Place tube on magnetic rack to capture. Wait 1 min then discard supernatant.

Add 600ul of Wash Buffer #1 to the tube and vortex for 10–20s.

Wait 1 min then place tube on magnetic rack to capture. Wait 1 min then discard supernatant.







