

# Kleargene

## Quick guide: Kleargene plant 96-well plate DNA extraction kit



### Sample disruption

- Homogenise samples, + 75 µL buffer L1 ●, incubate (55°C / 10 min)
- + 150 µL buffer C1 ●, mix, incubate (55°C / 10 min)
- Centrifuge (3,000 x g / 2 min) to clear lysate

### DNA binding

- Place filter plate on top of collection plate/ reservoir
- Transfer cleared lysate to filter plate wells, incubate at room temperature for 2 min
- Centrifuge (3,000 x g / 2 min) then clear collection plate

### 1st wash

- + 150 µL buffer A1 ● to filter plate wells, incubate at room temperature for 2 min
- Centrifuge (3,000 x g / 2 min) then clear collection plate

### 2nd wash

- + 75 µL buffer W1 ● to filter plate wells, incubate at room temperature for 2 min
- Centrifuge (3,000 x g / 2 min) then clear collection plate

### Ethanol wash

- + 75 µL 100% EtOH to filter plate wells, incubate at room temperature for 2 min
- Centrifuge (3,000 x g / 2 min) then clear collection plate

### Drying & elution

- Incubate filter plate (55°C / 10 min), then place on an elution plate
- + 75 µL buffer E1 ● (@ 55°C) to filter plate wells, incubate (55°C / 5min)
- Centrifuge (3,000 x g / 2 min)

Need help, please contact our technical support team on  
+44 (0)1992 470 757 or email us at [tech.support@lgcgenomics.com](mailto:tech.support@lgcgenomics.com)

[www.lgcgenomics.com](http://www.lgcgenomics.com)

Germany • United Kingdom • United States