DNA extraction protocol (QIAGEN DNeasy96)

Chang-Yu Chang

2021-02-08

Aims

Extract DNA from pellets in 96-well plates.

Materials

- PBS 36 mL
- proteinase K 2.5 mL
- $\bullet~$ buffer AL 20 mL
- ethanol 20 mL
- $\bullet~$ buffer AW1 50 mL
- buffer AW2 50 mL
- $\bullet~$ buffer AE 10 mL
- DNeasy96 plate 1
- S-block or DW96 1
- PCR plate 1

Procedures

Thaw pellet plates at room temperature.
Dissolve pellets in $180\mu L$ of PBS.
Transfer the dissolved pellets to DNeasy microtube plates.
Add $25\mu L$ proteinase K and $200\mu L$ Buffer AL.
Seal the plate with caps and mix by vortexing. Incubate at 56C for 30 min.
Add $200\mu L$ pure ethanol.
Seal the plate with caps and mix thoroughly by vortexing.
Assemble the DNeasy 96 plates on top of DW96s. Mark the DNeasy 96 plates.
Use mP1000 set at $625\mu L$ to transfer the lysate of each sample from DW96s to each well of DNeasy96
plates (maximum $900\mu L$).
Seal DNeasy96 with AirPore film. Use vacuum to collect the flow-through.
Add $500\mu L$ of Buffer AW1 to each sample.
Seal DNeasy96 with AirPore film. Use vacuum to collect the flow-through.
Add $500\mu L$ of Buffer AW2 to each sample.
Seal DNeasy96 with AirPore film. Use vacuum to collect the flow-through.
Label two PCR plates. Place DNeasy96s on these PCR plates.
Add $100\mu L$ Buffer AE to each sample, and seal the DNeasy 96 plates with new AirPore Tape Sheets
(provided).
Incubate for 1 min at room temperature (15–25°C). Centrifuge for 2 min at 6000 rpm.
Cover the PCR plates with aluminum foil and save it in -20C freezer.