

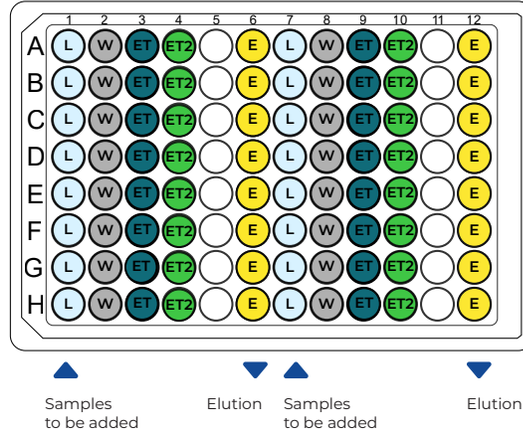
Pathogen/ Viral Nucleic Acid Isolation Kit - Instructions for use with the NorMag16

1. Prepare Lysis-bead mix

Prepare a bulk mix of Lysis/ Binding/ proteinase K as in the table below. We suggest calculating a 10 % overage to compensate for pipetting errors and inaccuracies. Vortex the beads and proteinase K before use, and vortex the mixture before dispensing to ensure that the magnetic beads are resuspended.

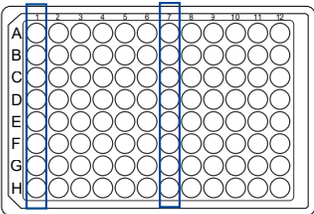
| Solution | Volume per sample | Volume for 16 samples |
|-----------------------------|-------------------|-----------------------|
| Nucleic Acid Magnetic Beads | 20 µL | 352 µL |
| Lysis Buffer B | 300 µL | 5.28 mL |
| Proteinase K | 10 uL | 176 µL |

2. Add Reagents as in Diagram Below

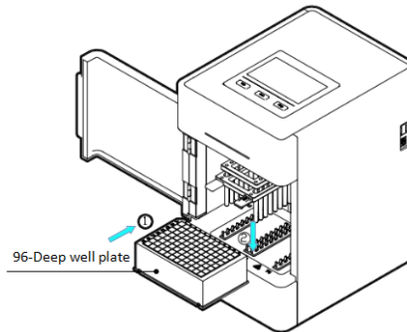


- L Lysis-bead mix (330µl)
Columns 1 & 7
- W Wash WN (1 mL)
Columns 2 & 8
- ET 80% Ethanol (1 mL)
Columns 3 & 9
- ET2 80% Ethanol (500µl)
Columns 4 & 10
- E Elution Buffer F (50-100µl)
Columns 6 & 12
- Leave Empty
Columns 5 & 11

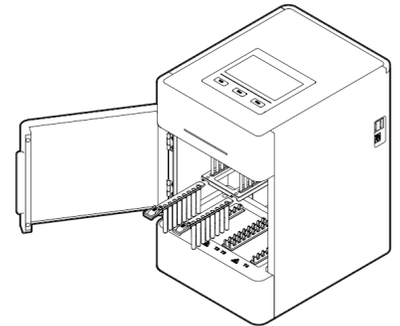
3. Add up to 16 samples to lysis binding mix (200µL).



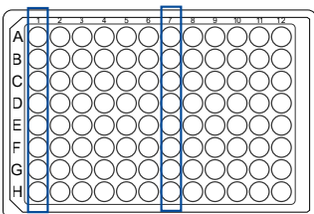
4. Place plate on machine. Ensure connect orientation.



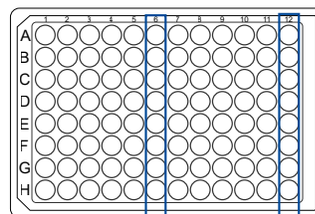
5. Insert two combs and select the protocol NormagVTNA & press run.



7. At pause remove the plate & add 500 µL 100% ethanol to the samples. Replace the plate and press run.



8. At the end of the run, columns 6 & 12 contain the purified DNA/RNA elution.



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