

Nanotrap[®] Microbiome A; Automated Protocol using NucleoMag[®] Kit and the KingFisher[™] Apex

Objective: This protocol uses Nanotrap Microbiome A Particles and Nanotrap Enhancement Reagent 1 to capture and concentrate microbes in environmental water samples. It is optimized for microbe capture from 10 mL samples and is compatible with MACHEREY-NAGEL NucleoMag DNA/RNA Water Kit. The automated script can process up to 24 samples at once and can be amended for the throughput in your lab.

Materials and equipment:

Sample Type Environmental Water Samples	
Nanotrap Microbiome A Particles	Ceres Nanosciences; SKU# 44202
Nanotrap Enhancement Reagent 1 (ER1) ¹	Ceres Nanosciences; SKU# 10111
Extraction Kit	Vendor
NucleoMag DNA/RNA Water Extraction Kit	MACHEREY-NAGEL; REF 744220.1
Materials/Equipment	Vendor
KingFisher™ Apex with 96 DW Head	Thermo Fisher Scientific; Cat# 5400930
KingFisher Apex 24 Combi head	Thermo Fisher Scientific; Cat# 24079940
KF Apex 96 KF heating block	Thermo Fisher Scientific; Cat# 24075920
KF Apex 24 DW heating block	Thermo Fisher Scientific; Cat# 24075940
KingFisher 24 Deep-well Plate, Barcoded	Thermo Fisher Scientific; Cat#95040470B
KingFisher 24 Deep-Well Tip Comb & Plate, Barcoded	Thermo Fisher Scientific; Cat#97002610B
KingFisher 96 Deep-well Plate, Barcoded	Thermo Fisher Scientific; Cat# 95040450B
KingFisher 96 Plate (200 μL), Barcoded	Thermo Fisher Scientific; Cat# 97002540B
General Reagents	Vendor
Molecular grade water	VWR; 45001-044

¹ Precipitate can form in ER1 if stored below room temperature. Allow ER1 to return to room temperature to dissolve the precipitate (can invert 2-3 times to help resuspend, do not heat).

Capture and Extract Microbes using Nanotrap Microbiome Particles

Procedure:

1. Nanotrap Microbiome A NucleoMag KingFisher Apex Procedure-Part 1

- 1. Prepare "Sample Plate 1" and "Sample Plate 2"
 - 1. Invert environmental water sample 5 times to mix. After inverting, place on a flat surface for 45 seconds.
 - 2. Add 4,875 μL of environmental water sample to one well (one well per sample) of a new KingFisher24 Well Deep Well Plate.
 - 3. Add another 4,875 µL of environmental water sample to the same well location on a second KingFisher 24 Well Deep Well Plate.
 - a) For example, if you loaded a sample into well A1 of the first plate, load the second volume of that sample into well A1 of the second plate.
 - 4. Add 50 μ L of Nanotrap Enhancement Reagent 1 (ER1) Solution to each sample on the two KingFisher 24 Well Deep Well sample plates (100 μ L total).
 - 5. Add 75 μL of Nanotrap Microbiome A Particles to each sample on the two KingFisher™ 24 Well Deep Well sample plates (150 μL total).
- 2. Prepare "Lysis Plate"
 - 1. Add 500 μ L of Lysis Buffer MWA1 to a new (the third) KingFisher 24 Well Deep Well Plate matching the number and location of the "Sample Plate" wells.
- 3. Prepare "Tip Plate"
 - 1. Insert a new tip comb into a new KingFisher 24 Well Deep Well Plate.
- 4. Run NT Script (Request file at sales @ceresnano.com)
 - 1. Run NT Microbiome A NucleoMag® 24 w heat.kfx
 - 2. Follow the on-screen instructions loading the previously prepared plates at the appropriate time.
- 5. Once the protocol is completed, the "Lysis Plate" will contain lysate that is ready to proceed to Part 2 (*caution* sample may be hot).

2. Nanotrap Microbiome A NucleoMag KingFisher Apex Procedure-Part 2

- 1. Prepare "NM Binding" Plate
 - 1. To a new KingFisher 96 Deep Well Plate, add 450 µL of the cleared lysate (NT lysate) from each well of the lysis plate used in "Part 1 step 5" of the protocol. Keep track of which well contains which sample in this new bead binding plate.
 - 2. Add 475 μL of Binding Buffer MWA2 to each well in which lysate was added.

- 3. Vortex the NucleoMag B-beads thoroughly and add 25 µL to each well.
 - a) Note: Binding mix (MWA2 + B-beads) can be pre-mixed before their addition to the plate.
- 2. Prepare "1st Wash MWA3" Plate
 - 1. Add 850 µL of Wash Buffer MWA3 to a new KingFisher 96 Deep Well Plate matching the number and location of the KingFisher 96 Deep Well Plate- "NM Binding" Plate wells.
- 3. Prepare "2nd Wash MWA3" Plate
 - 1. Add 850 µL of Wash Buffer MWA3 to a new KingFisher 96 Deep Well Plate matching the number and location of the KingFisher 96 Deep Well Plate- "NM Binding" Plate wells.
- 4. Prepare "3rd Wash MWA4" Plate
 - 1. Add 850 µL of Wash Buffer MWA4 to a new KingFisher 96 Deep Well Plate matching the number and location of the KingFisher 96 Deep Well Plate- "NM Binding" Plate wells.
- 5. Prepare "Elution" Plate
 - 1. Add 100 μ L of Rnase-free water to a new KingFisher 96- 200 μ L plate matching the number and location of the KingFisher 96 Deep Well Plate-"NM Binding" Plate wells.
- 6. Prepare "Tip Plate"
 - Insert the KingFisher™ 96 Deep Well Comb into a new KingFisher 96 Deep Well Plate
- 7. Run Extraction Script (Request file at sales @ceresnano.com)
 - Run

NucleoMag_DNA_RNA_Water_CeresNanoTrap_Apex_Rev02.kfx

- 2. Follow the on-screen instructions loading the previously prepared plates at the appropriate time.
- 3. Once the protocol is completed, the KingFisher 96-Elution Plate contains eluates that are ready for downstream analysis or can be stored at -80°C.

Note: Multiple freeze-thaw cycles may cause degradation.

Attachments: 2

KingFisher™ Apex

- 1. NT_Microbiome_A_NucleoMag®_24_w_heat.kfx
- 2. NucleoMag_DNA_RNA_Water_CeresNanoTrap_Apex_Rev02.kfx