

Automated Nanotrap[®] Wastewater Protocol using IDEXX Water DNA/RNA Magnetic Bead Kit

Objective: This protocol uses Nanotrap[®] Magnetic Virus Particles and Nanotrap[®] Enhancement Reagent 2 to capture and concentrate viruses in wastewater samples. It is optimized for viral capture from 10 mL samples of wastewater and is compatible with IDEXX Water DNA/RNA Magnetic Bead Kit and RT-PCR Kit. This method has been verified with SARS-CoV-2 viral samples.

Materials and equipment:

1. Wastewater sample
2. Nanotrap[®] Enhancement Reagent 2 (ER2) (Ceres Nanosciences SKU 10112)
 - a. Using ER2 improves viral detection by 1-2 Ct values when used with Nanotrap[®] Magnetic Virus Particles, but it is ok to skip addition of ER2 in this protocol if it is not available.
3. Nanotrap[®] Magnetic Virus Particles (Ceres Nanosciences SKU 44202)
4. Thermo Scientific KingFisher[™] Apex
5. IDEXX Water Nucleic Acid Extraction Kit (IDEXX Cat# 98-0014719-00)
 - a. Binding Solution
 - b. Beads
 - c. Wash Buffer 1
 - d. Wash Buffer 2
 - e. Elution Buffer
6. Buffer AVL (Qiagen Cat# 19073)
7. 1x Phosphate Buffered Saline, without Calcium and Magnesium (PBS) (Lonza cat# 17-516F)
8. 3-24 well-KF Deep Well Plates
9. 1-24 well-KF Deep Well Comb
10. 4-96 well-KF Deep Well Plates
11. 1-96 well-KF (200 μ L) Plate
12. 1-96 well-KF Deep Well Comb

Procedure:

Capture and Concentrate Virus using Nanotrap[®] Particles

1. Ceres Nanotrap KF Procedure - Part 1

a. Prepare Sample Plates

- i. Add 4,875 μ L of wastewater sample from wastewater bottle to one well (one well per sample) of a new KingFisher[™] 24 Well Deep Well Plate.
- i. Add another 4,875 μ L of each sample to the same well on a second KingFisher[™] 24 Well Deep Well Plate. 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.
- ii. Add 50 μ L of Nanotrap[®] Enhancement Reagent 2 (ER2) Solution to each wastewater sample on the two KingFisher[™] 24 Well Deep Well sample plates.
 1. Using ER2 improves viral detection by 1-2 Ct values when used with Nanotrap[®] Magnetic Virus Particles, but it is ok to skip addition of ER2 and proceed to the next step.
- iii. Add 75 μ L of Nanotrap[®] Magnetic Virus Particles to each wastewater sample on the two KingFisher[™] 24 Well Deep Well sample plates.

- iv. Insert the comb into one of the sample plates. This will be “Sample Plate 1” while the other plate will be “Sample Plate 2”.
- b. *Prepare Lysis Plate*
 - i. Add 160 μL of AVL lysis buffer and 40 μL of PBS to the third Kingfisher™ 24 Well Deep Well Plate matching the number and location of the “Sample Plate” wells.
- c. *Run NT KingFisher™ Protocol (See attached file)*
 - i. Follow the on-screen instructions loading the previously prepared plates at the appropriate time.
- d. Once the protocol is completed, the KingFisher™ plate to which a lysis solution was added will contain lysate that is ready to be run on the Kingfisher IDEXX extraction protocol.

2. IDEXX KF Extraction Procedure -Part 2

- a. *Prepare Wash Plate 1*
 - i. Add 500 μL of Wash Buffer 1 to a new KingFisher™ 96 Deep Well Plate matching the number and location of the KingFisher™ 96 Deep Well Plate- Bead Binding Plate wells.
 - b. *Prepare Wash Plate 2*
 - i. Add 500 μL of Wash Buffer 2 to a new KingFisher™ 96 Deep Well Plate matching the number and location of the KingFisher™ 96 Deep Well Plate- Bead Binding Plate wells.
 - c. *Prepare Wash Plate 3*
 - i. Add 500 μL of Wash Buffer 2 to a new KingFisher™ 96 Deep Well Plate matching the number and location of the KingFisher™ 96 Deep Well Plate- Bead Binding Plate wells.
 - d. *Prepare Elution Plate*
 - i. Add 50 μL of Elution Buffer to a new KingFisher™ 96- Plate (200 μL) matching the number and location of the KingFisher™ 96 Deep Well Plate- Bead Binding Plate wells.
 - e. *Prepare IDEXX Bead Binding Plate*
 - i. To a new KingFisher™ 96 Deep Well Plate, add 200 μL of the lysate from each well of the lysis plate used in Part 1 of the protocol. Keep track of which well contains which sample in this new Bead Binding Plate.
 - ii. Add 500 μL of Binding Buffer to each well in which lysate was added.
 - iii. Vortex the IDEXX Beads thoroughly and add 20 μL to each well.
 - iv. Insert the KingFisher™ 96 Deep Well Comb into the bead binding plate.
 - f. *Run IDEXX KingFisher™ Protocol (See attached file)*
 - i. 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 will contain purified viral RNA that is ready to be loaded onto a PCR plate.

Attachments: 2

KingFisher™ Apex

1. *KF-006-WW-Nanotrap-24.kfx*
2. *KF-006-WW-IDEXX-96.kfx*