# HELINI MagPure Viral Nucleic acid Purification Kit

[VTM/Serum/Plasma]

Cat. No: 2501 – 96 Prep

Compatible with: Manual magnetic stand separator & Reputed open type Automatic DNA/RNA purification system

### Introduction

The HELINI MagPure Viral Nucleic acid purification Kit is designed for rapid manual and automated purification of Viral Nucleic acid from human biological samples including nasal swabs, buccal swabs and urogenital swabs and other clinical samples. The Nucleic acid purified using the HELINI MagPure Nucleic acid purification kit contains high quality RNA and free of proteins, nucleases, and other contaminants or inhibitors. They are, therefore, suitable for direct use in many different downstream applications, such as qPCR (quantitative PCR), RT-qPCR (reverse transcription qPCR), and several other enzymatic reactions.

### **Intended Use**

For the purification of Viral Nucleic acid/total nucleic acid from human clinical samples such as Plasma/Serum and VTM.

# **Principle and Procedure**

The HELINI MagPure Viral Nucleic acid purification Kit uses magnetic-particle technology for Nucleic acid purification. The HELINI Biomolecules MagPure technology combines the speed and efficiency of nucleic acids purification with easy handling of magnetic particles. The purification process requires no phenol/chloroform extraction and needs very little hands-on time. The HELINI MagPure Magnetic Beads are highly reactive, super paramagnetic beads. The first step of the protocol lyses the sample, after which the nucleic acids can bind to the surface of the Magnetic Beads. The following three effective wash steps dispose of proteins, cell debris, and any residual contaminants, while the nucleic acids bound to the MagPure Magnetic Beads are transferred through the wash steps. High-quality nucleic acids are eluted into the nuclease-free water, and are ready for subsequent downstream processes.

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# Kit components

Components per plate	Qty	Storage
Magnetic beads	1.5ml	4°C
Proteinase K	2ml	-20C
Lysis buffer	20ml	RT
Wash Buffer	120ml	RT
Elution Buffer	10ml	RT
Instruction manual		

# **Safety Information**

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. Discard sample and assay waste according to your local safety regulations.

# **Technical Assistance**

For technical assistance and more information, please contact; 0091-9382810333 0091-44-24490433 helinibiomolecules@gmail.com

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# Material required:

### For Manual:

- 1. Micro pipettes variable all standard ranges and micro tips
- 2. Magnetic stand separator
- 3. Micro centrifuge -1.5/2ml rotors
- 4. 1.5/2.0 micro centrifuge tubes

### For automation:

- 5. Reputed any make Open system Automated DNA/RNA purification system
- 6. Plastic wares & combes or tips compatible with automated purification system
- 7. Reagent reservoir 5 Nos. Label them and use dedicated for that particular reagent only.
- 8. 8 channel variable micro pipette range 30 to 300μl
- 9. 8 channel variable micro pipette range 5 to  $50\mu l$
- 10.8 channel variable micro pipette range 100 to  $1000\mu l$  [optional]
- 11. Tips for micro pipettes all above range.
- 12. 1.5/2.0 micro centrifuge tubes
- 13. Plate sealing film [optional]

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### **Procedure - Manual method**

- 1. Transfer 250μl of Lysis buffer into sterile 1.5ml centrifuge tube.
- 2. If you are using internal control template to monitor extraction efficiency, please pipette mix manufacturer indicated volume of internal control template.
- 3. Add 10μl of Magnetic beads. [Invert mix or vortex mix well the beads tube before pipetting]
- 4. Add 200μl of VTM / 150μl of Plasma/Serum. Mix well by pipetting or pulse vortex for 10 seconds.
- 5. Add 20µl of Proteinase K and Mix well by pipetting or pulse vortex for 10seconds. Brief spin to bring down the liquid attached in the micro tube cap.
- 6. Incubate at 56C for 10min. [Intermediate brief vortex for two to three times is recommended].
- 7. Place it on the Magnetic stand separator for 10seconds. Discard the supernatant. [Decant or Pipette it out]
- 8. Takeout the tube from Magnetic stand separator and add 600μl of Wash buffer and vortex well for 10secs. Centrifuge the tubes for 5secs and incubate at room temperature for 10seconds.

- 9. Place on the Magnetic stand separator for 10seconds. Discard the supernatant completely. [Decant or pipette it out]
- 10. Takeout from the Magnetic stand and add 600μl of Wash buffer and vortex well for 10secs. Centrifuge the tubes for 5secs and incubate at room temperature for 10seconds.
- 11. Place on the Magnetic stand separator for 10seconds. Discard the supernatant completely. [Make sure there is no residual liquid in the tube, if any, use a micropipette to remove them]
- 12. Add 100μl of Elution buffer and vortex well for 10secs. Centrifuge the tubes 5 seconds and incubate at 56°C for 3mins.
- 13. Place on the Magnetic stand separator for 1min and carefully transfer elute in to fresh 1.5ml centrifuge tube.
- 14. Use the Viral Nucleic acid immediately for qPCR/PCR or store at -20C/-80C for later use.

### **Recommendation:**

Use  $10\mu l$  for qPCR /PCR assay

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# **Procedure**

# For Automation

- 1. Please refer the machine manufacturer recommended plastic wears and tips/combs for filling the reagents.
- 2. Use the following reagents volume, steps timings, mixing speed and heating steps;

Buffers	Volume Per well	Mixing Time	Heating Temp	Mixing Speed	Magnetic Beads pickup Time	Magnetic Beads Pickup speed
Lysis buffer	250μ1	***	65°C	Moderate	10secs	Moderate
Wash buffer	600μ1	2min		Moderate	10secs	Moderate
Wash buffer	600μ1	1min		Moderate	10secs	Moderate
Elution buffer	100μ1	3min	60°C	Moderate	10secs	Moderate

<sup>\*\*\*5</sup>min for VTM

# Manufactured and Marketed by

# HELINI Biomolecules,

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<sup>\*\*\*10</sup>min for Plasma/serum

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