

MEBEP TECH(HK) Co., Limited

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Virus Genomic DNA/RNA Fast Kit II

Product Number: RNK2201

Shipping and Storage

Storage at room temperature for 12 months does not affect the effectiveness of use, and each solution should be covered tightly in a timely manner after use. Poly Carrier can be transported at room temperature and stored at 4 °C for one month. It can be stored for a long time at -20 °C.

Components

Component	Storage	RNK2201
		50 Preps
Buffer VLB	RT	20ml
Poly Carrier	-20°C	200µl
Buffer RE	RT	25 ml
Buffer RW	RT	10ml
RNase-free H ₂ O	RT	10ml
RNase free adsorption column RA and collection tube	RT	50set

Description

The virus DNA/RNA extraction kit is suitable for rapidly extracting high-purity virus DNA/RNA from cell-free body fluids, including plasma, serum, ascites, cultured cell supernatant, cerebrospinal fluid, and urine, using a centrifugation adsorption column that specifically binds to virus DNA/RNA and a unique buffer system. This product can meet the simultaneous extraction requirements of most viral RNAs/DNA, such as viral RNAs: HCV (hepatitis C virus), HIV (HIV), and HTLV (human T-lymphotropic virus); Virus DNA: HBV (hepatitis B virus) and CMV (cytomegalovirus), etc. After virus lysis, DNA/RNA selectively adsorbs onto the silica matrix membrane in a highly dissociated salt state (especially equipped with Poly Carrier, which can easily capture trace amounts of nucleic acids from the system). Then, impurities such as salt, cellular metabolites, and proteins are removed through a series of rapid rinsing centrifugation steps. Finally, the pure virus DNA/RNA is eluted from the silica matrix membrane using a low salt elution buffer. The purified viral nucleic acid is free of impurities and PCR inhibitors, and can be directly used for PCR/RT-PCR analysis.

Features

- 1. No toxic reagents such as phenol are required, and no steps such as ethanol precipitation are required.
- 2. Time saving, simple, and single sample operation can generally be completed within 20 minutes.
- 3. Multiple column washes ensure high purity, and the extracted virus DNA/RNA has high purity, stable and reliable quality. It can be used for various routine operations, including PCR/RT-PCR, enzyme digestion, sequencing, Southern hybridization, etc.

Note

- 1. All centrifugation steps are completed at room temperature using a traditional desktop centrifuge with a speed of up to 13000rpm.
- 2. Preheat the required water bath to a specific temperature before starting the experiment.
- 3. Buffer VLB and Buffer RE contain irritating compounds. Wear latex gloves during operation to avoid contact with skin, eyes, and clothing. If it gets on the skin or eyes, rinse with plenty of water or physiological saline.

4. Poly Carrier:

Usage of Poly Carrier: If the initial processing volume is small, we recommend using Poly Carrier. If a large amount of nucleic acid production is expected, users can choose whether to join Poly Carrier according to their needs. When using, add 4µl Poly

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Carrier storage solution to the Buffer VLB required for each sample extraction, and mix the Buffer VLB and Poly Carrier solution completely upside down (Buffer VLB is easy to form foam, do not use vortex oscillation to mix). You can also add the required Poly Carrier to the total required Buffer VLB according to the number of samples and mix well for later use. The mixture is stable at room temperature for 24 hours.

Protocol(Please read the precautions before the experiment)

Please add the specified amount of anhydrous ethanol to the Buffer RW bottle before first use!

 200µl of serum and other body fluids (need to be returned to room temperature, insufficient can be supplemented with 0.9% NaCl or PBS) are transferred into the aforementioned 1.5ml centrifuge tube, and 400µl of Buffer VLB is added. Immediately, vortex and shake thoroughly to mix well.

If the sample size is small or the expected concentration of the virus is low, it is recommended to add 4µl Poly Carrier storage solution to 400µl Buffer VLB.

- 2. Place at room temperature (15-25°C) for 10 minutes, shake and mix well every 5 minutes.
- 3. Add 450µl of anhydrous ethanol and immediately vortex and shake thoroughly to mix well.
 - If the surrounding environment is above 25°C, ethanol needs to be pre cooled on ice before being added.
- 4. Add the above mixture to an adsorption column RA, centrifuge at 13000rpm for 30-60 seconds (with the adsorption column placed in the collection tube), and discard the waste liquid in the collection tube.

If the total volume exceeds 750µl, the solution can be added twice to the same adsorption column RA.

- 5. Add 500µl buffer RE, centrifuge at 12000rpm for 30 seconds, and discard the waste liquid.
- Add 500µl of Buffer RW (please check if anhydrous ethanol has been added first!), centrifuge at 12000rpm for 30 seconds, discard the waste liquid, add 500µl of Buffer RW, and repeat.
- 7. Put the adsorption column RA back into the empty collection tube, centrifuge at 13000rpm for 2 minutes, and try to remove Buffer RW as much as possible to avoid residual ethanol in Buffer RW inhibiting downstream reactions.
- 8. Take out the adsorption column RA and place it in an RNase free centrifuge tube. Add 30-50µl of RNase free H₂O in the middle of the adsorption membrane (better heating effect in a 65-70°C water bath beforehand). Leave at room temperature for 1 minute and centrifuge at 12000 rpm for 1 minute. If you want to obtain a large amount of DNA/RNA, you can add the obtained solution back to the centrifuge adsorption column and centrifuge at 12000 rpm for 1 minute.

The larger the elution volume, the higher the elution efficiency. If a high concentration of DNA/RNA is required, the elution volume can be appropriately reduced, but the minimum volume should not be less than 20µl. If the volume is too small, it will reduce the elution efficiency and DNA/RNA production.

9. Virus DNA can be stored at 2-8°C, and if it needs to be stored for a long time, it can be stored at -20°C. It is recommended to use viral RNA immediately, otherwise it should be stored at -70°C for short-term use.