

1. Identification**Product name****HigherPurity™ Viral RNA Extraction Kit****Cat. No****AN0805 (100 reactions)****Cat. No****AN0805-XL (500 reactions)****2. Description**

HigherPurity™ Viral RNA Extraction Kit is designed for the rapid simultaneous purification of viral RNA from cell –free samples such as serum, plasma, urine, cell free body fluids, cell culture supernatants and rinse liquid from swabs samples.

The viral RNA molecules bind to the silica-based media and impurities such as proteins and nucleases are removed by thorough washing with Wash Buffer. The RNA is then eluted in sterile, RNase free water. The isolated viral RNA is ready to use and should be stored at - 70°C.

The procedure can be used for isolation of viral RNA from a broad range of viruses. However, performance cannot be guaranteed for every virus species and must be validated by the customer. The amount of purified viral RNA depends on the sample type, the virus titer, sample source, transport, storage, and age. The Kit also includes carrier RNA that improves binding and recovery of low-concentrated viral RNA.

3. Composition

Item	Quantity	
	AN0805	AN0805-XL
Minispin-columns	100	500
Collection tubes (2 ml)	200	1000
Carrier RNA (Lyophilized)*	8 x 0.2 mg	8 mg
BLY-Buffer	40 ml	200 ml
WB1 Buffer**	33 ml	165 ml
WB2 Buffer**	20 ml	100 ml
RNase-free Water	10 ml	50 ml

* Add appropriate volume RNase-free Water (included with the kit) to lyophilized Carrier RNA to obtain 4 µg/µL Carrier RNA stock solution. Prepare aliquots of 100 µl and store -20°C. For 25 purifications, thaw one vial of 100 µl Carrier RNA and mix thoroughly with 10 ml BLY-Buffer. Mark the label of the bottle to indicate that Carrier RNA was added.

**Add the volume ethanol (96%-100%) specified [Not included] to WB1 and WB2 Buffer prior to initial use (see bottle label for volume). After ethanol has been added, mark the bottle to indicate that this step has been completed.

4. Storage

Store the Carrier RNA (Lyophilized) at -20 °C and all other components at room temperature (18 to 25 °C). If any kit reagent forms a precipitate, warm at 55–65 °C until the precipitate dissolves and allow cooling to room temperature before use.

5. Applications

The purified viral RNA is suitable for use in RT-PCR and qRT-PCR and can be used for:

- Viral load monitoring
- Viral detection
- Viral genotyping



DATA SHEET

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6. Quality Control

The quality of **HigherPurity™ Viral RNA Extraction Kit** is tested on a lot-to-lot basis by isolating viral RNA from a 200µl serum sample.

7. Further information

Product Use Limitations

This product is developed, designed, and sold exclusively only for research purposes use. The product was not tested for use in diagnostics or for drug development, nor is it suitable for administration to humans or animals.

Safety Information

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate material safety data sheets (MSDS). These are available online in convenient and compact PDF format at www.canvaxbio.com where you can find, view, and print the MSDS for each CANVAX kit.



DETAILED PROTOCOL

1. Transfer 200 µL of sample (plasma, serum, urine, body fluids or cell cultured supernatant) into a microcentrifuge tube (not provided).

• **Nasopharyngeal swab (NP) /oropharyngeal swab (OP):** If the swab is delivered in a transport media suitable for nucleic acid virus stabilization, transfer 200 µL directly into microcentrifuge tube. If you get a swab without transport media, place the swab with into microcentrifuge tube containing PBS and incubate for 15 minutes at room temperature. Afterwards shake the swab vigorously, squeeze it and remove the swab. Use a 200 µL aliquot of the liquid for viral RNA extraction.

2. Add 400 µL **BLY-Buffer (containing Carrier RNA)** and mix immediately.

• **Carrier RNA** enhances binding of viral RNA to the silica membrane and reduces the risk of viral RNA degradation.

• **Internal Extraction Control:** When performing RNA extraction, it is often advantageous to have an exogenous source of RNA template that is spiked into the lysis buffer (BLY-Buffer). This control RNA is then co-purified with the sample RNA and can be detected as a positive control for the extraction process. **DO NOT** add the internal control RNA directly to the biological sample as this will lead to degradation and a loss in signal strength.

• **Proteinase K treatment:** Although for isolation of viral RNA, Proteinase K treatment is usually not required, it is recommended for isolation from viscous samples (e.g., sputum samples). Add 25µL Proteinase K (20 mg/ml stock solution), to the lysis mixture and mix by vortexing vigorously for 20 seconds. Incubate for 5 min at 70°C. [Proteinase K not included in the kit].

3. Incubate the mix at room temperature for 10 minutes.

4. Place **Minispin-column** in a 2 ml Collection tube and transfer lysed sample from previous step. Centrifuge at 8000xg for 1 minute. Discard the flow-through.

• For successful nucleic acid purification it is important to obtain a homogeneous, clear, and nonviscous sample before loading onto the Minispin- Column.

5. Place the **Minispin-column** in a new Collection tube and add 500 µL of **WB1 Buffer**. Centrifuge at 8000xg for 1 minute. Discard the flow-through.

6. Place the **Minispin-column** in the same Collection tube and add 500 µL of **WB2 Buffer**. Centrifuge at 8000xg for 1 minute. Discard the flow-through.

7. Place the **Minispin-column** in the same Collection tube and add 500 µL of **WB2 Buffer**. Centrifuge at 8000xg for 1 minute. Discard the flow-through.

8. Centrifuge at full speed for an additional 3 min to dry the **Minispin-column**.

• This step will avoid the residual liquid to inhibit subsequent enzymatic reactions.

9. Place the **Minispin-column** into a new, labelled 1.5 microcentrifuge tube (not provided) and pipet 50 µL **RNase-free Water** directly into the membrane. Close the cap and incubate for 2 minutes at room temperature.

10. Centrifuge at full speed for 1 minute to elute. The eluate contains viral RNA. After extraction place the Elution Tube on ice. For long time storage place the nucleic acids at -80°C.

• Final eluates contain viral RNA and Carrier RNA; therefore, it is not possible to quantify the nucleic acids isolated with the kit by photometric or fluorometric methods.

