

sbeadex viral RNA purification kit

Protocol for manual extraction

*For Research Use Only.
Not for use in diagnostic procedures.*

General protocol

1. Preparation of samples

This protocol has been verified using swabs shaken in universal transport media (UTM) or sputum, both prepared following CDC guidelines.

2. Preparing the particle and buffer premix

The sbeadex™ particles suspension and Binding buffer SB can be added to the reaction(s) as a premix.

To prepare the premix for the sbeadex viral RNA purification protocol:

- a. Thoroughly mix the sbeadex particle suspension to fully resuspend the particles
- b. Add 20 µL sbeadex particle suspension to 160 µL Binding buffer SB.

If preparing premix for multiple reactions, multiply the volumes accordingly and allow sufficient overage for accurate pipetting.

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3. Viral RNA purification

1. Add the following to the reaction tube in the order listed below:
 - a. **Optional***: 20 µL Proteinase solution (PK) to the reaction tube/well
 - b. **Optional***: 1 µg carrier RNA
 - c. 100 µL of the liquid starting sample (see section 1)
 - d. 100 µL (1x) Lysis buffer SB.

***Optional:** LGC, Biosearch Technologies has generated scientific data with reference materials (AccuPlex™ Reference Material from LGC SeraCare Life Sciences) showing that the use of Proteinase K and carrier RNA was not required and has no influence on data quality.

2. Incubate at 55 °C for 10 minutes with constant shaking.
3. Allow the sample(s) to cool to room temperature.
4. Add 20 µL sbeadex particles suspension and 160 µL Binding buffer SB (these can be added as a 180 µL of premix – see section 2).
5. Mix thoroughly and incubate for 5 minutes at room temperature with, constant shaking.
6. Bring magnet into contact with the tube(s) for 2 minutes.
7. Remove the supernatant and discard.
8. Separate the magnet from the sample tube(s).
9. Add 400 µL Wash buffer BN1.
10. Incubate for 5 minutes at room temperature, with constant shaking.
11. Bring magnet into contact with the tube(s) for 2 minutes.
12. Remove the supernatant and discard.
13. Separate the magnet from the sample tube(s).
14. Repeat steps 9-13 with Wash buffer TN1.
15. Repeat steps 9-13 with Wash buffer TN2.
16. Add 100 µL Elution buffer AMP. Mix thoroughly.
17. Incubate for 10 minutes at 60 °C with periodic shaking.
18. Bring magnet into contact with the tube(s) for 3 minutes.
19. Transfer the eluate to a new tube by pipetting, avoiding the transfer of any sbeadex beads.

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4. Safety information

To access the SDS document for the components in this kit, please visit our [website](#). Work with infectious virus should be carried out according to the regulation of the country within which the kit is being used.

- Wear appropriate skin and eye protection throughout the preparation procedure.
- Lysis buffer SB, Binding buffer SB and Wash buffer TN1 contain high concentrations of detergent and salt.
- Binding buffer SB and Wash buffer TN1 contain up to 50% n-propanol, therefore keep away from naked flames.
- Ensure kit components are stored appropriately according to local safety guidance.
- In case of accidental contact, thoroughly rinse or flush the affected areas with water.
- Spillages can be removed using standard laboratory cleaning procedures.
- Safety data sheets are available for all kit components on request.






| Kit component | GHS symbol | Hazard phrases | Precaution phrases |
|------------------------------|--|-------------------------------|--|
| Lysis buffer SB |  Warning | H302/H315/H319/H400 | P101/P102/P103/P273/ P280/P305+P351+P338/ P301+P312/P332+P313/P501/ P301+P312 |
| Protease solution |  Danger | H334/H317 | P101/P102/P103/P261/ P304+P341/P501 |
| Binding buffer SB |  Danger | H226/H302/H315/H318/H336/H400 | P101/P102/P103/P210/ P241/P303+P361+P353/ P305+P351+P338/P310/P501 |
| sbeadex particles suspension | - | - | - |
| Wash buffer BN1 |  Danger | H226/H332/H315/H318/H336 | P101/P102/P103/P210/ P303+P361+P353/ P305+P351+P338/P310/ P405/P501 |
| Wash buffer TN1 |  Danger | H315/H318/H226/H336 | P101/P102/P103/P210/ P303+P361+P353/ P305+P351+P338/P310/P405/ P501 |
| Wash buffer TN2 | - | - | - |
| Elution buffer AMP | - | - | - |

Table 1. Safety information for sbeadex viral RNA purification kit components.

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5. Technical support

If you require additional information or technical assistance please feel free to email our Technical Support Team at: techsupport@lgcgroup.com.

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Accelerated science.**

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