

# sbeadex Pathogen Nucleic Acid Purification Kit

## Protocol for manual extraction

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

### 1. Preparation of samples

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

### 2. Preparing the particle and buffer premix

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

To prepare the premix for the sbeadex Pathogen Nucleic Acid Purification Kit 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. Nucleic acid purification

#### 3.1. Optional pre-lysis for bacterial samples

1. Add the following to the reaction tube/well in the order listed below:
  - a. Optional: 20  $\mu$ L Protease solution
  - b. Optional: 1  $\mu$ g carrier DNA/RNA
  - c. 100  $\mu$ L of the liquid starting sample
  - d. 100  $\mu$ L (1x) Lysis buffer SB
2. Incubate at 55 °C for 10 minutes with constant shaking.
3. Allow the sample(s) to cool to room temperature.
4. Proceed to the step-by-step protocol (section 3.2).

**NOTE:** Some bacterial species may require further treatment (i.e. heat inactivation at 90 °C and/or zirconium beads) to disrupt the cell wall.

#### 3.2. Step-by-step protocol for nucleic acid purification

1. Add 20  $\mu$ L sbeadex particle suspension and 160  $\mu$ L Binding buffer SB (these can be added as a 180  $\mu$ L of premix – see section 2).
2. Mix thoroughly and incubate for 5 minutes at room temperature with constant shaking.
3. Bring magnet into contact with the tube(s) for 2 minutes.
4. Remove the supernatant and discard.
5. Separate the magnet from the sample tube(s).
6. Add 400  $\mu$ L Wash buffer BN1.
7. Incubate for 5 minutes at room temperature with constant shaking.
8. Bring magnet into contact with the tube(s) for 2 minutes.
9. Remove the supernatant and discard.
10. Separate the magnet from the sample tube(s).
11. Repeat steps 9-13 with Wash buffer TN1.
12. Repeat steps 9-13 with Wash buffer TN2.
13. Add 100  $\mu$ L Elution buffer AMP. Mix thoroughly.
14. Incubate for 10 minutes at 60 °C with periodic shaking.
15. Bring magnet into contact with the tube(s) for 3 minutes.
16. 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 Pathogen Nucleic Acid 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](mailto:techsupport@lgcgroup.com).

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**Integrated tools.  
Accelerated science.**

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