Contents

| Introduction | 2 |
|---|---|
| Principle | 2 |
| Storage and Stability | 2 |
| Kit Contents | 3 |
| Before Starting | 3 |
| Mag-Bind [™] Viral DNA/RNA Protocol (200 µl) | 1 |
| Mag-Bind™ Viral DNA/RNA Protocol (400 μl) | 3 |
| Troubleshooting Guide | 3 |

Revised September 2008

Introduction

The E.Z.N.A. Mag-Bind[™]Viral DNA/RNA Kit is designed for rapid and reliable isolation of viral DNA and RNA from serum, plasma, and other cell-free body fluids. The Mag-Bind[™] Magnetic Beads technology provide high quality RNA and RNA, which is suitable for direct use in most downstream applications such as amplifications and enzymatic reactions. This system can be easily adapted with an automated system and the procedure can be scaled up or down.

Principle

If using the Mag-Bind[™] Viral DNA/RNA Kit for the first time, please read this booklet to become familiar with the procedure and its various modifications. Samples are lysed in a specially formulated buffer containing detergent and chaotropic salt. After adjust the buffer condition, nucleic acids (DNA/RNA) will form a complex with magnetic beads.. The beads/nucleic acids complex is separated from lysates using a magnet. Proteins and cellular debris are efficiently washed away by a washing step. Purified DNA and RNA are then eluted in nuclease-free water or low ionic strength buffer. DNA and RNA purified with this method can be directly used in most downstream applications without the need for further purification.

Storage and Stability

Mag-Bind[™] Viral DNA/RNA Kit are stable for at least 12 months from date of purchase. During shipment, or storage in cool ambient conditions, precipitates may form in the MVL Buffer. Mag-Bind[™] Particle R need to be stored at 4-8° C.

Kit Contents

| Product | M6245-00 | M6245-01 | M6245-02 |
|-----------------------|----------|-----------|------------|
| Purification | 5 | 50 | 200 |
| Mag-Bind™ Particles R | 100 µL | 2 mL | 8 mL |
| Buffer MVL* | 10 ml | 2 x 50 mL | 2 x 200 mL |
| Proteinase K | 3 mg | 60 mg | 240 mg |
| Buffer MBW | 6 ml | 100 ml | 400 ml |
| SPR Wash Buffer | 2.5 mL | 12 mL | 50 mL |
| DEPC-Water | 10 mL | 100 mL | 400 mL |
| Carrier RNA | 50 µg | 320 µg | 1600 µg |
| User Manual | 1 | 1 | 1 |

* MVL Contains guanidine salts. Do not use bleach as disinfectant.

Before Starting

| | SPR Wash Buffer must be diluted with absolute ethanol before use. | | | |
|-----------|---|-------------------------|--|--|
| IMPORTANT | M6245-00 | Add 10 mL 100% ethanol | | |
| | M6245-01 | Add 48 mL 100% ethanol | | |
| | M6245-02 | Add 200 mL 100% ethanol | | |
| | Dissolve Proteinase K as following and store at -20°C. | | | |
| | M6245-00 | Add 75 µl DEPC-Water | | |
| | M6245-01 | Add 750 ul DEPC-Water | | |
| | | | | |

Please take a few minutes to read this booklet thoroughly and become familiar with the protocol. Prepare all materials required before starting in order to minimize RNA degradation. *Wear gloves/protective goggles and take great care when working with chemicals*.

 Whenever working with RNA, always wear latex gloves to minimize RNase contamination. Use only clean RNase-free disposable plastic pipette tips when using the supplied reagents.

- During the procedure work carefully but quickly.
- Under cool ambient conditions, crystals may form in MBW Buffer This is normal and the bottle should be warmed to re-dissolve the salt.
- Prepare Carrier RNA: Dissolve Carrier RNA with DEPC-Water to the final concentration at 1mg/ml. Dissolve the dissolved Carrier RNA into convenient aliquot and store at -20°C. Add 3 µl Carrier RNA for each milliliter of MVL Buffer before use.

Material and Equipments supplied by User

Have the following reagents and supplies ready before starting procedure:

- Magnetic separation device for 1.5 OR 2.0 mL centrifuge tubes
- 1.5 mL or 2.0 Centrifuge Tubes
- Absolute ethanol (96-100%)
- Isopropanol
- Ethanol (96-100%)

Mag-Bind[™] Viral DNA/RNA Isolation Protocol (for 200ul serum or plasma)

The following protocol is designed for isolating viral DNA and RNA from 200 μ l cell-free body fluids such as plasma, or serum. The protocol can be easily scale up to process up to 400 μ l sample by using appropriate processing tube and suitable magnetic separation device.

- Add 200 ul sample (serum, plasma or other cell-free body fluids) into a 1.5 mL Centrifuge Tube followed by adding 10 ul Proteinase K solution and 10 ul Mag-Bind[™] Particle RV.
- 2. Add 200 µl MVL/Carrier RNA Buffer mix to the sample. Mix throughly by pipetting or vortexing . Incubate at room temperature for 10 minutes.

Note: see the instruction for preparing Carrier RNA on page 4.

- 3. Add 400ul isopropanol to each sample, mix throughly by pipetting or vortexing. Incubate 3-5minutes at room temperature.
- 4. Place the Centrifuge Tube on a magnetic separation device to magnetize the

magnetic particles. Leave the tube on the magnet until all the magnetic particles are pelleted. The capture time is depend on the sample type and magnetic stand used.

- 5. Carefully aspirate and discard the supernatant without disturbing the magnetic particles.
- 6. Remove the centrifuge tube containing the magnetic particles from the magnetic separation device. Add 300 μl of Buffer MBW and resuspend magnetic particles pellet by pipetting up and down 20 times or vortex the tube at maximum speed for 30 seconds.
- 7. Place the tube onto a magnetic separation device to magnetize the magnetic particles. Aspirate and discard the cleared supernatant after the lysate is cleared.
- Remove the tube containing the magnetic particles from the magnetic separation device. Add 400 µl of SPR Wash Buffer to each well and resuspend magnetic particles pellet by pipetting up and down or vortexing for 30 seconds.
- 9. Place the tube onto a magnetic separation device to magnetize the magnetic particles. Aspirate an discard the cleared supernatant after the lysate is cleared.
- 10. Add another 400 µl of SPR Wash Buffer and resuspend magnetic particles pellet by pipetting up and down for 20 times or vortexing at maximum speed for 30 seconds.
- 11. Place the tube onto a magnetic separation device to magnetize the Mag-Bind[™] particles.
- **12.** Aspirate and discard the cleared supernatant. Carefully remove any liquid drop from tube. Leave the tube on the magnet stand. Air dry the magnetic particles at room temperature for 7-10 minutes.
- 13. Add 20-50 µl DEPC Treated water and resuspend magnetic particles pellet by vortexing or pipetting up and down for 20 times. Incubate at room temperature for 3 minutes.
- 14. Place the tube onto a magnetic separation device to magnetize the Mag-Bind[™] particles.
- 15. Transfer the cleared supernatant contains purified DNA and RNA into a new RNasefree microtube .

Mag-Bind[™] Viral DNA/ RNA Isolation Protocol (for 400ul serum or plasma

The following protocol is designed for isolating viral DNA and RNA from 400 μ l cell-free body fluids such as plasma, or serum. This protocol should use 2.0 mL centrifuge tube and suitable magnetic separation device.

- 1. Add 400 ul sample (serum, plasma or other cell-free body fluids) into appropriate Processing Tube (not supplied) followed by adding 10 ul Proteinase K solution and 10 ul Mag-Bind[™] Particle RV.
- Add 400 µl MVL/Carrier RNA Buffer mix to the sample. Mix throughly by pipetting or vortexing. Incubate at room temperature for 10 minutes.

Note: see the instruction for preparing Carrier RNA on page 4.

- 3. Add 800ul isopropanol to each sample, mix throughly by pipetting or vortexing. Incubate 3-5minutes at room temperature.
- 4. Place the centrifuge Tube on a magnetic separation device to magnetize the magnetic particles. Leave the tube on the magnet until all the magnetic particles are pelleted. The capture time is depend on the sample type and magnetic stand used.
- 5. Carefully aspirate and discard the supernatant without disturbing the magnetic particles.
- Remove the Processing Tube containing the magnetic particles from the magnetic separation device. Add 400 µl of Buffer MBW and resuspend magnetic particles pellet by pipetting up and down 20 times or vortex the tube at maximum speed for 1 minute.
- 7. Place the tube onto a magnetic separation device to magnetize the magnetic particles. Aspirate and discard the cleared supernatant after the lysate is cleared.
- 8. Remove the tube containing the magnetic particles from the magnetic separation device. Add 500 µl of SPR Wash Buffer to each well and resuspend magnetic particles pellet by pipetting up and down or vortexing for 30 seconds.
- 9. Place the tube onto a magnetic separation device to magnetize the magnetic particles. Aspirate an discard the cleared supernatant after the lysate is cleared.

- 10. Add another 500 µl of SPR Wash Buffer and resuspend magnetic particles pellet by pipetting up and down for 20 times or vortexing at maximum speed for 30 seconds.
- 11. Place the tube onto a magnetic separation device to magnetize the Mag-Bind[™] particles.
- **12.** Aspirate and discard the cleared supernatant. Carefully remove any liquid drop from tube. Leave the tube on the magnet stand. Air dry the magnetic particles at room temperature for 7-10 minutes.
- Add 50-100 µl DEPC Treated water and resuspend magnetic particles pellet by vortexing or pipetting up and down for 20 times. Incubate at room temperature for 3 minutes.
- 14. Place the tube onto a magnetic separation device to magnetize the Mag-Bind[™] particles.
- 15. Transfer the cleared supernatant contains purified DNA and RNA into a new RNasefree microtube .

Trouble Shooting

| Problem | Likely Cause | Suggestions |
|--|---|---|
| Low RNA yields | Incomplete resuspension of magnetic particles | Resuspend the magnetic particles by vortexing before use. |
| | RNA degraded during sample storage | Make sure the sample is properly stored and make sure the samples are processed immediately after collection or removal from storage. |
| | SPR Buffer were not prepared correctly. | Prepare the SPR Buffer by adding ethanol according to instruction |
| | Loss of magnetic beads during operation | Be careful not to remove the magnetic beads during the operation |
| | Undissolved particles in the cell lysate cause congregation of magnetic beads | make sure the lysate is clear of particles before adding magnetic beads. Do not over dry the magnetic beads |
| No RNA eluted. | SPR and MBW Wash Buffer Concentrate not diluted with absolute ethanol and isopropanol. | Prepare MBW and SPR Wash Buffer Concentrate as instructed on the label. |
| Problem with downstream application | | 1. RNA in the sample is already degraded; do not freeze and thaw the sample more than once or store at room temperature for too long |
| | | 2. Quantify the purified DNA/RNA accurately and use sufficient elute. |
| Carryover of the magnetic beads in the elution | Carryover from the magnetic beads in the eluted RNA will not effect downstream applications. | To remove the carryover magnetic beads from eluted DNA/RNA, simply magnetize the magnetic beads and carefully transfer to a new tube or tube. |