



PCRopsis™ Pre-Plated Lysis Beads

(NOT FOR RESALE)

v.20230926

INTENDED USE (Research use only. Not for use in diagnostic procedures.)

PCRopsis™ Pre-Plated Lysis Beads are intended to facilitate lysis of difficult to lyse microorganisms in a convenient 96-well plate.

PRINCIPLES OF THE PROCEDURE

PCRopsis™ Lysis Beads are solid beads coated with a mixture of molecules that facilitate lysis of difficult to lyse microorganisms and ensure compatibility with PCRopsis™ reagents. These beads outperform comparable, uncoated beads. Each well of the PCRopsis™ Pre-Plated Lysis Beads contains ~0.25g of PCRopsis™ Lysis Beads and the plate is sealed with an autoclaved silicon, piercable cover and aluminum coating for ease of use.

WARNINGS & PRECAUTIONS

For Research Use Only.

- Observe approved biohazard precautions and aseptic techniques to prevent contamination of the product.
- Directions should be read and followed carefully.
- Do not re-pack.
- Do not ingest.

Storage: This product is ready for use and no further preparation is necessary. The product should be transported and stored in its original container at 20°C - 30°C until used. Do not overheat or keep open in humid environments. Do not freeze prior to use. Improper storage will result in a loss of efficacy. Do not use after expiration date, which is printed on the label.

Product Deterioration: PCRopsis™ Pre-Plated Lysis Beads should not be used if (1) there is evidence of damage or contamination to the product, (2) there is evidence of moisture in the original container, (3) the color of the beads has changed from white prior to use, (4) the expiration date has passed, or (5) there are other signs of deterioration.

PROCEDURES

Materials Provided: PCRopsis™ Pre-Plated Lysis Beads

Materials Required But Not Provided: PCRopsis™ Reagent RVD with RVD Enhancer (see product IFU), PCRopsis™ Support for urine samples only (see product IFU), nuclease-free water for nail samples, thermal cycler, controlled heat block, plate vortexer, thin walled tube (0.2 ~ 0.6 mL) or 96-well PCR plate, plate sealer, pipette tips and test sample

Specimen: liquid sample containing microorganisms (e.g., dairy, waste water, centrifuged urine, oral rinse, beverages, swab in compatible medium, toenails in compatible medium, etc.)

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Procedure Outlines Based on Sample Types:

In all cases, samples should be free of debris. Remove debris from samples by letting the debris settle for ~10 minutes with gravity, centrifuging the sample at a low speed, or filtering the sample and only using the debris-free sample for processing. Firmly tap the plate on the bench three times to ensure the lysis beads are at the bottom of the wells before using PCRopsis™ Pre-Plated Lysis Beads.

For urine-based samples:

PCRopsis™ Reagent Prep:

1. Thoroughly mix 2 µL of PCRopsis™ Support with 1 mL PCRopsis™ Reagent RVD with RVD Enhancer (called **RVD-Support solution** from here on)
 1. This mixture is stable for at least 24 hours at room temperature

After Reagent Prep:

1. Centrifuge 1.5~25 mL of urine at >1,400xg for 10 minutes in a 15 mL or 50 mL conical
 1. Higher volumes of centrifuged urine is expected to result in higher sensitivity
2. Remove supernatant and leave <250 µL of residual urine
3. Resuspend cell pellet using residual urine
4. Sterilize the aluminum, top surface of the Pre-Plated Lysis Beads with 70% ethanol or suitable alternatives
5. Add 200 µL of the resuspended sample to a well in the Pre-Plated Lysis Beads plate by piercing the aluminum and silicon plate sealers with the pipette tip
 1. Both the aluminum and silicon plate sealer are piercable and should not be removed
6. Vortex plate on high (~3000 RPM) for 10 minutes to lyse microorganisms
 1. Vortexing can be reduced to 5~7 minutes if complete lysis is observed under these conditions
7. Mix 20 µL PCRopsis™ RVD-Support solution with 20 µL of the sample pre-processed with PCRopsis™ Pre-Plated Lysis Beads in a thin-walled tube or plate (0.2 ~ 0.6 mL)
 1. **For optimal results, the reagent needs to be added first to the tube/well before the sample is added.**
 2. Ratio of sample to PCRopsis™ reagent will remain 1:1, but volume can be increased if needed (example: 30 µL : 30 µL and so forth)
8. Pipette up & down to ensure complete mixing and then cap tube or apply plate sealer to plate to prevent evaporation

9. Heat diluted sample at 95°C for 15~20 minutes in a thermal cycler or controlled heat block with a lid, and let cool at room temperature for ~10 seconds before continuing
 1. Heating for a longer period of time may improve the nucleic acid extraction efficiency
 2. Make sure the heating device has reached the desired temperature before applying sample.
 3. You may need to increase the heating time if increasing the volume of sample and reagent past 100 µl of each
 4. Sample heating can be performed using a controlled heating block or thermal cycler; however a device lid is highly recommended to minimize popping of tube caps or unpeeling of the plate sealer
10. Mix heated sample and use lysed / stabilized sample in your desired downstream applications

For toenail samples:

PCRopsis™ Reagent Prep:

None needed. Please use PCRopsis™ Reagent RVD with RVD Enhancer directly.

After Reagent Prep:

1. Cut toe nail samples into small pieces measuring less than 0.5 cm in length to maximized exposed surface area
2. Remove the aluminum and silicon plate sealer covering the Pre-Plated Lysis Beads
3. Place nail pieces into a well of the Pre-Plated Lysis Beads
4. Add ~200 µL of nuclease-free water to the well with nail pieces and Lysis Beads
5. Seal the plate with the aluminum and silicon plate sealer by pressing on the plate sealer and ensuring a complete seal
6. Vortex plate on high (~3000 RPM) for 10 minutes to lyse microorganisms
 1. Vortexing can be reduced to 5~7 minutes if complete lysis is observed under these conditions
7. Mix 20 µL PCRopsis™ Reagent RVD with RVD Enhancer with 20 µL of the sample pre-processed with PCRopsis™ Lysis Beads in a thin-walled tube or plate (0.2 ~ 0.6 mL)
 1. **For optimal results, the reagent needs to be added first to the tube/well before the sample is added.**
 2. Ratio of sample to PCRopsis™ reagent will remain 1:1, but volume can be increased if needed (example: 30 µl : 30 µl and so forth)

8. Pipette up & down to ensure complete mixing and then cap tube or apply plate sealer to plate to prevent evaporation
9. Heat diluted sample at 95°C for 15~20 minutes in a thermal cycler or controlled heat block with a lid, and let cool at room temperature for ~10 seconds before continuing
 1. Heating for a longer period of time may improve the nucleic acid extraction efficiency
 2. Make sure the heating device has reached the desired temperature before applying sample.
 3. You may need to increase the heating time if increasing the volume of sample and reagent past 100 µl of each
 4. Sample heating can be performed using a controlled heating block or thermal cycler; however a device lid is highly recommended to minimize popping of tube caps or unpeeling of the plate sealer
10. Mix heated sample and use lysed / stabilized sample in your desired downstream applications

For select liquid samples (beverages, water, swab samples in transport mediums, oral rinse, saliva samples, etc.):

PCRopsis™ Reagent Prep:

None needed. Please use PCRopsis™ Reagent RVD with RVD Enhancer directly.

After Reagent Prep:

OPTIONAL:

1. Centrifuge 1.5~25 mL of sample at >1,400xg for 10 minutes in a 15 mL or 50 mL conical to concentrate cells
 1. Higher volumes of centrifuged sample is expected to result in higher concentrations of nucleic acids for downstream applications
 2. Remove supernatant and leave <250 µL of residual sample
 3. Resuspend the sample in residual liquid
1. Sterilize the aluminum, top surface of the Pre-Plated Lysis Beads with 70% ethanol or suitable alternatives
 2. Add 200 µL of the resuspended sample to a well in the Pre-Plated Lysis Beads plate by piercing the aluminum and silicon plate sealers with the pipette tip
 1. Both the aluminum and silicon plate sealer are piercable and should not be removed
 3. Vortex plate on high (~3000 RPM) for 10 minutes to lyse microorganisms
 1. Vortexing can be reduced to 5~7 minutes if complete lysis is observed under these conditions



4. Mix 20 μ L PCRopsis™ Reagent RVD with RVD Enhancer with 20 μ L of the sample pre-processed with PCRopsis™ Lysis Beads in a thin-walled tube or plate (0.2 ~ 0.6 mL)
 1. **For optimal results, the reagent needs to be added first to the tube/well before the sample is added.**
 2. Ratio of sample to PCRopsis™ reagent will remain 1:1, but volume can be increased if needed (example: 30 μ L : 30 μ L and so forth)
5. Pipette up & down to ensure complete mixing and then cap tube or apply plate sealer to plate to prevent evaporation
6. Heat diluted sample at 95°C for 15~20 minutes in a thermal cycler or controlled heat block with a lid, and let cool at room temperature for ~10 seconds before continuing
 1. Heating for a longer period of time may improve the nucleic acid extraction efficiency
 2. Make sure the heating device has reached the desired temperature before applying sample.
 3. You may need to increase the heating time if increasing the volume of sample and reagent past 100 μ L of each
 4. Sample heating can be performed using a controlled heating block or thermal cycler; however a device lid is highly recommended to minimize popping of tube caps or unpeeling of the plate sealer
7. Mix heated sample and use lysed / stabilized sample in your desired downstream application

AVAILABILITY – NOT FOR RESALE

Cat. #	Description
7520096	PCRopsis™ Pre-Plated Lysis Beads (96-well, deep well plate)

MANUFACTURER

Entopsis, Inc., 7600 NW 69th Ave, Medley, FL 33166, USA info@entopsis.com

Glossary of Symbols Used

RUO

Research use only



Storage temperature

REF

Manufacturer's catalog number

STERILE | A

Sterile through aseptic techniques

LOT

Lot number



Manufacturer



Expiration date (year/month)