

## SUGGESTED PROTOCOL: Extraction-free Processing of Nail Samples

v. 20230206

## Sample:

- Nail clippings
  - o Nails should be cut into small pieces to maximize exposed surface area

## Materials:

- Entopsis Inc.:
  - PCRopsis™ Reagent 123 (see product IFU)
  - o PCRopsis™ Lysis Beads (see product IFU)
- Test sample
- 2 mL round-bottom tubes
- Thin-walled PCR tubes
- Heat block or Thermocycler

## Methods:

- 1. Cut nail samples into small pieces measuring less than 0.5 cm in length to maximized exposed surface area
- 2. Add roughly 0.25 grams of Lysis Beads to a 1.5  $\sim$  2 mL vial or well in a deep, 96-well plate
  - 1. One full PCR*opsis*™ Lysis Bead Scoop holds ~0.3 grams of beads
- 3. Place nail pieces into a vial or well with Lysis Beads
  - 1. A 2 mL, round-bottom tubes works best
- 4. Add ~200  $\mu$ L of nuclease-free water to the vial or well with nail pieces and Lysis Beads
- 5. Cap tube or place a plate sealer on the deep well plate
- 6. Vortex on high for 5~10 minutes to lyse microorganisms
- 7. Mix 20 µL PCRopsis™ Reagent 123 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 PCR*opsis*™ 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 slightly longer period (extra 2~5 minutes) of time does not negatively affect results
  - 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