TB-LAMP Workflow

1. Sample transfer and lysis



> Remove the cap to open the heating tube of the Loopamp[™] PURE DNA Extraction Kit.



> Use the Pipette-60 to collect slowly the most purulent portion of each sputum sample. Rub the end of the tip on the bottom of the cup to avoid and cut strings.



> Transfer 60 µl of the sputum.

2. Loopamp[™] PURE DNA extraction



> Remove the cap of the adsorbent tube but do not discard it.



> Screw the heating tube onto the adsorbent tube.



> Mix the lysed sample with the powder in the adsorbent tube by shaking thoroughly.



> Shake the tube until a milky solution is obtained.

3. Loop-mediated isothermal amplification



> Incubate the tube upside down for 2 min (use timer) at room temperature to reconstitute the reagents in the cap.

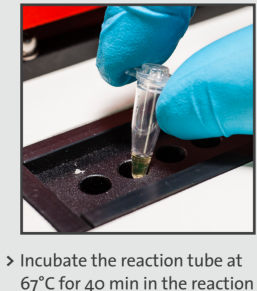
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> Mix the contents of the tube by inverting five times.



> Flick down the reaction tube until the reaction mixture is collected at the bottom.



unit. The reaction is automatically inactivated by a further incubation step at 80°C for 5 min.

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> Transfer the sample slowly into the heating tube. Slowly rinse the tip once to remove the sputum.



> Mix the contents of the tube by shaking.



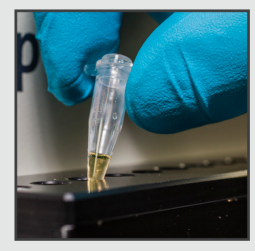
> Incubate the tube in the HumaLoop T heating unit at 90°C for 5 min.



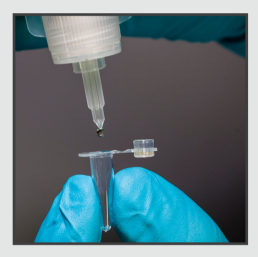


> Screw the injection cap onto the other side of the adsorbent tube.

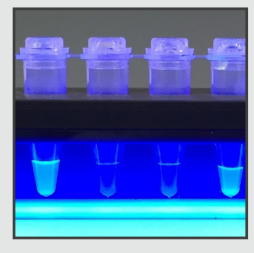
4. Result reading



> Insert the tube into the detection unit and turn on the UV light.



> Extract 30 µl of the DNA directly into the reaction tube by squeezing the adsorbent tube.



> Positive results fluoresce green.

A visualized workflow is shown. Please always refer to the latest versions of the Instructions for Use.



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