

## Protocol 1, part A: Fluid Sample Preparation

Kit 1: Buffers & Consumables (+18 to +25°C) Kit 3: Consumables (+18 to +25°C)

- Pipette 10ml of the fluid specimen in the 50ml Centrifuge tube (Kit 3).
- In case of less sample volume available, pipette the fluid into the *50ml Centrifuge tube* and fill up to 10ml with buffer *SU* (Kit 1) (use the measure line of the tube).

Continue with the instructions of the scheme Ultra-Deep Microbiome Prep10 / Protocol 1, part B: DNA Isolation (page 3, short manual).



\*<u>Please note:</u> Before using this scheme inform yourself of the details of the procedure. Please consult the manual. Special care is required for working under DNA-free conditions and secure working conditions, please consult the manual for more information.

**Ultra-Deep Microbiome Prep10 Short Manual** 

Molzym GmbH & Co.KG; Mary-Astell-Str. 10; 28359 Bremen; Germany; www.molzym.com; info@molzym.com



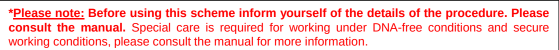
**Tissue** 

## Protocol 2, part A: Tissue Sample Preparation

Kit 1: Buffers & Consumables (+18 to +25°C) Kit 2: Enzymes & Reagents (-15 to -25°C)

- Transport the specimen under conditions avoiding contamination to the laboratory for analysis.
- Per specimen, place a *Sample tube* (*ST*, Kit 1,) in a rack and mark. Pipette 180µl of buffer *PKB* (Kit 1) into the *Sample tube* (*ST*).
- Transfer the specimen to a sterile support (e.g., Petri dish). Cut the specimen (~0.5x0.5cm) into small pieces by using a sterile scalpel.
- Transfer the cut specimen to the Sample tube (ST) filled with buffer PKB.
  Add 20µl of Proteinase K (Kit 2), vortex for 15s.
  Incubate at 56°C, 10min, 1,000rpm (thermomixer).
- Fill up to 1ml with the transport solution, if available, or with buffer *TSB* (use the measure line of the tube).

Continue with the instructions of the scheme Ultra-Deep Microbiome Prep10 / Protocol 2, part B: DNA Isolation (page 5, short manual).



**Ultra-Deep Microbiome Prep10 Short Manual** 



**Removal of Human DNA** 

Lysis of Pathogens

Kit 1: Buffers & Consumables (+18 to +25°C) Kit 2: Enzymes & Reagents (-15 to -25°C)

Arrange bottles according to the sequence of steps as below: *CM – DB1 – RS – RL – RP – CS – AB – WB – 70% Ethanol – Deionized Water* 

Continued from Ultra-Deep Microbiome Prep10 / Fluid Sample Preparation, part A (page 1, short manual).

Per sample:

- Add **4ml buffer** *CM*, vortex for 15s. Let stand at room temperature (+18 to +25°C) for 5min.
- Add 4ml buffer DB1.
  Add 10µl MolDNase B (Kit 2), vortex for 15s.
  Incubate at room temperature (+18 to +25°C) for 15min.
- 3. Centrifuge at 9,500xg, 10min. Remove supernatant by pipetting and discard.
- 4. Resuspend pellet in **1ml buffer RS** by pipetting.
- Transfer the suspension by pipetting into a Sample tube (ST, Kit 1) and centrifuge at ≥12,000xg, 5min. Remove supernatant by pipetting. (Optional: freeze pellet at -15 to -25°C for storage).
- 6. Resuspend pellet in **80µl buffer** *RL*, briefly centrifuge tube.
- Add 20µl BugLysis (Kit 2).
  Add 1.4µl ß-mercaptoethanol (Kit 2), vortex for 15s.
  Take care not to inhale.
  Incubate at 37°C, 30min, 1,000rpm (thermomixer).
- Briefly centrifuge. Add 150µl buffer *RP*. Add 20µl *Proteinase K* (Kit 2), vortex for 15s. Incubate at 56°C, 10 min, 1,000 rpm (thermomixer). Continue on page 4

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> atnogens Lysis

> > **DNA** Purification

DNA Elutior

During 10 min incubation: Kit 1: Buffers & Consumables

Unpack Spin columns (SC), 2 ml Collection tubes (CT) and 1.5 ml Elution tubes (ET), label; heat Deionized Water (100µl each sample) vial to 70°C (thermomixer).

- 9. Briefly centrifuge. Add 250µl buffer CS, vortex for 15s.
- 10. Briefly centrifuge. Add 250µl buffer AB, vortex for 15s.
- 11. Briefly centrifuge to clear lid. Pipette lysate into a Spin column. Centrifuge:  $\geq 12,000 \times g$ , 30 to 60s.
- 12. Remove column and place in a new 2 ml Collection tube. Add 400µl buffer WB. Centrifuge: ≥12,000x*q*, 30 to 60s.
- 13. Remove column and place in a new 2 ml Collection tube. Add 400µl 70% Ethanol. Centrifuge:  $\geq 12,000 \times g$ , 3min.
- 14. Carefully remove column and place in a 1.5 ml *Elution tube*.
- 15. Add 100µl Deionized Water heated to 70°C. Incubate at room temperature (+18 to +25°C) for 1min. Centrifuge:  $\geq$ 12,000xg, 1min. Discard column. close lid of *Elution tube*.
- 16. Store eluted DNA (1.5 ml Elution tube) at -15 to -25°C.

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Version 04



**Removal of Human DNA** 

Lysis of Pathogens

Kit 1: Buffers & Consumables (+18 to +25°C) Kit 2: Enzymes & Reagents (-15 to -25°C)

Arrange bottles according to the sequence of steps as below: *CM – DB1 – RS – RL – RP – CS – AB – WB – 70% Ethanol – Deionized Water* 

Continued from Ultra-Deep Microbiome Prep10 / Tissue Sample Preparation, part A (page 2, short manual).

Per sample:

- Add **250µl buffer** *CM*, vortex for 15s. Let stand at room temperature (+18 to +25°C) for 5min.
- Briefly centrifuge.
  Add 250µl buffer DB1.
  Add 10µl MolDNase B (Kit 2), vortex for 15s.
  Incubate at room temperature (+18 to +25°C) for 15min.
- Centrifuge at ≥12,000xg, 10min.
  Remove supernatant by pipetting and discard.
- 4. Resuspend pellet in **1ml buffer RS**.
- Centrifuge at ≥12,000xg, 5min.
  Remove supernatant by pipetting.
  (Optional: freeze pellet at -15 to -25°C for storage).
- Resuspend pellet in 80µl buffer *RL*, briefly centrifuge tube. Add 20µl *BugLysis* (Kit 2). Add 1.4µl *ß-mercaptoethanol* (Kit 2), vortex for 15s. *Take care not to inhale.* Incubate at 37°C, 30min, 1,000rpm (thermomixer).
- Briefly centrifuge.
  Add 150µl buffer RP.
  Add 20µl Proteinase K (Kit 2), vortex for 15s.
  Incubate at 56°C, 10 min, 1,000 rpm (thermomixer).

## Continue on page 6

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During 10 min incubation: Kit 1: Buffers & Consumables

Unpack Spin columns (SC), 2 ml Collection tubes (CT) and 1.5 ml Elution tubes (ET), label; heat Deionized Water (100µl each sample) vial to 70°C (thermomixer).

- 8. Briefly centrifuge. Add 250µl buffer CS, vortex for 15s.
- 9. Briefly centrifuge. Add 250µl buffer AB, vortex for 15s.
- 10. Briefly centrifuge to clear lid. Pipette lysate into a Spin column. Pipette the fluid phase in the column. Avoid transfer of any unresolved particles! Centrifuge: ≥12,000x*q*, 30 to 60s.
- 11. Remove column and place in a new 2 ml Collection tube. Add 400µl buffer WB. Centrifuge:  $\geq 12,000xq$ , 30 to 60s.
- 12. Remove column and place in a new 2 ml Collection tube. Add 400µl 70% Ethanol. Centrifuge:  $\geq 12,000xq$ , 3min.
- 13. Carefully remove column and place in a 1.5 ml *Elution tube*.
- 14. Add 100µl Deionized Water heated to 70°C. Incubate at room temperature (+18 to +25°C) for 1min. Centrifuge:  $\geq 12,000xq$ , 1min. Discard column, close lid of Elution tube.
- 15. Store eluted DNA (1.5 ml Elution tube) at -15 to -25°C.

\*<u>Please note:</u> Before using this scheme inform yourself of the details of the procedure. Please consult the manual. Special care is required for working under DNA-free conditions and secure working conditions, please consult the manual for more information.

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Version 04



**DNA Elutior** 





**DNA** Purification