

October 2023



DANAGENE MICROBIOME SOIL DNA KIT

Ref. 0621 50 PREPS

1. INTRODUCTION

The DANAGENE MICROBIOME Soil DNA kit has been designed for the isolation of high molecular weight genomic DNA from microorganisms like Gram positive and Gram negative bacteria, archaea and fungi **from environmental samples of all soil types.**

The kit uses a new **Microbiome Lysis Buffer and Microbiome Precipitation Buffer** specifically designed for use with environmental samples containing high humic substances content, including difficult soil types.

In this procedure, the microorganisms are efficiently lysed by a combination of heat, chemical and mechanical disruption with specialized beads. Proteins and Inhibitors are eliminated by precipitation using a new proprietary inhibitor removal buffer and subsequently pelleted by centrifugation together with the beads and undissolved sample material. The purified lysate is mixed with the Binding Buffer and then applied to a Microbial DNA column. The DNA that is bound to the column undergoes a two wash step. After a drying step, ready to use DNA can be eluted with Elution Buffer (5 mM Tris/HCl, pH 8.5).

Features:

- **Designed for a fast and easy purification microbial DNA from different types of soil samples.**
- **Optimized lysis method-combination of heat, chemical and mechanical lysis via bead-based homogenization enables isolation of DNA from archaea, fungi, Gram-negative and Gram-positive bacteria.**
- **Eliminates inhibitory substances as humic substances and others inhibitors.**
- **No phenol/chloroform extraction or ethanol precipitation is necessary.**
- **The eluted DNA is ready to use for all standard downstream applications. In most cases the concentrated DNA can be used as PCR template without further dilution for highest sensitivity**

Applications:

- **Microbiome analysis.**
- **PCR and Real-time applications.**
- **Microarrays.**
- **Mutation analysis.**

2. COMPONENTS KIT

	50 preps	Storage
Microbiome Lysis Buffer	60 ml	Room temperature
Microbiome Precipitation Buffer	12 ml	4°C
Binding Buffer Soil	45 ml	Room temperature
Desinhibition Buffer	18 ml	Room temperature
Wash Buffer	10 ml	Room temperature
Elution Buffer	5 ml	Room temperature
Bead Microtubrs	50 units	Room temperature
Proteinase K *	30 mg	-20°C
Microbial DNA Columns	50 units	Room temperature
Collection Tubes	150 units	Room temperature

(*) **This solution must be prepared as indicated in the Preliminary Preparations section of the protocol.**

PRECAUTIONS: The Lysis Buffer Soil and Binding Buffer contains guanidium salts, and should be handled with care. Guanidinium salts form highly reactive compounds when combined bleach.

Intended Use

All DANAGENE products are intended for molecular biology applications. These products are not intended for the diagnosis, prevention, or treatment of a disease.

2.1 Equipment and additional reagents required

Reagents

- 96 – 100 % ethanol

Consumables

- 1.5 mL microcentrifuge tubes
- Disposable pipette tips

Equipment

- Manual pipettors (50–1000 µl)
- Heat block, dry bath, or water bath (70°C)
- Centrifuge for microcentrifuge tubes
- Equipment for sample disruption and homogenization (see section 3.2). **Vortex-Genie® 2**
- Personal protection equipment (e.g., lab coat, gloves, goggles)

3. PROTOCOL

3.1 Preliminary Preparations

- Dissolve the proteinase K in **1.3 ml of nuclease-free water** and store at -20°C . It is recommended to do several aliquots to avoid many thaw/freeze cycles. At this temperature it is stable for 1 year.
- **Pre-heat the Elution Buffer at 70°C .**

3.2 General Remarks

Amount of starting material

DANAGENE MICROBIOME SOIL DNA kit is suitable for processing sample material of 250 mg. Very dry material can soak up large volumes of lysis. In this case, either reduce the amount of sample material or add additional lysis buffer.

The kit contains sufficient Microbiome Lysis Buffer for 50 samples using up to 1200 μl per sample. However, fill the Bead Tube (including the beads) to ensure sufficient head space for an efficient mechanical disruption.

Usually a reduction of starting material also helps to improve the lysis efficiency and to increase the purity of the DNA.

If possible remove foreign material like leaves, stones, or twigs (e.g., by sieving) as well as excess of water (e.g., by discarding the supernatant after spinning down sediment samples).

Sample Homogenization

The procedure is optimized by using "beads" in a vortex with horizontal agitation (Vortex Genie 2 or similar). Make sure that the vortex adapter allows horizontal agitation; **This is a cost-effective method for recovery of high-quality microbial DNA.**

Adapters with a vertical tube orientation may not agitate properly.

Processing time will vary based on sample input and bead beater. Times may be as little as 5 minutes when using high-speed cell disrupters (FastPrep) or as long 10-20 minutes when using lower speeds (Vortex-Genie® 2).

The DANAGENE MICROBIOME SOIL DNA kit does not require homogenization using a high-velocity bead beater. However, if the microorganism of interest requires stronger homogenization than provided by a vortex, or if using a bead beater is desired **you can use "Bead mil" homogenizers such as FastPrep, Precellys and others** but following the manufacturer's instructions to optimize the lysis of the sample. IMPORTANT: Many modern disruption devices can cause very high energy input in bead tubes. Depending on bead tube type and content (beads, liquid volume, sample type), especially high frequency of shaking and / or long shaking duration can cause breaking up of the bead tubes! **It is the responsibility of the user to perform initial stability test for the used bead tubes under the conditions used!** Perform initial test with water instead of lysis buffer and moderate machine setting (low frequency, short time) in order to avoid spillage of chaotropic lysis buffer in case of tube breakage.

3.3 Protocol for DNA extraction from 250 mg of environmental samples

1. Add up to **250 mg of soil sample** in a 2.0 ml **bead microtube** and add **850 µL of Microbiome Lysis Buffer**. Vortex briefly to mix.

Depending on the nature of the soil, it may be necessary to increase the Microbiome Lysis Buffer proportionally with the aim of recovering **500-600 µl of supernatant in point 4** (dry samples). The kit contains sufficient Microbiome Lysis Buffer for 50 samples using 1200 µl per sample.

For very wet material: Remove excess liquid before addition of lysis buffer, if necessary, after spinning down the sample.

2. **Add 25 µl of Proteinase K. Incubate at 70°C for 10 minutes.**

3. Homogenize samples thoroughly using one of the explained methods

Homogenize by bead beating for **10 minutes** at maximum speed on the [Vortex Genie 2](#) or [similar](#) using a **horizontal adapter**.

*The lysis time should be as short as necessary to avoid shearing of DNA and to minimize the release of humic acids. **Depending on the sample**, however, it might be advantageous to increase the lysis time to 10, 20, or 30 min.*

4. **Centrifuge at 16.500 x g for 2 minutes.** Transfer **the supernatant** to a clean 1.5 ml microcentrifuge tube.
IMPORTANT: Expect 500-600 µl of supernatant. A layer of debris may be present on top of the bead pellet. Avoid transfer of this debris with the supernatant.
5. Add **200 µl Microbiome Precipitation Buffer. Vortex.**
6. **Centrifuge at 16.500 x g for 2 minutes.** Transfer **600 µl of supernatant** in a new 1.5 ml microtube avoiding touching the pellet.
7. Add **900 µl of Binding Buffer Soil** and vortex briefly.
8. Load **750 µl** mixture sample into reservoir of a combined Microbial DNA column-collection tube assembly. **Centrifuge at 15.000 x g for 1 minute.**
9. Discard the flow-through and place the Spin Column back into the same 2 ml Collection Tube, repeat step 8 with the remaining sample mixture.

Ensure that the entire sample mixture has passed into the collection tube by inspecting the column. If sample remains in the column, centrifuge again at 15.000 x g for 1 minute.

10. Carefully place the Spin Column into a clean 2 ml Collection Tube (provided). Avoid splashing any flow-through onto the Spin Column.
11. Add **500 µl of Desinhibition Buffer. Centrifuge at 15.000 x g for 1 minute.** Remove the collection tube.
12. Add **700 µl of Wash Buffer. Centrifuge at 16.000 x g for 1 minute.** Remove the collection tube.
13. Carefully place the Spin Column into a clean 2 ml Collection Tube (provided). Avoid splashing any flow-through onto the Spin Column.
14. **Dry silica membrane.** Centrifuge at 16.500 x g for **2 minutes.**

15. Place the Microbial DNA Column into a 1.5 mL nuclease-free tube (not provided) and add **50-100 µL Pre-heat the Elution Buffer** at 70°C on the centre of the white filter membrane. Incubate **at room temperature** for **2 minutes**.
16. **Centrifuge** the spin column-tube assembly **at 15.000 x g for 1 minute**, then discard the column. The DNA is now ready for downstream applications.

4. PROBLEM GUIDE AND POSSIBLE ANSWER

Due to the great environmental samples variety that can be treated, it becomes difficult to generalize possible problems and answers. For this reason, we recommend to contact **DANAGEN-BIOTED** Laboratory Technical Service for any question regarding the protocols, specific soil samples or any problem you may have during the process.

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