

## Blood and Tissue DNA Mini Kit, 50 preps

**Product Number: DNK0701**

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### Shipping and Storage

1. Buffer TL and Buffer CB may precipitate and precipitate at low temperatures. They can be dissolved again in a 37°C water bath for a few minutes to restore clarity. After cooling to room temperature, they can be used.
2. Proteinase K is stored in a ready-to-use glycerol buffer and transported at room temperature. Upon receipt, it should be stored at room temperature for at least 6 months, at 4°C for 12 months, and at -20°C for 2 years
3. Avoid volatilization, oxidation, and pH changes caused by prolonged exposure of reagents to the air. Each solution should be covered tightly after use.

### Components

Component	Storage	DNK0701 50 Preps	DNK0702 100 Preps	DNK0703 200 Preps
Balance Buffer	RT	5 ml	10 ml	20 ml
Buffer TL	RT	11 ml	20 ml	40 ml
Buffer CB	RT	11 ml	20 ml	40 ml
Buffer PE	RT	16 ml	32 ml	64 ml
Buffer WB	RT	13 ml	25 ml	50 ml
Buffer EB	RT	15 ml	15 ml	20 ml
Proteinase K	4°C	1 ml	1 ml×2	1 ml×4
Adsorption column AC	RT	50	100	200
Collection tube (2ml)	RT	50	100	200

### Description

The unique Buffer CB/Proteinase K rapidly lyse cells and use silica gel membrane centrifugation columns to specifically adsorb DNA, without the need for toxic reagents such as phenol chloroform or time-consuming alcohol precipitation, maximizing the removal of proteins and other inhibitory impurities. Suitable for efficiently extracting genomic DNA from various materials such as animal tissue cells, mouse tails, insects, etc. The extracted DNA can be directly used for experiments such as enzyme digestion, PCR, Southern Blot, and virus detection.

### Features

1. There is no need to use toxic reagents such as phenol, nor do steps such as ethanol precipitation.
2. Fast and simple, single sample operation can generally be completed within 30 minutes.
3. Multiple column rinsing ensures high purity, with a typical OD<sub>260</sub>/OD<sub>280</sub> ratio of 1.7-1.9 and a length of up to 30kb-50kb. It can be directly used for PCR, Southern blot, and various enzyme digestion reactions.

### Application

Suitable for extracting genomic DNA from tissues/cells/mouse tails/insects.

### Note

1. All centrifugation steps are completed at room temperature using a traditional desktop centrifuge with a rotational speed of up to 13000rpm.
2. You need to bring your own isopropanol (for the steps of adding isopropanol, it is recommended to use isopropanol first, and ethanol can also be used as a substitute without isopropanol). RNase A (optional).

**For Research Use Only**

3. Before the experiment, preheat the required water bath to 70°C for later use.

**Protocol(Please read the precautions before the experiment)**

Tip: 1) Before the first use, please add anhydrous ethanol according to the instructions on the labels of Buffer WB and Buffer PE bottles, mix thoroughly, and mark the box with a hook indicating that ethanol has been added in a timely manner to avoid multiple additions!

1. **Column equilibrium:** Add 100µl Balance Buffer to the adsorption column AC, centrifuge at 13000 rpm for 1 minute, discard the filtrate, and set aside for later use.

**Note:** Balance Buffer can enhance the adsorption capacity of silica gel membrane for nucleic acids. Please use the adsorption column processed on the same day.

2. **Tissue cultured cells**

- 2.1. Collect approximately  $10^5$ - $10^6$  suspended cells into a 1.5ml centrifuge tube; For adherent cells, they should be digested with trypsin before being blown down and collected.
- 2.2. Centrifuge at 13000rpm for 10 seconds to allow cells to precipitate. Discard the supernatant, leaving behind cell clusters.
- 2.3. Add 200µl of 1×PBS (or physiological saline 0.9% sodium chloride solution) to resuspend and wash the cells, Centrifuge at 000 rpm for 10 seconds to allow the cells to precipitate. Completely discard the supernatant and resuspend the cell pellet in 180µl of 1×PBS.
- 2.4. Add 20µl of Proteinase K, mix thoroughly, and then add 200µl of Buffer CB. **Immediately shake and mix thoroughly**, and let it stand at 70°C for 10 minutes.

**Optional steps:** If there is a large amount of RNA residue that needs to be removed, 5µl of RNase A (100mg/ml) solution can be added before adding 200µl of Buffer CB, shaken well, and left at room temperature for 5-10 minutes.

- 2.5. After cooling, add 100µl of isopropanol (which can also be replaced by anhydrous ethanol, the same below), immediately vortex and shake thoroughly to mix well. At this time, flocculent precipitation may appear.
- 2.6. Add the previous mixture (including possible precipitates) to an adsorption column AC, centrifuge at 13000 rpm for 60 seconds (with the adsorption column placed in the collection tube), and discard the waste liquid from the collection tube.
- 2.7. **Continue to step 5 of the operation.**

3. **Animal and plant tissues (such as mouse liver and brain or plant leaves)**

- 3.1. Cut 20-50mg of fresh or thawed tissue into small pieces using a dissecting knife (cutting into small pieces can increase yield) or grind the tissue into fine powder in liquid nitrogen. Transfer the tissue into a 1.5ml centrifuge tube containing 180µl of buffer TL and mix well with a large caliber gun.
- 3.2. Join 20µl of Proteinase K, **immediately vortex shake and thoroughly mix.**
- 3.3. Place the lysate in a 55°C water bath for 1-3 hours or until the tissue is fully digested, gently shaking it several times during this period to aid in lysis.

**Optional steps:** If there is a large amount of RNA residue that needs to be removed, 5µl of RNase A (100mg/ml) solution can be added before adding 200µl of Buffer CB, shaken well, and left at room temperature for 5-10 minutes.

- 3.4. Add 200µl Buffer CB, **immediately shake thoroughly with vortex** and let stand at 70°C for 10 minutes.
- 3.5. After cooling, add 100µl of isopropanol and **immediately mix thoroughly through vortex** oscillation. At this point, flocculent precipitation may occur.
- 3.6. Suck the mixture with a 1ml gun tip, add the mixture to an adsorption column AC, centrifuge at 13000rpm for 60 seconds (the adsorption column is placed in the collection tube), and pour out the waste liquid in the collection tube.

**Note:** If there is any insoluble tissue that may block the gun head, the gun head can be gently rubbed onto absorbent paper to remove the insoluble matter; If there is a small amount of mixture sucked up, the gun head and insoluble matter can be discarded together. This method is to remove the insoluble matter and avoid clogging the centrifuge column.

- 3.7. **Continue to step 5 of the operation.**

4. **Animal tissue (rat tail)**

- 4.1. Cut the 0.2-0.5cm tip of the mouse tail (i.e. 20-50mg) into small pieces (**it is necessary to cut the tip within the range**

of 0-2cm, otherwise the cracking effect will not be good), or grind the tissue into fine powder in liquid nitrogen, transfer it into a 1.5ml centrifuge tube containing 180µl buffer TL, and use a large caliber gun to blow and mix.

- 4.2. Join 20µl of Proteinase K, **immediately vortex oscillate and thoroughly mix.**
- 4.3. Place the lysate in a 55°C water bath for 3 hours or until the tissue is fully digested, gently shaking it several times during this period to aid in lysis.

Optional steps: If there is a large amount of RNA residue that needs to be removed, 5µl of RNase A (100mg/ml) solution can be added before adding 200µl of Buffer CB, shaken well, and left at room temperature for 5-10 minutes.

Optional: Use a large diameter suction head to pump the cracked product 2-3 times to assist in cracking.

- 4.4. Add 200µl Buffer CB and 100µl isopropanol, immediately vortex shake and mix thoroughly.
- 4.5. **Centrifuge at 13000rpm for 5 minutes**, add the supernatant to an adsorption column AC, (the adsorption column is placed in the collection tube) Centrifuge at 13000rpm for 1min, and discard the waste liquid in the collection tube.
- 4.6. **Continue to step 5 of the operation.**

**Note:** In the above steps, it is very important to immediately vortex or blow thoroughly mix. Insufficient mixing seriously reduces production. If necessary, if the sample is sticky and difficult to mix, vortex oscillation can be used for 15 seconds to mix well.

5. Add 500µl Buffer PE (please check if anhydrous ethanol has been added first!), 13, Centrifuge at 000 rpm for 30 seconds and discard the waste liquid.
6. Add 600µl Buffer WB (please check if anhydrous ethanol has been added first!), centrifuge at 13000 rpm for 30 seconds, and discard the waste liquid.
7. Repeat step 6.
8. Place the adsorption column AC back into the empty collection tube, centrifuge at 13000rpm for 2 minutes, and try to remove Buffer WB as much as possible to prevent residual ethanol from inhibiting downstream reactions in Buffer WB.
9. Take out the adsorption column AC and place it in a clean centrifuge tube. Add 50-100µl Buffer EB (Buffer EB can be preheated in an 80-100°C water bath to increase yield) to the middle of the adsorption membrane. Let it stand at room temperature for 3-5 minutes, Centrifuge at 000 rpm for 1 minute.

**Note:** 1) The solution obtained from the first elution can be re added to the centrifuge column, left at room temperature for 2 minutes, and centrifuged at 13000 rpm for 1 minute. It can increase the concentration by about 10%.

2) The larger the elution volume, the higher the elution efficiency. If a higher DNA concentration is required, the elution volume can be appropriately reduced, but the minimum volume should not be less than 30µl. Small volume reduces DNA elution efficiency and reduces DNA production.

10. DNA can be stored at -20°C, and if it needs to be stored for a long time, it can be stored at -70°C.