



DANAGENE CIRCULATING DNA MIDIKIT

Ref. 0614.2 5 preps

Ref. 0614.3 50 preps

1. INTRODUCTION

DANAGENE Circulating DNA Midikit provides a fast, reliable and convenient method to purify high quality, high purity and inhibitor-free **cell-free circulating DNA from fresh and frozen plasma / serum samples and other body fluids from samples of 3 ml**

This kit use a specially formulated buffer system allows circulating DNA to bind to the MidiSpin columns based on spin column chromatography proprietary resin separation matrix and MicroSpin for concentrate the circulating cell-free DNA.

Normally the circulating DNA is highly fragmented 50-1000 bp. The degree of fragmentation depends on several parameters such as the origin of DNA (fetal, tumor, microbial DNA), health blood donor, procedure blood collection, handling and storage of the sample.

Features:

- **Efficient recovery and concentration of fragmented DNA (circulating cell-free DNA) with high input and low elution volume.**
- **Sample size: 3000 ul fresh and frozen plasma/serum and other body fluids.**
- **No organic extraction or ethanol precipitation.**
- **Removal of contaminants and inhibitors.**
- **Yield: 0.1-100 ng / ml plasma or serum. Variable because each donor and disease status.**
- **Circulating DNA purified is ready for applications such PCR o real-time PCR, microarrays and Next generation sequencing.**

Applications:

- **Biomarker research and validation for blood-based cancer detection.**
- **Ideal for detection of biomarkers in different diseases like autoimmune diseases, infection diseases, stroke, sepsis, trauma and hematologic disorders. .**
- **Analysis of fetal DNA from maternal plasma.**

2. KIT COMPONENTS

	Ref.0614.2 5 preps	Ref.0614.3 50 preps	T ^a Stock
Lysis Buffer PS	40 ml	375 ml	15-25°C
Proteinase K*	400 µl	2 x 100 mg	-20°C
Desinhibition Buffer*	3 ml	18 ml	15-25°C
Wash Buffer*	3 ml	10 ml	15-25°C
Elution Buffer	3 ml	30 ml	15-25°C
Funnel Columns	5 units	50 units	15-25°C
MicroSpin Columns	5 units	50 units	15-25°C
Collection Tubes	10 units	100 units	15-25°C

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

Both the Lysis Buffer PS , Union Buffer PS, Desinhibition Buffer contain Guanidine salts which is an irritant agent , for this reason we recommend to use gloves and glasses for its manipulation. Guanidine salts can form highly reactive compounds when combined with bleach.

Equipment and additional reagents required

- Water bath
- Votexer
- Ethanol 100 % and isopropanol
- Microcentrifuge capable of 13.000 x g
- Nuclease-free 1.5-2 ml microcentrifuge tubes

3.PROTOCOL

3.1 Collection and sample storage

Plasma/serum samples , blood should be cooled and centrifuged within one hour after blood collection.

Plasma preparation :

1. Centrifuge fresh blood at 1600-2000 x g for 10 minutes.
2. Recover the plasma care not to add the layer containing the cells.
3. Keep the plasma at -80 ° C until DNA extraction.
4. **Thaw the plasma and centrifuge at maximum speed for 5 minutes to remove possible contaminating DNA coming from residual cells. Use the supernatant for DNA extraction.**

3.2 Preliminary preparations

These previous preparations refer to 50 preparations kit (ref.0614.3), the kit of 5 preparations (ref.0614.2) comes with all the solutions ready to use.

- Dissolve the proteinase K in **2 x 1.90 ml** in 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.
- Add **40 ml** of Ethanol 100 % to the Wash Buffer . Keep the container closed to avoid the ethanol evaporation.
- Add **10 ml** of Ethanol 100 % to the Desinhibition Buffer. Keep the container closed to avoid the ethanol evaporation.
- **Pre-heat Elution Buffer at 70°C .**

3.3 Protocol for purification circulating cell-free DNA from 3000 μl plasma/serum and body fluids samples

1. **Pipette 3 ml of plasma / serum** in a 15 ml tube (not supplied) that has previously been centrifuged to remove possible contamination of cells. It is recommended to centrifuge at high speed for 5 minutes, for example 2 samples of 1.5 ml and then join them.
2. Add **7 ml Lysis Buffer PS + 75 μl Proteinase K. Incubate at 55°C** for 30 minutes. Mix by inverting and shaking every 5 minutes.
3. Add **2250 μl of Isopropanol. Mix well.**
4. Insert a MicroSpin column into a funnel. It has to be pressed firmly so that it does not come off during centrifugation. Place the funnel + MicroSpin column in a 50 ml centrifuge tube to collect the waste liquid.
5. Transfer the **12.50 ml of the mixture** from step 3 to the funnel + MicroSpin column.
6. **Centrifuge at 165 x g (1000 r.p.m.) for 5 minutes.** If all the liquid has not passed, centrifuge again at 1100 rpm for 4 minutes. If the microcolumn after centrifugation has been separated from the funnel, it must be firmly replaced and all the 12.5 ml of lysate added again.
7. Place the MicroSpin column in a new collection tube and add **500 μl Desinhibition Buffer** to the reservoir. **Centrifuge at 12.000 rpm for 60 seconds.** Remove the liquid.
8. **Add 500 μl of Wash Buffer** into reservoir of Spin column. **Centrifuge at 14.000 rpm for 60 seconds.** Remove the liquid.
9. **Centrifuge at maximum speed for 3 minutes to remove the residual ethanol.**
10. Remove the collection tube and insert the Spin Column in a 1.5 ml microtube . **Add 35 μl elution buffer (preheated at 70°C)** into reservoir of Spin column. Dispense buffer directly onto the silica membrane. **Incubate at room temperature for 2 min.**

It is very important to add the elution buffer in the center of the membrane to be completely wet.

17. Centrifuge at 10.000 rpm for 60 seconds. Collect 35 μ l and redeposit in the center of the membrane. This increases yield.

18. Incubate 2 minutes and centrifuge at maximum speed for 1 minute.

The microtube now contains the circulating cell-free DNA. The total cf DNA isolated can be quantified using our **Cell-free human DNA detc-qPCR Test** designed to target a conserve sequence region of a gene repeated more than a hundred times in the human genome. Are individuals ready-to-use tubes containing all the components needed to perform the quantitative PCR assay.

4. PROBLEM GUIDE AND POSSIBLE ANSWER

For any doubt or additional consultation on the protocol, do not hesitate to contact with the technical service of DANAGEN-BIOTED S.L info@danagen.es