CatchGene

CatchGene[®] Catch-cfDNA Serum/Plasma 1000 Kit

 Cat. No.
 Rxn

 MC10002_1000
 2

 MC10030_1000
 30

 MC10100_1000
 100

Kit Content

	2rxn	30rxn	100rxn	
LV Module (with 50ml tube)	2	30	100	set
LVR Column	2	30	100	pcs
Collection Tube (2 ml)	4	60	200	pcs
Buffer AE	1.5	2	10	ml
Carrier RNA	12	140	140	μg
Proteinase K	11	11x3	130	mg
Buffer DCL	3	35	144	ml
Buffer CW1 (concentrated)	3.5	50	165	ml
Buffer CW2 (concentrated)	1.68	25	-75	ml
Elution Buffer	0.48	2	7.2	ml

Important Notice !

"LVR Column" should be stored at 4°C upon arrival for long term storage. "Carrier RNA and Proteinase K" should be stored at -20°C upon arrival for long term storage.

Kit Preparation

1. Prepare 20 mg/ml Proteinase K

For 11 mg Proteinase K, please add 0.55 ml Buffer AE into tube and vortex thoroughly for dissolving For 130 mg Proteinase K, please add 6.5 ml Buffer AE into tube and vortex thoroughly for dissolving After dissolving into solvent, please store in 4°C for 6 month or -20°C for 1 year.

Prepare 0.5 μg/μl Carrier RNA For 12 μg Carrier RNA, please add 24 μl Buffer AE into the bottom of tube and mix thoroughly for dissolving. For 140 μg Carrier RNA, please add 280 μl Buffer AE into the bottom of tube and mix thoroughly for dissolving. After dissolving, please aliquot into smaller volume and store at -20°C. Do not freeze-thaw more than three times.

3. Prepare Buffer CW1

Add equal volume of 100% EtOH into Buffer CW1 (concentrated) to get Buffer CW1. After adding 100% EtOH, please check the sticker on the bottle and close the cap tightly.

4. Prepare Buffer CW2

Add equal volume of 100% EtOH into Buffer CW2 (concentrated) to get Buffer CW2. After adding 100% EtOH, please check the sticker on the bottle and close the cap tightly.

Sample Pretreatment

The half life of cfDNA in whole blood is very short. So, after sampling by blood tube, please must perform pretreatment as soon as possible.

For Whole Blood Sample

- 1. Centrifuge whole blood at 3,000 x g for 10 minute at room temperature.
- 2. Transfer upper layer to the 1.5 ml micro-centrifuge tube (not provided). Please avoid aspirating any cell debris or WBC, and intermediate layer, otherwise will co-extract gDNA form intact cell
- 3. Centrifuge at 11,000 x g for 10 min and transfer the supernatant for following extraction.

Please keep samples into -20°C or -70°C if extraction won't be performed immidiately after pretreatment.

General Protocol

For 1ml Serum/Plasma sample

- 1. Add 50 μl Proteinase K (20 mg/ml) into the bottom of 15 ml tube.
- 2. Add 2.5 μl Carrier RNA (0.5 $\mu g/\mu l)$ into the 15 ml tube.
- 3. Transfer 1ml of sample (already centrifuged with high speed) to 15 ml tube, vortex 10 sec to let Proteinase K, Carrier RNA and samples mix thoroughly.

Important! Avoid mixing Proteinase K with Buffer DCL directly. It will cause malfunction of Proteinase K and shortage of cfDNA. Please must mix Proteinase K with sample first, then add Buffer DCL .

- 4. Add 1 ml Buffer DCL to 15 ml tube, vortex 30 sec.
- 5. 56°C incubate for 30 min, then cool down to room temperature (25°C)
- 6. Add 1 ml 100% EtOH, vortex 15 sec.
- 7. Connect LV Module with LVR Column to become LV Column Module. Please refer to the illustration in next page.
- 8. Transfer all lysate into LV Column Module, centrifuge at 2,700 x g for 2 min, discard the flow-through.
- 9. Add 3ml CW1 Buffer into LV Column Module, centrifuge at 2,700 x g for 2 min, discard the flow-through.
- 10. Take LV Column Module out of 50 ml tube. Disconnect the LVR Column from the LV Module, then place the LVR Column on a 2 ml Collection Tube. Please refer to the illustration in next page.
- 11. Add 700 μl CW2 Buffer into spin column, centrifuge at 11,000 x g for 1 min, discard the flow-through.
- 12. Add 700 μ l CW2 Buffer into spin column, centrifuge at 11,000 x g for 1 min, discard the flow-through.
- 13. Add 700 μ l 100% EtOH into spin column, centrifuge at 11,000 x g for 1 min, discard the flow-through.
- 14. Place spin column on a new 2 ml Collection Tube, centrifuge at 11,000 x g for 3 min to eliminate any remaining EtOH.
- Place spin column on a new 1.5 ml micro-centrifuge tube. Add 30-60 μl Elution Buffer, incubation at room temperature for 5 min, and then centrifuge at 11,000 x g for 1 min for elution.

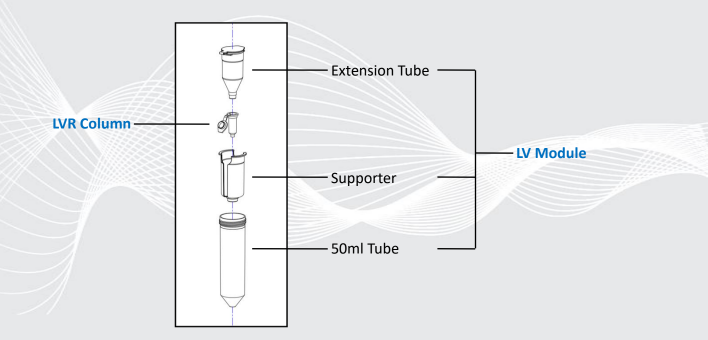
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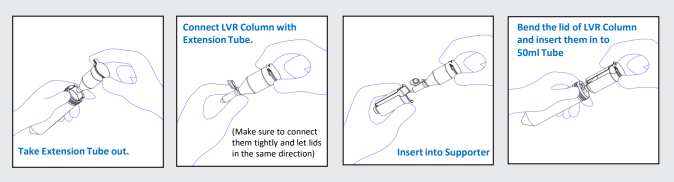


Kit Storage

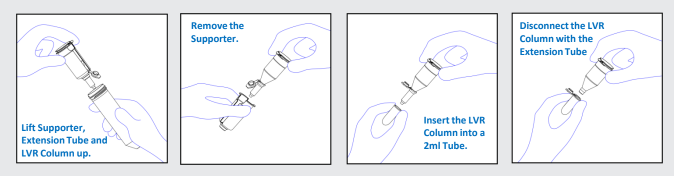
- 1. LVR Column is able to ship at ambient temperature for 2 weeks. After receiving the kit, if you won't use them immediately, please take them out and store at 4°C for long term storage. (Do not freeze into -20 °C)
- 2. Proteinase K and Carrier RNA are lyophilized powder able to ship at ambient temperature for 2 weeks. After receiving the kit, if you won't use them immediately, please take them out and store at -20 °C for long term storage.
- Buffer, solvent and consumables, please store at 15-25 °C.
 If a precipitate has formed in Buffer DCL, please dissolve by incubating at 60°C.



Connect LV Module with LVR Column



Disconnect LVR Column from LV Column Module



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