

# CatchGene® Catch-miRNA Serum/Plasma Kit

Cat. No.	Rxr		
MC20004	4		
MC20050	50		
MC20250	250		

### **Kit Description**

The *Catch-miRNA* Serum/Plasma kit enables purification of 19-24 nucleotides miRNA, small RNA and less than 1000 nucleotides RNA from serum/plasma or urine samples. Based on optimized reagent buffer and silica membrane column, *Catch-miRNA* Serum/plasma kit is able to get high quality and purity of miRNA, which can be used in wide range of downstream application such as qPCR, Microarray and NGS. It provides a convenient and eco-friendly protocol without using phenol or chloroform for RNA purification.

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		4rxn	50rxn	250rxn	
	MC20 Column	4	50	250	pcs
	Collection Tube (2ml)	12	150	750	pcs
	Buffer RCL1	0.36	4.5	22.5	ml
	Buffer RCL2	0.12	1.5	7.5	ml
	Buffer CRW1 (concentrated)	0.48	6	30	ml
	Buffer CRW2 (concentrated)	0.96	12	60	ml
	RNase-Free H <sub>2</sub> O	0.96	12	60	ml

## **Kit Storage**

Upon arrival,

- Please store MC20 Column at 4°C for long term storage.
- 2. Buffer, solvent and consumables, please store at 15-25 °C.

# **Kit Preparation**

#### 1. Prepare Buffer CRW1

Add 4 volume of 100% EtOH into concentrated Buffer CRW1 to get Buffer CRW1. After adding 100% EtOH, please tick the sticker on the bottle and close the cap tightly.

#### 2. Prepare Buffer CRW2

Add 4 volume of 100% EtOH into concentrated Buffer CRW2 to get Buffer CRW2. After adding 100% EtOH, please tick the sticker on the bottle and close the cap tightly.

### **General Protocol**

- 1. Pipette 250  $\mu$ l serum/plasma sample into 1.5 ml micro-centrifuge tube and add 75  $\mu$ l Buffer RCL1. Pulse-vortexing for 10 sec , brief spin down then incubate at 25°C (room temperature) for 3 min.
- 2. Add 25 μl Buffer RCL2, pulse-vortexing for 10 sec, brief spin down then incubate at 25°C (room temperature) for 1 min.
- 3. Centrifuge at 11,000 x g for 3 min.
- Transfer clear supernatant to a new 1.5 ml micro-centrifuge tube, add 330 μl Isopropanol, pulse-vortexing for 10 sec then briefly spin down.
- 5. Transfer all mixture to Spin Column (with 2ml Tube), incubate at 25°C (room temperature) for 2 min.
- 6. Centrifuge at 11,000 x g for 1 min.
- 7. Change a new collection tube, add 500 µl Buffer CRW1 into spin column, centrifuge at 11,000 x g for 1 min.
- 8. Discard the flow-through, add 500  $\mu$ l Buffer CRW2 into spin column, centrifuge at 11,000 x g for 1 min.
- Repeat step 8.
- 10. Change a new collection tube, centrifuge at 11,000 x g for 3 min.
- 11. Place the spin column into 1.5 ml micro-centrifuge tube, add 30-100  $\mu$ l RNase-Free H<sub>2</sub>O and incubate at 25°C (room temperature) for 2 min.
- 12. Centrifuge at 11,000 x g for 1 min for elution.