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# cfKapture™ Kit (200-400 µl)

Manual Revision v1.0

Catalog Nos. CFK-DI10-400UL , CFK-DI50-400UL

## Isolation kit for circulating cell-free DNA from plasma

# PROTOCOL

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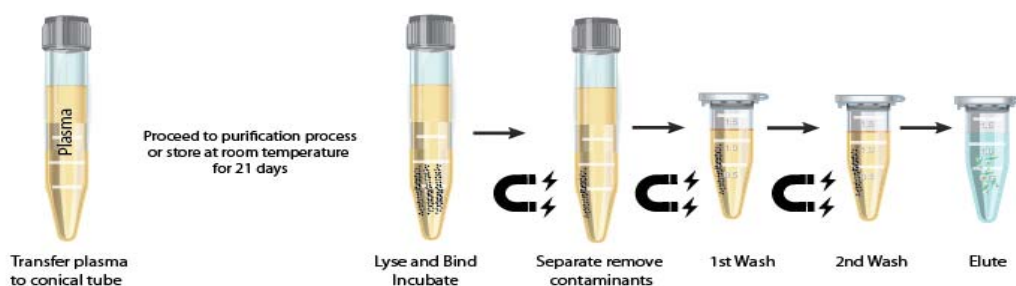
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## Product Description

cfKapture™ kit is a system for purification of circulating, cell free nucleic acids from plasma. It is designed for the purification of circulating cell free DNA(ccfDNA) from maternal and cancer patient's whole blood.

The isolated cfDNA can be directly used for real time-PCR and DNA library preparation suitable for next generation sequencing. **This kit is designed for research purposes only.**

## Process



## Kit Contents and Storage

cfKapture Kit Catalog No.	CFK-DI10-400UL	CFK-DI50-400UL	STORAGE
Number of Preps	10	50	
CFL Buffer	5 ml	22 ml	15-25°C
CFW1 Buffer <sup>1</sup>	8 ml	40 ml	15-25°C
CFW2 Buffer <sup>1</sup>	5 ml	25 ml	15-25°C
Elution Buffer	600 µl	3 ml	15-25°C
Pro K Solution <sup>2</sup>	220 µl	1.1 ml	2-8°C
MAG-CFB Particles	220 µl	1.1 ml	2-8°C

<sup>1</sup> Ethanol must be added prior to use. See Preparation of Reagents

## Stability

All components are stable for 12 months when stored accordingly.

- Pro K Solution comes in a ready to use solution. Component is stable for 1 year when stored at 15-25°C. For storage longer than 1 year, storage at 2-8°C is recommended.
- During shipment or storage in cool ambient conditions, precipitates may form in some buffers. Dissolve such deposits by warming the solution at 37°C and then gently shaking the buffer.

## Safety Information

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate material safety data sheets (MSDSs). MSDS can be downloaded from the "Product Resource" tab when viewing the product kit.

## Equipments Required

Catalog No.	Description
<b>MBMS-10</b>	Magnetic Separation Device for 2ml microcentrifuge tube

## Preparation of Reagents

Prepare the following components for each kit before use:

Catalog No.	Component	Add 100% Ethanol	Storage
<b>CFK-DI10-400UL</b>	CFW1 Buffer	10 ml	15-25°C
	CFW2 Buffer	13 ml	15-25°C
Components are stable for 1 year when stored closed at room temperature			

Catalog No.	Component	Add 100% Ethanol	Storage
<b>CFK-DI50-400UL</b>	CFW1 Buffer	50 ml	15-25°C
	CFW2 Buffer	65 ml	15-25°C
Components are stable for 1 year when stored closed at room temperature			



## CF-Kapture™ 21 Kit : 200 µl plasma sample protocol

### Equipment and Reagents to Be Supplied by User

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate material safety data sheets (MSDSs) from each product supplier.


- Nuclease-free 1.5 ml microcentrifuge tubes
- 15 ml conical tubes
- magnetic separation device for 1.5 ml microcentrifuge tube (CAT# MBMS-10)
- 100% Ethanol
- Vortex
- Tube rotator
- Water bath, incubator or heat block capable of 60°C

### Things to do before starting

- Prepare CFW1 Buffer, CFW2 Buffer according to the "Preparation of Reagents" section on page 2
- Preheat and warm water bath, incubator or heat block to 60°C
- Warm Elution Buffer to 60°C

### Isolation steps

#### Binding Steps

- 1. Transfer 200 µl plasma to 1.5 ml conical tube.**
- 2. Add 20 µl Pro K Solution, and mix well by vortexing at maximum speed for 20 seconds.**
- 3. Incubate sample at 60°C for 10 minutes in a water bath. Mix by inverting the tube once during incubation.**
- 4. Add 200 µl CFL Buffer and mix well by vortexing at maximum speed for 60 seconds, and incubate sample at room temperature for 5 minutes.**
- 5. Add 150 µl 100% ethanol and 20 µl MAG-CFB Particles. Mix immediately by vortexing at maximum speed for 20 seconds.**  
 Complete resuspension of the MAG-CFB Particles is crucial for obtaining purity.
- 6. Incubate sample tube on a tube rotator for 20 minutes at room temperature at 10 rpm.**  
Adjust rotator angle to approximately 45 degrees for better mixing.
- 7. Remove the tube from rotator and place on a compatible 1.5 ml magnetic separation device (CAT# MBMS-10) to magnetize the MAG-CFB Particles for 5 minutes or until the magnetic particles are completely cleared from solution.**
- 8. With the tube on the magnetic separation device, remove and discard the cleared supernatant by pipetting.** Do not disturb the attracted beads while aspirating the supernatant.

### Wash Steps

9. **Remove the sample tube off the magnetic separation device.**
10. **Add 800 µL CFW1 Buffer and resuspend the magnetic particles by vortexing at maximum speed for 1 minute or by pipetting up and down 10 times.**
11. **Place the 1.5 ml sample tube on a compatible 1.5 ml magnetic separation device (CAT#MBMS-10) to magnetize the MAG-CFB Particles at room temperature for 5 minutes or until the magnetic particles are completely cleared from solution.**
12. **Remove and discard the cleared supernatant.** Do not disturb the attracted beads while aspirating the supernatant.
13. **Remove the sample off the magnetic separation device and repeat wash by adding 800µl CFW1 Buffer and resuspend the magnetic particles by vortexing at maximum speed for 1 minute or by pipetting up and down 10 times.**
14. **Place the 1.5 ml sample tube back on the magnetic separation device to magnetize the MAG-CFB Particles at room temperature for 5 minutes or until the magnetic particles are completely cleared from solution.**
15. **With the sample still on the magnetic separation device, remove and discard the cleared supernatant.** Do not disturb the attracted beads while aspirating the supernatant.
16. **Remove the sample tube off the magnetic separation device.**
17. **Add 800 µL CFW2 Buffer and resuspend the magnetic particles by vortexing at maximum speed for 1 minute or by pipetting up and down 10 times.**
18. **Place the 1.5 ml sample tube on the magnetic separation device to magnetize the MAG-CFB Particles at room temperature for 5 minutes or until the magnetic particles are completely cleared from solution.**
19. **With the sample still on the magnetice separation device, remove and discard the cleared supernatant.** Do not disturb the attracted beads while aspirating the supernatant.
20. **Repeat Steps 19-22 for a second CFW2 Buffer wash step.**
21. **With the sample tube still on the magnetic separation device, air dry the MAG-CFB Particles for 10 minutes. Remove any residual liquid with a pipette.**  
Note: It is critical to completely remove all liquid from the tube.

### Elution Steps

22. **Remove the sample tube containing the MAG-CFB Particles off the magnetic separation device.**
23. **Add 30-50 µl Elution Buffer and completely resuspend the MAG-CFB Particles by vortexing**

**at maximum speed for 10 seconds.**

Note: Heat Elution Buffer at 60°C to improve yield.

- 24. Incubate at room temperature for 10 minutes.**
- 25. Place the tubes back on the magnetic separation device and wait 5 minutes or until the magnetic particles are completely cleared from elution buffer.**
- 26. Transfer the clear supernatant containing the ccfDNA to a new 1.5 ml microcentrifuge tube and store at -20°C.**

## cfKapture™ Kit : 400 µl plasma sample

### Equipment and Reagents to Be Supplied by User

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate material safety data sheets (MSDSs) from each product supplier.


- Nuclease-free 2.0 ml microcentrifuge tubes
- 15 ml conical tubes
- magnetic separation device for 2.0 ml microcentrifuge tube (CAT# MBMS-10)
- 100% Ethanol
- Vortex
- Tube rotator
- Water bath, incubator or heat block capable of 60°C

### Things to do before starting

- Prepare CFW1 Buffer, CFW2 Buffer according to the “Preparation of Reagents” section on page 2
- Preheat and warm water bath, incubator or heat block to 60°C
- Warm Elution Buffer to 60°C

### Isolation steps

#### Binding Steps

- 1. Transfer 400µl plasma sample to a 2 ml centrifuge tube.**
- 2. Add 20 µl Pro K Solution, and mix well by vortexing at maximum speed for 20 seconds.**
- 3. Incubate sample at 60°C for 10 minutes in a water bath. Mix by inverting the tube once during incubation.**
- 4. Add 400µl CFL Buffer and mix well by vortexing at maximum speed for 60 seconds, and incubate sample at room temperature for 10 minutes.**
- 5. Add 300µl 100% ethanol and 20 µl MAG-CFB Particles. Mix immediately by vortexing at maximum speed for 20 seconds.**  
 Complete resuspension of the MAG-CFB Particles is crucial for obtaining purity.
- 6. Incubate sample tube on a tube rotator for 20 minutes at room temperature at 10 rpm.**  
Adjust rotator angle to approximately 45 degrees for better mixing.
- 7. Remove the tube from rotator and place on a compatible 2 ml magnetic separation device (CAT# MBMS-10) to magnetize the MAG-CFB Particles for 5 minutes or until the magnetic particles are completely cleared from solution.**
- 8. With the tube on the magnetic separation device, remove and discard the cleared supernatant by pipetting.** Do not disturb the attracted beads while aspirating the supernatant.



### Wash Steps

9. **Remove the sample tube off the magnetic separation device.**
10. **Add 800  $\mu$ L CFW1 Buffer and resuspend the magnetic particles by vortexing at maximum speed for 1 minute or by pipetting up and down 10 times.**
11. **Place the 2 ml sample tube on a compatible 2 ml magnetic separation device (CAT#MBMS-10) to magnetize the MAG-CFB Particles at room temperature for 5 minutes or until the magnetic particles are completely cleared from solution.**
12. **Remove and discard the cleared supernatant.** Do not disturb the attracted beads while aspirating the supernatant.
13. **Remove the sample off the magnetic separation device and repeat wash by adding 800 $\mu$ l CFW1 Buffer and resuspend the magnetic particles by vortexing at maximum speed for 1 minute or by pipetting up and down 10 times.**
14. **Place the 2 ml sample tube back on the magnetic separation device to magnetize the MAG-CFB Particles at room temperature for 5 minutes or until the magnetic particles are completely cleared from solution.**
15. **Remove and discard the cleared supernatant.** Do not disturb the attracted beads while aspirating the supernatant.
16. **Remove the sample tube off the magnetic separation device.**
17. **Add 800  $\mu$ L CFW2 Buffer and resuspend the magnetic particles by vortexing at maximum speed for 1 minute or by pipetting up and down 10 times.**
18. **Place the 2 ml sample tube on the magnetic separation device to magnetize the MAG-CFB Particles at room temperature for 5 minutes or until the magnetic particles are completely cleared from solution.**
19. **Remove and discard the cleared supernatant.** Do not disturb the attracted beads while aspirating the supernatant.
20. **Repeat Steps 19-22 for a second CFW2 Buffer wash step.**
21. **With the sample tube on the magnetic separation device, air dry the MAG-CFB Particles for 10 minutes. Remove any residual liquid with a pipette.**  
Note: It is critical to completely remove all liquid from the tube.

### Elution Steps

22. **Remove the sample tube containing the MAG-CFB Particles off the magnetic separation device.**
23. **Add 30-50  $\mu$ L Elution Buffer and completely resuspend the MAG-CFB Particles by vortexing**

**at maximum speed for 10 seconds.**

Note: Heat Elution Buffer at 60°C to improve yield.

- 24. Incubate at room temperature for 10 minutes.**
- 25. Place the tubes back on the magnetic separation device and wait 5 minutes or until the magnetic particles are completely cleared from elution buffer.**
- 26. Transfer the clear supernatant containing the ccfDNA to a new 1.5 ml microcentrifuge tube and store at -20°C.**

## Troubleshooting guide

Please use this guide to troubleshoot any problems that may arise. For further assistance, please contact technical support via:

Phone: 1-855-262-4246 (in US), outside US, 1-301-302-0144

Email: [support@magbiogenomics.com](mailto:support@magbiogenomics.com)

Symptoms	Possible Causes	Comments

## Ordering Information

Catalog No.	Product	Description	Preps
CFK-DI10-400UL	cfKapture - 400µl (10 preps)	Purification of cell-free DNA (cfDNA) from 400 µl plasma	10
CFK-DI5-2ML	cfKapture - 2ml (5 preps)	Purification of cell-free DNA (cfDNA) from 2 ml plasma	5
CFK-DI5-5ML	cfKapture - 5ml (5 preps)	Purification of cell-free DNA (cfDNA) from 5 ml plasma	5
CFK-DI50-400UL	cfKapture - 400µl (50 preps)	Purification of cell-free DNA (cfDNA) from 400 µl plasma	50
CFK-DI50-2ML	cfKapture - 2ml (50 preps)	Purification of cell-free DNA (cfDNA) from 2 ml plasma	50
CFK-DI50-5ML	cfKapture - 5 ml (50 preps)	Purification of cell-free DNA (cfDNA) from 5 ml plasma	50

## Magnetic Separation Devices

Catalog No.	Description
MBMS-31550	15ml and 50ml magnetic stand combo. (3x15ml and 3x50ml)
MBMS-10	MagStip magnetic stand (1.5mL x 10)









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