## **User's Manual and Instructions**

## cfPure<sup>®</sup> Cell-Free DNA Extraction Kit

### Catalog Number: K5011610, K5011625, K5011625MA

### **Storage Conditions**

Store all of the contents of this kit at room temperature

### Shelf Life

1 year from the date of receipt under proper storage conditions

### Features

- Non-toxic chemicals
- High Cell-Free DNA recovery
- Short and Scalable Protocol
- Fully automatable using KingFisher Flex Purification System and Robotic Liquid Handlers
- Purified DNA is suitable for NGS, PCR, Bisulfite sequencing, etc

### Description

BioChain's cfPure<sup>®</sup> Cell-Free DNA extraction kit allows for fast and efficient Cell-Free DNA (cfDNA) isolation from plasma/serum samples. The magnetic bead-based extraction protocol is ideally suited for use with robotic liquid handlers and King Fisher Flex Purification System. This kit may also be applied on Hamilton's Pestle system with the appropriate program. DNA extracted using this kit is suitable for PCR, qPCR, next generation sequencing (NGS), and other applications.

### Contents

All necessary reagents for cfDNA isolation from human plasma samples are provided. There are 3 package sizes for this kit, which contains sufficient reagents for isolating cfDNA from up to 100 ml, 250 ml, or 250 ml's of sample in 5 - 10 ml increments.

### **Quality Control**

Each component has been tested for purity and efficacy.

### **Important Notes**

<u>Blood Collection:</u> The cfPure<sup>®</sup> Cell-Free DNA extraction kit has been optimized for use with samples collected in Streck Cell-Free DNA BCT, EDTA tubes and Acid Dextrose Acid (ACD) tubes.

<u>Starting Material</u>: Both fresh and frozen plasma can be used with the Cell-Free DNA isolation protocol. Fresh plasma, however, tends to have higher yields.

<u>Quantification:</u> Plasma will yield 1-100 ng of Cell-Free DNA per ml of plasma. Therefore, quantification by absorbance measurement (eg. Nanodrop) may not be sensitive enough to accurately determine yield. Instead, we suggest using Qubit<sup>™</sup> dsDNA High Sensitivity Assay..

<u>Recommendations for PCR</u>: Due to the highly fragmented nature of the nucleic acids obtained from plasma, care should be taken in the design of primers. Cell-free DNA tends to have a small size (~170bp). Therefore, PCR primers should be designed to produce amplicons of 150 bp or less. Given the low concentration of cfDNA in plasma taken from healthy individuals, 40 amplification cycles may be needed in some cases.

F-753-3UMRevD

K50116XXUC

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<u>Streck Cell-Free DNA BCT Tube(s)</u>: Plasma from blood collected with Streck Cell-Free DNA BCT Tube(s) must go through a Proteinase K treatment prior to Cell-Free DNA isolation to ensure optimal yields. Forgoing Proteinase K treatment may decrease yields by 50%

### Equipment and Reagents to be Supplied by User

- o Pipettes
- Vortex-Genie 2 or similar vortexing mixer\*
- Magnet stand for molecular applications (e.g. DynaMag<sup>™</sup>-15 or DynaMag<sup>™</sup>-2)
- 1.5 ml non-stick eppendorf tube(s)
- Fresh 100% EtOH

\* Contact BioChain<sup>®</sup> Technical Service for additional recommendations for high throughput or automated mixing.



### Prior to Initial Use

The Lysis/Binding and Wash Buffer are shipped as a concentrate. If precipitate is present in either solution, incubate solution at 37°C for 30 minutes. 100% EtOH must be added to both solutions before the first use. Once EtOH is added, these buffers are stable for one year if stored at 4°C. Be sure to close the bottle tightly for long term storage.

### For 100 ml kit (K5011610)

- Add 23 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 51 ml of fresh 100% ethanol to each bottle of Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction For 250 ml kits (K5011625)
- Add 19 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 51 ml of fresh 100% ethanol to each bottle of Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction For 250 ml Max kits (K5011625MA)
- Add 19 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 60 ml of fresh 100% ethanol to Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction

Before starting the protocol, determine the amount of plasma to be used for extraction and calculate the amount of buffer and beads needed. Any amount from 100  $\mu$ l to 10 ml of plasma can be used. Scale buffer and bead volumes accordingly using the table below.

Plasma	Lysis/Binding Buffer	Bead Solution*	Tube(s) size
x (x=ml of plasma)	1.25x	0.025x	n/a
5 ml	6.25 ml	125 µl	15 ml or 50ml**
7 ml	8.75 ml	175 µl	50 ml

## Small (0.2 ml to 7.9 ml) Sample Protocol

 \* Important: for sensitive PCR applications where inhibition is a concern, use 0.008x instead (e.g. 40 µl of bead solution for 5 ml plasma and 56 µl of bead solution for 7 ml plasma
 \*\*Using a 50 ml tube(s) for 5 ml or more of plasma is recommended over a 15 ml tube(s). While a 15 ml tube(s) will work it may lead to slightly lower yields

### Proteinase K Treatment

If samples were collected using a **Streck Cell-Free DNA BCT tube(s)**, Proteinase K treatment is required to ensure optimal yields. If blood was not collected with **Streck Cell-Free DNA BCT tube(s)** proceed to the Lysis/Binding step.



Plasma	Proteinase K	20% SDS Solution
x (x=ml of plasma)	0.015 x	0.050 x
5 ml	75 µl	250 μl
7 ml	105 µl	350 µl

- 1. Add the appropriate amount of plasma to an appropriately sized tube(s)
- 2. Add 15 µl of Proteinase K (20 mg/ml) for every 1 ml of plasma used
- 3. Add 50 µl of 20% SDS solution for every 1 ml of plasma used
- 4. Mix by inverting gently 5 times
- 5. Incubate at 60°C for 20 minutes
- 6. After incubation place tube(s) on ice for 5 minutes to cool tube(s) to room temperature
- 7. Once tube(s) are at room temperature proceed to step 2 of the Lysis/Binding section

### Lysis/ Binding

- 1. Add the appropriate amount of plasma to appropriately sized tube(s)
- 2. Add 1.25 ml of cfPure Lysis/Binding Buffer for every 1 ml of plasma used
- 3. Add 25 µl (or 8 µl) of cfPure Magnetic Bead Solution for every 1 ml of plasma Important: Mix beads well prior to adding. There should be no visible sedimentation at the bottom of the solution after mixing. Beads will settle quickly, so be sure to mix the magnetic bead solution after adding to each sample. Failure to do so may result in inconsistent yields
- Vortex or shake tube(s) vigorously for 10 minutes at room temperature
  \* To obtain high yields, ensure that plasma/buffer solution is mixing vigorously in tube(s). A vortexing mixer with a tube(s)-holder that allows for walk-away mixing will make this easier.
- 5. Place tube(s) into a magnet stand for 2 to 5 minutes, or until solution clears
- **6.** While keeping the tube(s) on the magnet stand, remove supernatant. Be careful not to remove magnetic particles
- 7. Keep tube(s) on magnet stand for 1 minute, and remove residual supernatant

### **First Wash**

- 8. Add 1000 µl of cfPure Wash Buffer to lysis/binding tube(s)
- 9. Resuspend beads by vortexing for 10 seconds or pipetting up and down 10 times
- 10. Transfer magnetic particle suspension into 1.5 ml micro tube(s) on magnet stand
- 11. Allow beads to attach to magnet stand for 10-30 seconds
- 12. Pipette supernatant from 1.5 ml tube(s) and use the supernatant to wash the



lysis/binding tube(s)

- 13. Transfer the rest of the magnetic particles in lysis/binding tube(s) to the 1.5 ml tube(s)
- **14.** Keep tube(s) on magnet stand for 10-30 seconds or until solution is clear
- 15. Remove as much buffer as possible using a 1000 µl pipette
- **16.** Tap magnet stand on bench 5 times and remove remaining wash buffer with 200 µl pipette
- 17. Transfer tube(s) to non-magnetic rack and add 1000 µl of cfPure Wash Buffer
- 18. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 19. Centrifuge tube(s) briefly

\*Centrifuge steps are needed when using vortexing to resuspend beads. Only a brief spin is recommended to remove solution from tube(s) lid

- 20. Place tube(s) on magnet stand for 10-30 seconds
- 21. Remove as much buffer as possible using a 1000 µl pipette
- 22. Tap magnet stand on bench 5 times and remove remaining wash buffer with 200  $\mu I$  pipette

### Second Wash

- 23. Transfer tube(s) to non-magnetic rack and add 1000 µl of 80% EtOH
- 24. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 25. Centrifuge tube(s) briefly
- 26. Place on magnet stand for 10-30 seconds or until solution clears
- 27. Remove as much buffer as possible using a 1000 µl pipette
- 28. Tap magnet stand on bench 5 times and remove remaining EtOH with 200 µl pipette
- 29. Transfer tube(s) to non-magnetic rack and add 1000 µl of 80% EtOH
- 30. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 31. Centrifuge tube(s) briefly
- 32. Place on magnet stand for 10-20 seconds
- 33. Remove as much EtOH as possible using a 1000 µl pipette and leave cap open
- 34. Tap magnet stand with tube(s) on bench 5 times
- 35. Remove remaining EtOH with 200 µl pipette
- **36.** Leave tube(s) open on magnet stand for two minutes and then tap tube(s) on bench 5 times and remove any remaining EtOH with 20 µl pipette
- **37.** Allow magnetic particles to dry for an additional 1-3 minutes \*Be careful to not over dry or beads may stick to tube(s)

### **Elution Step**

38. Transfer microtube(s) to non-magnetic rack and add desired volume of cfPure



### Elution Buffer and resuspend beads

**Important::** A minimum of 12.5 µl of cfPure Elution Buffer per ml of plasma is recommended to elute DNA to ensure optimal yields

- 39. Vortex or shake tube(s) vigorously for 5 minutes
- 40. Centrifuge tube(s) briefly
- **41.** Place tube(s) on magnetic rack for 10 to 30 seconds
- 42. Transfer elute into a new 1.5 ml tube(s)

## Large (8 ml to 20 ml) Sample Extraction Protocol

This protocol is optimized for use with samples with volumes of 8 ml or larger.

Plasma	Lysis/Binding Buffer	Bead Solution*	Tube(s) size
x (x=ml of plasma)	1.25x	0.020x	n/a
8 ml	10.00 ml	160 µl	50 ml
10 ml	12.50 ml	200 µl	50 ml

\* **Important:** for sensitive PCR applications where inhibition is a concern, use 0.008x instead (e.g. 64 µl of bead solution for 8 ml plasma and 80 µl of bead solution for 10 ml plasma

### **Proteinase K Treatment**

If samples were collected using a **Streck Cell-Free DNA BCT tube(s)**, Proteinase K treatment is required to ensure optimal yields. If blood was not collected with **Streck Cell-Free DNA BCT tube(s)** proceed to the Lysis/Binding step.

Plasma	Proteinase K	20% SDS Solution
x (x=ml of plasma)	0.015 x	0.050 x
8 ml	120 µl	400 µl
10 ml	150 µl	500 µl

- 1. Add the appropriate amount of plasma to an appropriately sized tube(s)
- 2. Add 15 µl of Proteinase K (20 mg/ml) for every 1 ml of plasma used
- 3. Add 50 µl of 20% SDS solution for every 1 ml of plasma used
- 4. Mix by inverting gently 5 times
- 5. Incubate at 60°C for 20 minutes



- 6. After incubation place tube(s) on ice for 5 minutes to cool tube(s) to room temperature
- 7. Once tube(s) are at room temperature proceed to step 2 of the Lysis/Binding section

### Lysis/ Binding

- 1. Add appropriate amount of plasma to a 50 ml conical tube(s)
- 2. Add 1.25 ml of cfPure Lysis/Binding Buffer for every 1 ml of plasma used
- 3. Add 20 µl (or 8 µl) of cfPure Magnetic Bead Solution for every 1 ml of plasma used Important: Mix beads well prior to adding. There should be no visible sedimentation at the bottom of the solution after mixing. Beads will settle quickly, so be sure to mix the magnetic bead solution after adding to each sample. Failure to do so may result in inconsistent yields
- Vortex or shake tube(s) vigorously for 10 minutes at room temperature
  \* To obtain high yields, ensure that plasma/buffer solution is mixing vigorously in tube(s). A vortexing mixer with a tube(s)-holder that allows for walk-away mixing will make this easier.
- 5. Place tube(s) into a magnet stand for 5 to 10 minutes, or until solution clears
- **6.** While keeping the tube(s) on the magnet stand, remove supernatant. Be careful not to remove magnetic particles
- 7. Keep tube(s) on magnet stand for 1 minute, and remove residual supernatant

### First Wash

- 8. Add 2000 µl of cfPure Wash Buffer to lysis/binding tube(s)
- 9. Resuspend beads by vortexing for 20 seconds or by manually swirling tube(s)
- 10. Place 50 ml tube(s) on to a magnetic stand for 2 to 5 minutes, or until solution clears
- 11. Using a 1000 ul pipette wash the tube(s) using the supernatant within the tube Important: Washing the side of the tube(s) will ensure all beads are attached to the magnet. The cap of the tube may also need to be washed if beads have stuck to the cap.
- 12. Keep 50 ml tube(s) on to a magnetic stand for an additional 2 minutes
- **13.** While keeping the tube(s) on the magnet stand, remove supernatant. Be careful not to remove magnetic particles
- 14. Transfer tube(s) to non-magnetic rack and add 1000 µl of cfPure Wash Buffer
- 15. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 16. Transfer magnetic particle suspension into 1.5 ml micro tube(s) on a magnet stand
- 17. Allow beads to attach to magnet stand for 10-30 seconds
- **18.** Pipette supernatant from 1.5 ml tube(s) and use the supernatant to wash the lysis/binding tube(s)
- **19.** Transfer the rest of the magnetic particles in lysis/binding tube(s) to the 1.5 ml tube(s)



- 20. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 21. Centrifuge tube(s) briefly \*Centrifuge steps are needed when using vortexing to resuspend beads. Only a brief spin is recommended to remove solution from tube(s) lid
- 22. Place tube(s) on magnet stand for 10-30 seconds or until solution is clear
- 23. Remove as much buffer as possible using a 1000 µl pipette
- 24. Tap magnet stand on bench 5 times and remove remaining wash buffer with 200 µl pipette

### Second Wash

- 25. Transfer tube(s) to non-magnetic rack and add 1000 µl of 80% EtOH
- 26. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 27. Centrifuge tube(s) briefly
- 28. Place on magnet stand for 10-30 seconds or until solution clears
- 29. Remove as much buffer as possible using a 1000 µl pipette
- 30. Tap magnet stand on bench 5 times and remove remaining EtOH with 200 µl pipette
- 31. Transfer tube(s) to non-magnetic rack and add 1000 µl of 80% EtOH
- 32. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 33. Centrifuge tube(s) briefly
- **34.** Place on magnet stand for 10-20 seconds
- 35. Remove as much EtOH as possible using a 1000 µl pipette and leave cap open
- 36. Tap magnet stand with tube(s) on bench 5 times
- 37. Remove remaining EtOH with 200 µl pipette
- **38.** Leave tube(s) open on magnet stand for two minutes and then tap tube(s) on bench 5 times and remove any remaining EtOH with 20 μI pipette
- **39.** Allow magnetic particles to dry for an additional 4 minutes \*Be careful to not over dry or beads may stick to tube(s)

### **Elution Step**

**40.** Transfer microtube(s) to non-magnetic rack and add desired volume of **cfPure Elution Buffer** and resuspend beads

Important: For optimal yields a minimum of 100 µl of cfPure Elution Buffer should be used.

- **41.** Vortex or shake tube(s) vigorously for 5 minutes
- 42. Centrifuge tube(s) briefly
- 43. Place tube(s) on magnetic rack for 10 to 30 seconds
- **44.** Transfer eluate into a new 1.5 ml tube(s)



## Alternative Low Elution Volume Protocol

The low elution volume protocol is recommended when 50 ul or less of elution when extracting from 10 ml samples.

Plasma	Lysis/Binding Buffer	Bead Solution	Tube(s) size
10 ml	12.50 ml	75 µl	50 ml

### Proteinase K Treatment

If samples were collected using a **Streck Cell-Free DNA BCT tube(s)**, Proteinase K treatment is required to ensure optimal yields. If blood was not collected with **Streck Cell-Free DNA BCT tube(s)** proceed to the Lysis/Binding step.

Plasma	Proteinase K	20% SDS Solution
x (x=ml of plasma)	0.015 x	0.050 x
10 ml	150 µl	500 µl

- 1. Add the 10 ml's of plasma to an appropriately sized tube(s)
- 2. Add 150 µl of Proteinase K (20 mg/ml)
- 3. Add 500 µl of 20% SDS solution
- 4. Mix by inverting gently 5 times
- 5. Incubate at 60°C for 20 minutes
- 6. After incubation place tube(s) on ice for 5 minutes to cool tube(s) to room temperature
- 7. Once tube(s) are at room temperature proceed to step 2 of the Lysis/Binding section

### Lysis/ Binding

- 1. Add the 10 ml of plasma to 50 ml tube(s)
- 2. Add 12.5 ml of cfPure Lysis/Binding Buffer to the tube(s)
- 3. Add 75 µl of cfPure Magnetic Bead Solution for to the tube(s) Important: Mix beads well prior to adding. There should be no visible sedimentation at the bottom of the solution after mixing. Beads will settle quickly, so be sure to mix the magnetic bead solution after adding to each sample. Failure to do so may result in inconsistent yields
- Vortex or shake tube(s) vigorously for 10 minutes at room temperature
  \* To obtain high yields, ensure that plasma/buffer solution is mixing vigorously in tube(s). A vortexing mixer with a tube(s)-holder that allows for walk-away mixing will make this easier.
- 5. Place tube(s) into a magnet stand for 5 to 10 minutes, or until solution clears



- **6.** While keeping the tube(s) on the magnet stand, remove supernatant. Be careful not to remove magnetic particles
- 7. Keep tube(s) on magnet stand for 1 minute, and remove residual supernatant

### **First Wash**

- 8. Add 1000 µl of cfPure Wash Buffer to lysis/binding tube(s)
- **9.** Resuspend beads by swirling tube or pipetting up and down 10 times
- 10. Transfer magnetic particle suspension into 2 ml micro tube(s) on magnet stand
- 11. Allow beads to attach to magnet stand for 20-30 seconds or until solution clears
- **12.** Pipette supernatant from 2 ml tube(s) and use the supernatant to wash the lysis/binding tube(s)
- 13. Transfer the rest of the magnetic particles in lysis/binding tube(s) to the 2 ml tube(s)
- 14. Keep tube(s) on magnet stand for 20-30 seconds or until solution is clear
- 15. Remove as much buffer as possible using a 1000 µl pipette
- 16. Tap magnet stand on bench 5 times and remove remaining wash buffer with 200 µl pipette
- 17. Transfer tube(s) to non-magnetic rack and add 1000 µl of cfPure Wash Buffer
- 18. Resuspend beads by vortexing for 25 seconds or pipetting up and down 10 times
- 19. Centrifuge tube(s) briefly
  \*Centrifuge steps are needed when using vortexing to resuspend beads. Only a brief spin is recommended to remove solution from tube(s) lid
- 20. Place tube(s) on magnet stand for 20-30 seconds or until solution clears
- 21. Remove as much buffer as possible using a 1000 µl pipette
- 22. Tap magnet stand on bench 5 times and remove remaining wash buffer with 200 µl pipette

### Second Wash

- 23. Transfer tube(s) to non-magnetic rack and add 1000 µl of 80% EtOH
- 24. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 25. Centrifuge tube(s) briefly
- 26. Place on magnet stand for 20-30 seconds or until solution clears
- 27. Remove as much buffer as possible using a 1000 µl pipette
- 28. Tap magnet stand on bench 5 times and remove remaining EtOH with 200 µl pipette
- 29. Transfer tube(s) to non-magnetic rack and add 1000 µl of 80% EtOH
- 30. Resuspend beads by vortexing for 20 seconds or pipetting up and down 10 times
- 31. Centrifuge tube(s) briefly
- 32. Place on magnet stand for 20-30 seconds or until solution clears



- **33.** Remove as much EtOH as possible using a 1000 µl pipette and leave cap open
- **34.** Tap magnet stand with tube(s) on bench 5 times
- 35. Remove remaining EtOH with 200 µl pipette
- **36.** Leave tube(s) open on magnet stand for two minutes and then tap tube(s) on bench 5 times and remove any remaining EtOH with 20 µl pipette
- **37.** Allow magnetic particles to dry for an additional 4 minutes \*Be careful to not over dry or beads may stick to tube(s)

### **Elution Step**

- **38.** Transfer microtube(s) to non-magnetic rack and add 50 ul of **cfPure Elution Buffer** and resuspend beads
- **39.** Vortex or shake tube(s) vigorously for 5 minutes
- **40.** Centrifuge tube(s) briefly
- **41.** Place tube(s) on magnetic rack for 20 to 30 seconds or until solution clears
- **42.** Transfer eluate into a new 1.5 ml tube(s)

## Kit Components cfPure cfDNA Extraction, 100ml Kit (K5011610)

Item	Cat#	Amount	Storage
1. Lysis/Binding Buffer	K5011610-1	1 x 115 ml	Room Temp
2. Wash Buffer	K5011610-2	2 x 55 ml	Room Temp
3. Elution Buffer	K5011610-3	1 x 6 ml	Room Temp
4. Magnetic Bead Solution	K5011610-4	2 x 1.33 ml	Room Temp

## Kit Components cfPure cfDNA Extraction, 250ml Kit (K5011625)

Item	Cat#	Amount	Storage
1. Lysis/Binding Buffer	K5011625-1	3 x 95 ml	Room Temp
2. Wash Buffer	K5011625-2	5 x 55 ml	Room Temp
3. Elution Buffer	K5011625-3	1 x 15 ml	Room Temp
4. Magnetic Bead Solution	K5011625-4	5 x 1.33 ml	Room Temp

## Kit Components cfPure MAX cfDNA Extraction, 250ml Kit (K5011625MA)

Item	Cat#	Amount	Storage
1. Lysis/Binding Buffer	K5011625MA-1	3 x 95 ml	Room Temp
2. Wash Buffer	K5011625MA-2	1 x 65 ml	Room Temp
3. Elution Buffer	K5011625MA-3	1 x 15 ml	Room Temp
4. Magnetic Bead Solution	K5011625MA-4	5 x 1.33 ml	Room Temp



# cfPure<sup>™</sup> Cell-Free DNA Extraction Kit

Isolation of cfDNA from 1 - 5 ml of sample using KingFisher<sup>TM</sup> Flex Magnetic Processor 24DW

### Catalog Number: K5011610, K5011625

### **Product Description**

Biochain's new cfPure Cell-Free DNA Extraction Kit has been designed to isolate circulating DNA from human plasma and serum. The cfPure utilizes Silica coated magnetic bead technology allowing for scalability and easy automation. The cfPure Cell-Free DNA extraction Kit can be used to isolate cfDNA from up to 24 samples of 1 - 5 ml of plasma or serum using the KingFisher<sup>™</sup> Flex Magnetic Processor with 24 Deep Well Head. This guide describes the use of the cfPure kit with the KingFisher<sup>™</sup> Flex Magnetic Processor 24DW to process samples of 1 - 5 ml.

### Kit Contents and Storage

CfPure cfDNA Extraction Kit , 100ml (K5011610)

Item	Amount	Storage
cfPure Lysis/Binding Buffer	1 x 115 ml	
cfPure Wash Buffer	2 x 55 ml	
cfPure Elution Buffer	1 x 6 ml	Room
cfPure Magnetic Bead	2 x 1.33 ml	Temp.
Solution	2 X 1.33 IIII	

### CfPure cfDNA Extraction Kit, 250ml (K5011625)

Item	Amount	Storage
cfPure Lysis/Binding Buffer	3 x 95 ml	
cfPure Wash Buffer	5 x 55 ml	
cfPure Elution Buffer	1 x 15 ml	Room
cfPure Magnetic Bead Solution	5 x 1.33 ml	Temp.

### Equipment and Reagents to be Supplied by User

ltem	Source	
Equipment		
Multi-channel micropipettors	Any	
Adjustable Micropipettors	Any	
Vortexor	Any	
Magnetic Particle Processor		
KingFisher <sup>™</sup> Flex Magnetic Particle	Thermofisher	
Processor	5400630	
Magnetic Head		
24 Deep-Well Plates for KingFisher <sup>™</sup>	Thermofisher	
Flex Magnetic Particle Processor	24074440	
Deep-Well Plates		
KingFisher <sup>™</sup> Flex 24 deep well plate,	Thermofisher	
sterile	95040490	
Tip Combs		
King Fisher Flex 24 Deep Well Tip	Thermofisher	
Comb and Plate	97002610	

Item	Source
Consumables	
Aerosol-resistant pipette tips	Any
Nonstick, RNase-free Microfuge tubes (1.5ml)	Any
MicroAmp <sup>™</sup> Clear Adhesive Film	Any
Reagent Reservoirs	Any
Reagents	
Ethanol, 200 proof (Absolute)	Any
SDS, 20% Solution (Only required for Proteinase K treament)	Any
Proteinase K solution (20mg/ml) (Only required for Proteinase K treament)	BioChain Z5050002

### Download KingFisher<sup>™</sup> Flex Program

On cfPure Webpage scroll down to Manual Section.
 Click cfPure\_4-5ml\_Flex and/or cfPure\_1-2ml\_Flex to download program to your computer
 Refer to KingFisher<sup>™</sup> Flex manual for instructions for installing program on the instrument Important Notes

<u>Starting Material:</u> Both fresh and frozen plasma can be used with the Cell-Free DNA isolation protocol. Fresh plasma, however, tends to have higher yields. <u>Streck Cell-Free DNA BCT Tube(s)</u>: Plasma from blood collected with Streck Cell-Free DNA BCT Tube(s) must go through a Proteinase K treatment prior to Cell-Free DNA isolation to ensure optimal yields. Forgoing Proteinase K treatment may decrease yields by 50%.



#### Protocol

### Prior to Initial Use

The Lysis/Binding and Wash Buffer are shipped as a concentrate. If precipitate is present in either solution, incubate solution at 37°C for 30 minutes. 100% EtOH must be added to both solutions before the first use. Once EtOH is added, these buffers are stable for one year if stored at 4°C. Be sure to close the bottle tightly for long term storage.

### For 100 ml kit (K5011610)

- Add 23 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 51 ml of fresh 100% ethanol to each bottle of Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction For 250 ml kit (K5011625)
- Add 19 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 51 ml of fresh 100% ethanol to each bottle of Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction

### **Proteinase K Treatment**

If samples were collected using a **Streck Cell-Free DNA BCT tube(s)**, Proteinase K treatment is required to ensure optimal yields. If blood was not collected with **Streck Cell-Free DNA BCT tube(s)** proceed to the Lysis/Binding step.

Plasma	Proteinase K	20% SDS Solution
x (x=µl of plasma)	0.015 x	0.050 x
1 ml	15	50 µl
2 ml	30 µl	100 μl
4 ml	60 μl	200 µl
5 ml	75 μl	250 μl

8. Add the appropriate amount of plasma to an appropriately sized tube(s)

- 9. Add 15 µl of Proteinase K (20 mg/ml) for every 1 ml of plasma used
- **10.** Add 50 μl of 20% SDS solution for every 1 ml of plasma used
- **11.** Mix by inverting gently 5 times
- 12. Incubate at 60°C for 20 minutes
- 13. After incubation place tube(s) on ice for 5 minutes to cool tube(s) to room temperature



### Plate Set up for 1 or 2 ml samples

Set up 24 Deep Well Plates by adding appropriate reagents according to table below

Plate ID	Diata Turna	Plate Position	Boogont	Volume p	er well
Plate ID	Plate Type	Plate Type Plate Position Reagent		1 ml	2 ml
Lucic (Dinding Diato	24 DW Plate	1	cfPure Lysis/Binding Buffer	1.25 ml	2.5 ml
Lysis/Binding Plate	24 DW Plate	1	cfPure Magnetic Bead Solution	25 μl*	50 µl*
Wash Plate 1	24 DW Plate	2	cfPure Wash Buffer	1 ml	
Wash Plate 2	24 DW Plate	3	cfPure Wash Buffer	1 ml	
Wash Plate 3	24 DW Plate	4	80% Ethanol	2 ml	
Wash Plate 4	24 DW Plate	5	80% Ethanol	1 ml	
Elution Plate	24 DW Plate	6	cfPure Elution Buffer	50 - 100 μl	
Tip Comb	24 DW Plate	7	Place a 24 Deep-Well Tip Comb in Plate		

\* **Important:** for sensitive PCR applications where inhibition is a concern, use 8  $\mu$ l or 16  $\mu$ l of beads for 1 ml or 2 ml of plasma, respectively.

### Plate Set up for 4 or 5 ml samples

Set up 24 Deep Well Plates by adding appropriate reagents according to table below

Plate ID	Plate Ture Plate Pecitic	Plate Position	Boagont	Volume per well			
Plate ID	Plate Type	Plate Position	Reagent	4 ml	5 ml		
Lysis/Binding Plate 1				1	cfPure Lysis/Binding Buffer	2.5 ml	3.125 ml
	24 DW Plate 1	cfPure Magnetic Bead Solution	50 μl*	62.5 μl*			
Lysis/Binding Plate 2	24 DW Plate 2	cfPure Lysis/Binding Buffer	2.5 ml	3.125 ml			
		24 DW Plate	cfPure Magnetic Bead Solution	50 μl*	62.5 μl*		
Wash Plate 1	24 DW Plate	3	cfPure Wash Buffer	1 ml			
Wash Plate 2	24 DW Plate	4	cfPure Wash Buffer	1 ml			
Wash Plate 3	24 DW Plate	5	80% Ethanol	2 ml			
Wash Plate 4	24 DW Plate	6	80% Ethanol	1 ml			
Elution Plate	24 DW Plate	7	cfPure Elution Buffer	50 - 100 μl			
Tip Comb	24 DW Plate	8	Place a 24 Deep-Well Tip Comb in Plate				

\* **Important:** for sensitive PCR applications where inhibition is a concern, use 16  $\mu$ l or 20  $\mu$ l of beads in each Lysis/Binding plate above for 4 ml or 5 ml of plasma, respectively.



- Gently shake Lysis/Binding Plate(s) to mix the reagents
- If extracting cfDNA from a 2 ml sample add enitre sample to a well on Lysis/Binding Plate
- If extracting cfDNA from a 4 or 5 ml sample add half of sample to a well on Lysis/Binding Plate 1 and the other half of sample to the same well on Lysis/Binding Plate 2

### Instrument Set up

- Place 24 Deep-Well magnetic head on to machine according the manuals protocol
- Select cfPure\_4-5ml\_Flex on the instrument for 4 or 5 ml extraction or cfPure\_1-2ml\_Flex for 1 or 2 ml extractions
- Start the run and follow on screen prompts to load processing plates in their respective positions

• At the end of the run remove elution plate from machine and cover plate or transfer eluate to new tubes Isolated cfDNA is ready for immediate use or can be stored at -20°C



# cfPure<sup>™</sup> Cell-Free DNA Extraction Kit

Isolation of cfDNA using KingFisher<sup>™</sup> Flex Magnetic Processor 96DW

### Catalog Number: K5011610, K5011625

### **Product Description**

Biochain's new cfPure Cell-Free DNA Extraction Kit has been designed to isolate circulating DNA from human plasma and serum. The cfPure utilizes Silica coated magnetic bead technology allowing for scalability and easy automation. The cfPure Cell-Free DNA extraction Kit can be used to isolate cfDNA from up to 96 samples of 600 µl of plasma or serum using the KingFisher<sup>™</sup> Flex Magnetic Processor with 96 Deep Well Head. This guide describes the use of the cfPure kit with the KingFisher<sup>™</sup> Flex Magnetic Processor 96DW to process samples of 1000 µl or less.

### **Kit Contents and Storage**

CfPure cfDNA Extraction Kit , 100ml (K5011610)

		,
Item	Amount	Storage
cfPure Lysis/Binding Buffer	1 x 115 ml	
cfPure Wash Buffer	2 x 55 ml	
cfPure Elution Buffer	1 x 6 ml	Room
cfPure Magnetic Bead	2 x 1.33 ml	Temp.
Solution	2 X 1.33 mi	

### CfPure cfDNA Extraction Kit , 250ml (K5011625)

Item	Amount	Storage
cfPure Lysis/Binding Buffer	3 x 95 ml	
cfPure Wash Buffer	5 x 55 ml	
cfPure Elution Buffer	1 x 15 ml	Room
cfPure Magnetic Bead Solution	5 x 1.33 ml	Temp.

### Equipment and Reagents to be Supplied by User

ltem	Source				
Equipment					
Multi-channel micropipettors	Any				
Adjustable Micropipettors	Any				
Vortexor	Any				
Magnetic Particle Processor					
KingFisher <sup>™</sup> Flex Magnetic Particle	Thermofisher				
Processor 96DW	5400630				
Deep-Well Plates					
96 Deep-Well Plates for KingFisher <sup>™</sup>	Thermofisher				
Flex Magnetic Particle Processor	95040460				
Standard Plates					
96 Standard Plates for KingFisher <sup>™</sup>	Thermofisher				
Flex Magnetic Particle Processor	97002540				
Tip Combs					
96 Deep-Well Tip Combs for	Thermofisher				
KingFisher <sup>™</sup> Flex Magnetic Particle	97002534				

Item	Source
Consumables	
Aerosol-resistant pipette tips	Any
Nonstick, RNase-free Microfuge tubes (1.5ml)	Any
MicroAmp <sup>™</sup> Clear Adhesive Film	Any
Reagent Reservoirs	Any
Reagents	
Ethanol, 200 proof (Absolute)	Any
SDS, 20% Solution (Only required for Proteinase K treament)	Any
Proteinase K solution (20mg/ml) (Only required for Proteinase K treament)	BioChain Z5050002

### Download KingFisher<sup>™</sup> Flex Program

1.On cfPure Webpage scroll down to Manual Section.2. Click cfPure\_600ul\_Flex to download program to your computer

**3.** Refer to KingFisher<sup>™</sup> Flex manual for instructions for installing program on the instrument

### Important Notes

Starting Material: Both fresh and frozen plasma can be used with the Cell-Free DNA isolation protocol. Fresh plasma, however, tends to have higher yields. Streck Cell-Free DNA BCT Tube(s): Plasma from blood collected with Streck Cell-Free DNA BCT Tube(s) must go through a Proteinase K treatment prior to Cell-Free DNA isolation to ensure optimal yields. Forgoing Proteinase K treatment may decrease yields by 50%



### Protocol

### Prior to Initial Use

The Lysis/Binding and Wash Buffer are shipped as a concentrate. If precipitate is present in either solution, incubate solution at 37°C for 30 minutes. 100% EtOH must be added to both solutions before the first use. Once EtOH is added, these buffers are stable for one year if stored at 4°C. Be sure to close the bottle tightly for long term storage.

### For 100 ml kit (K5011610)

- Add 23 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 51 ml of fresh 100% ethanol to each bottle of Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction For 250 ml kit (K5011625)
- Add 19 ml of fresh 100% ethanol to each bottle of Lysis/Binding Buffer and mix by inverting gently
- Add 51 ml of fresh 100% ethanol to each bottle of Wash Buffer and mix by inverting gently
- Prepare fresh 80% ethanol solution prior to each extraction

### **Proteinase K Treatment**

If samples were collected using a **Streck Cell-Free DNA BCT tube(s)**, Proteinase K treatment is required to ensure optimal yields. If blood was not collected with **Streck Cell-Free DNA BCT tube(s)** proceed to the Lysis/Binding step.

Plasma	Proteinase K	20% SDS Solution
x (x=µl of plasma)	0.015 x	0.050 x
600 μl	9.0 μl	30 µl
1000 μl	15 μl	50 µl

- 14. Add the appropriate amount of plasma to an appropriately sized tube(s)
- 15. Add 15  $\mu$ l of Proteinase K (20 mg/ml) for every 1 ml of plasma used
- 16. Add 50  $\mu l$  of 20% SDS solution for every 1 ml of plasma used
- **17.** Mix by inverting gently 5 times
- 18. Incubate at 60°C for 20 minutes
- **19.** After incubation place tube(s) on ice for 5 minutes to cool tube(s) to room temperature



### Plate Set up

Set up 96 Plates by adding appropriate reagents according to table below

	_	Plate		Volume	e per well
, , , , , , , , , , , , , , , , , , ,	Plate Type	Position on Instrument	Reagent	600 ul	1000 ul
Lysis/Binding	96 Deep-Well	1	cfPure Lysis/Binding Buffer	375 μl	625 μl
Plate 1	Plate	I	cfPure Magnetic Bead Solution	7.5 μl*	12.5 μl*
Lysis/Binding	96 Deep-Well	2	cfPure Lysis/Binding Buffer	375 μl	625 μl
Plate 2 Plate	Plate		cfPure Magnetic Bead Solution	7.5 μl*	12.5 μl*
Wash Plate 1	96 Deep-Well Plate	3	cfPure Wash Buffer	1 ml	1 ml
Wash Plate 2	96 Deep-Well Plate	3	cfPure Wash Buffer	1 ml	1 ml
Wash Plate 3	96 Deep-Well Plate	4	80% Ethanol	1 ml	1 ml
Wash Plate 4	96 Deep-Well Plate	5	80% Ethanol	500 μl	500 μl
Elution Plate	96 Standard Plate	6	cfPure Elution Buffer	30 - 50 μl	30 - 50 μl
Tip Comb	96 Deep-Well Plate	7		-	-Well Tip Comb in late

\* **Important:** for sensitive PCR applications where inhibition is a concern, use 2.4  $\mu$ l or 4  $\mu$ l of beads in each Lysis/Binding plate above for 600  $\mu$ l or 1000  $\mu$ l of plasma, respectively.

- Gently shake Lysis/Binding Plate 1 and 2 to mix the reagents
- Add half of plasma sample to the same wells of Lysis/Binding Plate 1 and 2

### Instrument Set up

- Place 96 well Deep-Well magnetic head on to machine, and select cfPure\_1ml\_96\_Flex on the instrument
- Start the run and follow on screen prompts to load processing plates in their respective positions
- At the end of the run remove elution plate and cover immediately

Isolated cfDNA is ready for immediate use

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