

# Reagent Kit Selection Guide

## Reagent Kit Selection Guide (From Target Nucleic Acids)

Target	Type	Origin	Scale	Reagent Kits
DNA	Total DNA	Human, animal blood (fresh, old, dried, frozen whole blood with common anticoagulants, Buffy Coat	100-400 µl whole blood	AnaPrep Blood DNA Extraction Kit 200
			400-1000 µl whole blood	AnaPrep Blood DNA Extraction Kit 1200 *especially for the granulocytes-rich blood samples (white blood cell no. more than $2 \times 10^4$ cells/µl)
DNA	Virus	Whole Blood	100-400 µl whole blood	AnaPrep Blood DNA Extraction Kit 200
DNA/RNA	Virus	Cell culture supernant, human serum, plasma, urine, cerebrospinal fluid, and other cell-free body fluids	See Reagent Handbook	AnaPrep Viral Nucleic Acid Extraction Kit
DNA	Virus/ Bacteria	Genital tract specimen (collected by cervical brush or genital swab), cervicovagina lavage, urine specimens	See Reagent Handbook	AnaPrep HPV DNA Extraction kit for swab samples
DNA	Total DNA	<ul style="list-style-type: none"> <li>Human and animal tissue (fresh and frozen tissues), Rodent tails</li> <li>Insects (fresh and frozen tissue)</li> <li>Dried blood</li> <li>Dried Swab Material (buccal, nasal, pharyngeal, vaginal, eye swab or saliva)</li> </ul>	See Reagent Handbook	AnaPrep Tissue DNA Extraction Kit
DNA	Total DNA	FFPE (formalin fixed paraffin embedded) tissue sections	See Reagent Handbook	AnaPrep FFPE DNA Extraction Kit
DNA	Total DNA	Cell culture, plasma, serum, bone marrow, buffy coat (fresh or frozen serum/plasma, cells in adherent/suspension culture, lavage)	See Reagent Handbook	AnaPrep Cultured Cell DNA Extraction Kit
DNA	Bacteria	Bacteria species (from different kinds of starting materials), bacteria pellets, liquid transport media, swabs and urine, colony	See Reagent Handbook	AnaPrep Bacterial DNA Extraction Kit *Special item: AnaPrep TB DNA Extraction Kit
DNA	Total DNA	Forensic material (whole blood, clotted blood, bones, teeth, ancient bones, hair roots, forensic surface and contact swabs, saliva, chewing gum, cigarette butts, stamps, envelops, tissue, etc.)	See Reagent Handbook	AnaPrep Forensic DNA Extraction Kit

# Introduction

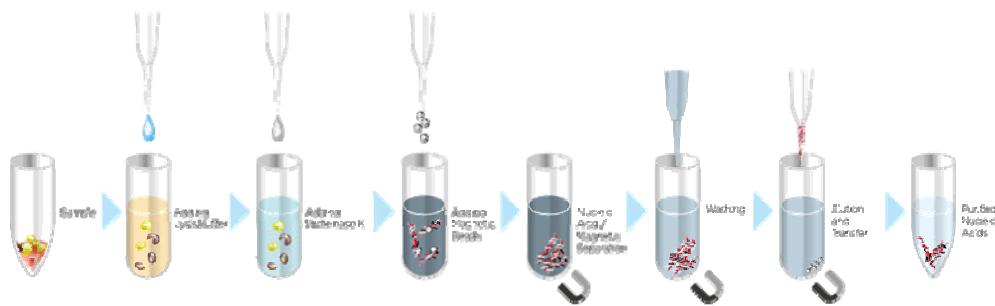
## The BioChain Nucleic Acid Preparation Technology

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### Introduction

BioChain Institute Inc. specializes in developing advanced, efficient and reliable technologies in nucleic acid preparation, to enable successful delivery of extraction results from varied sample types.

The AnaPrep technology is a state of the art platform that uses magnetic beads to extract nucleic acids from samples. The platform commits to a truly walk-away automation for nucleic acid purification from samples to results. The purification processes contain steps of lysis, binding, washing and elution (see figure below).



magnetic bead extraction process

# Product information

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## **Intended use**

AnaPrep Kits are intended to be used on the AnaPrep 12 and 24 instruments for the preparation of nucleic acids from biological specimens. The AnaPrep instruments and AnaPrep reagent kits are not intended for use as part of a specific in vitro diagnostic test.

The nucleic acids purified using the AnaPrep instruments and reagent kits are suitable for a variety of polymerase chain reaction (PCR) tests. The AnaPrep instruments and reagent kits are intended for research use only.

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## **Warranty**

BioChain is committed to providing our customers with high-quality products and services. Our goal is to ensure that every customer is 100% satisfied with our products and services. If you have questions or concerns about our products or services, contact our Technical Support Representatives.

BioChain guarantees the performance of all products according to specifications stated on our product literature. The purchaser/user must determine the suitability of the product for its particular use. We reserve the right to change, alter, or modify any product to enhance its performance and design.

This warranty limits BioChain Institute's liability only to the cost of the product. No warranty is granted for products beyond their listed expiration date. No warranty is applicable unless all product components are stored in accordance with instructions.

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**Satisfaction  
Guarantee**

For any product that fails to perform satisfactorily due to any reason other than misuse, BioChain will replace it free of charge. Simply call BioChain or your distributor to get a replacement.

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**Technical  
Support**

For technical assistance and more information, please visit our website at [www.biochain.com](http://www.biochain.com) or call BioChain's Technical Service Department or your local distributor.

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**Safety  
Information**

When working with chemicals or samples, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate material safety data sheets (MSDS). You can find, download, view, and print them from our website [www.biochain.com](http://www.biochain.com).

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**Manufacturer  
Information****Manufacturer:**

BioChain Institute Inc.

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**Tel:** 1-510-783-8588 **Fax:** 1-510-783-5386

**Mail:** [info@biochain.com](mailto:info@biochain.com)

**Country of Origin:** USA

# AnaPrep Cultured Cell DNA Extraction Kit

**Cat. No. Z1322005**

**Process Time: 45 minutes**

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## Intended Use

AnaPrep Cultured Cell DNA Extraction Kit is used with the AnaPrep 12 or 24 instruments for extraction of genomic DNA from cultured cells or buffy coat.

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## Application

Nucleic acids extracted with the Cultured Cell DNA Extraction kit can be used in a number of downstream applications including: PCR, qPCR, Sequencing (NGS), Microarray, RFLP, and Southern Blot Analysis.

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**Number Of Tests** 48 extractions

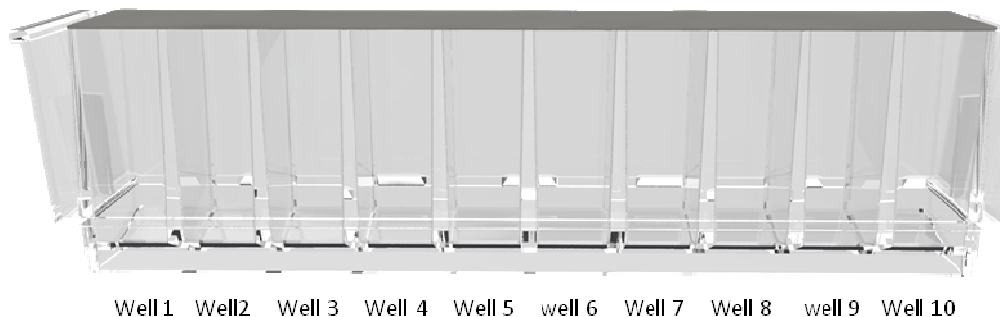
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## Kit Components

Kit Contents	Z1322005-48
Reagent Cartridge	48 pcs
Reaction Chamber	48 pcs
Tip Holder	48 pcs
Filtered Tip	50 pcs
Piercing Pin	50 pcs
Sample Tube (2 ml)	50 pcs
Elution Tube (1.5 ml)	50 pcs
Protocol Barcodes	1 pc

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## Reagent Cartridge Content



well-1	Proteinase K solution	40 $\mu$ l
well-2	Lysis Buffer 2	720 $\mu$ l
well-3	Binding Buffer 1	720 $\mu$ l
well-4	Magnetic Bead Solution	800 $\mu$ l
well-5	Washing Buffer 1	1000 $\mu$ l
well-6	Washing Buffer 2	1000 $\mu$ l
well-7	Washing Buffer 3	1000 $\mu$ l
well-8	Elution Buffer 1	1000 $\mu$ l
well-9	Elution Buffer 2	1000 $\mu$ l
well-10	Empty	

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### Storage

- ◆ AnaPrep Cultured Cell DNA Extraction Kit should be stored at room temperature (15-25 °C). Do not freeze the reagent cartridges. The Kits are stable for 12 months under the proper storage conditions.
- ◆ Store the purified DNA at 4 °C (short- term) or aliquot and store at -70°C (long-term).

## Starting Material

1. Cultured cells in suspension and monolayer
  2. Cells from Buffy coat (without red blood cells)
- ◆ If the buffy coat is removed directly from whole blood, then the AnaPrep Blood DNA Extraction Kit (Z1322001; Z1322002) is recommended for the complete removal and lysis of red blood cells.
  - ◆ Do not use more than  $4 \times 10^6$  cells with a normal set of chromosomes. However, due to differences in the sizes of cells, **we suggest that customers start with lower number of cells and slowly titrate the cell numbers if needed.**
    - a. For suspension cells, start with  $2.5 \times 10^6$  cells
    - b. For adherent cells, start with  $1 \times 10^6$  cells
  - ◆ Too many cells may clog the tips.
  - ◆ The cell number can be determined by using a Hemocytometer<sup>1,2</sup> (Petroff-Hauser Chamber ) and an automated cell counter ( e.g. TC10™, Countess®, Cellometer®, and Scepter™ automated cell counters).
  - ◆ The types and amounts of starting material for use in the AnaPrep Cultured Cell DNA purification procedures are shown in the Table listed below.
  - ◆ References:
    1. <http://www.smccd.edu/accounts/case/biol230/algae/hemocytometer1.pdf>
    2. <http://web.mnstate.edu/provost/CountingCellsHemocytometer.pdf>

<p><b>Cells grown in suspension</b></p>	<p>◆ <b>Suspension culture:</b></p> <ol style="list-style-type: none"> <li>1. Determine the number of cells. (<u>see Starting Material section for suggested cell number</u>).</li> <li>2. Centrifuge the appropriate number of cells for 5 min at 300 x g in a 1.5 ml microcentrifuge tube.</li> <li>3. Remove the supernatant completely and discard, without disturbing the cell pellet.</li> <li>4. Resuspend the cell pellet in PBS to a final volume of 200 µl.</li> </ol>
<p><b>Cells grown in a monolayer</b></p>	<p>◆ Cells grown in a monolayer can be detached from the culture flask by either trypsinization or by using a cell scraper.</p> <p>◆ <b>Using trypsin:</b></p> <ol style="list-style-type: none"> <li>1. Aspirate the medium and wash the cells with PBS.</li> <li>2. Aspirate the PBS, and add 0.10–0.25% trypsin, incubate at 37°C.</li> <li>3. After cells have detached from the dish or flask, collect them in medium, and transfer the appropriate number of cells to a 1.5 ml microcentrifuge tube.</li> <li>4. Centrifuge for 5 min at 300 x g.</li> <li>5. Remove the supernatant completely and discard, without disturbing the cell pellet.</li> <li>6. Resuspend the cell pellet in PBS. Determine the number of cells (<u>see Starting Material section</u>). Bring final volume to 200 µl.</li> </ol> <p>◆ <b>Using a cell scraper:</b></p> <ol style="list-style-type: none"> <li>1. Detach cells from the dish or flask by scraping.</li> <li>2. Harvest and transfer the cells to a 1.5 ml microcentrifuge tube and centrifuge for 5 min at</li> </ol>

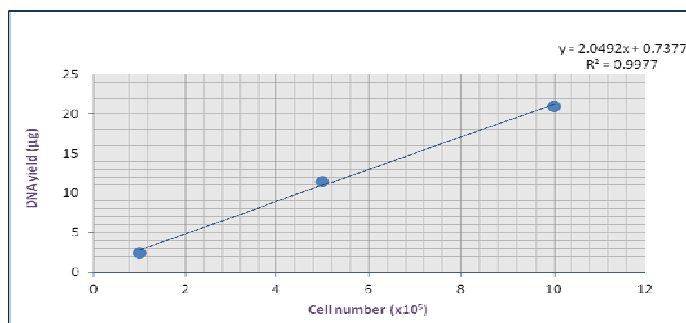


	<p>300 x g.</p> <ol style="list-style-type: none"> <li>3. Remove the supernatant completely and discard, without disturbing the cell pellet.</li> <li>4. Resuspend the cell pellet in PBS. Determine the number of cells (<u>see Starting Material section</u>). Bring final volume to 200 <math>\mu</math>l.</li> </ol>
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## Yield Of Purified DNA

DNA yields depend on the sample type, number of nucleated cells in the sample, and the protocol used for purification of DNA.

For example, the average DNA yield from the HT29 colon adenocarcinoma cell line at the different concentrations (in the range from  $1 \times 10^5$  to  $10^6$  cells) is about  $22\mu\text{g}/10^6$  cells (see below).



## Quality Control

In accordance with BioChain's ISO-certified Quality Management System, each lot of AnaPrep Cultured Cell DNA Extraction Kits is tested to ensure consistency in product quality.

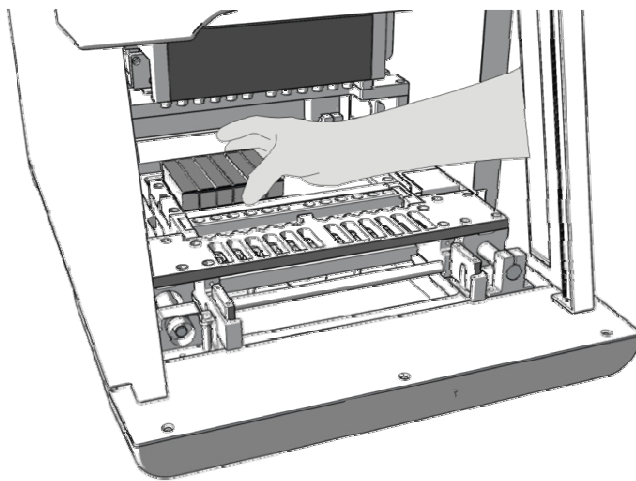
## Protocol of extraction

1. Turn the power switch on and wait for the LCD screen to light up and display “AnaPrep 12 System Stand-By”, or “AnaPrep 24 System Stand-By” (figures shown here are from AnaPrep 12 and both systems operate the same way).
2. Press the “Start” button  
(The system will process self-testing, and then go to steady mode).

**Note:**

The system will block main functions before the completion of the self-testing process.

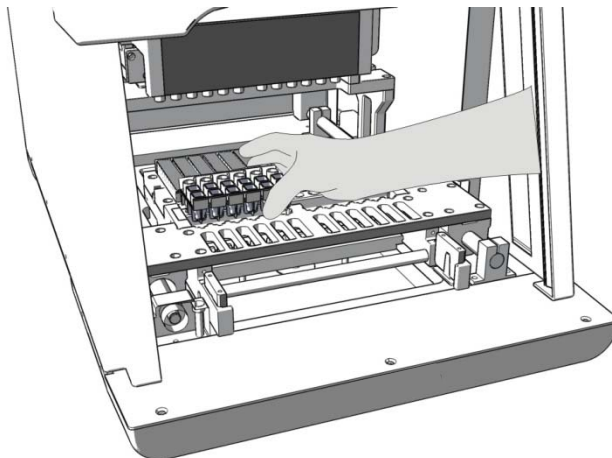
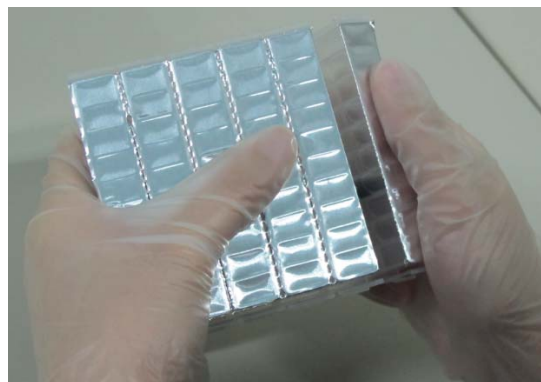
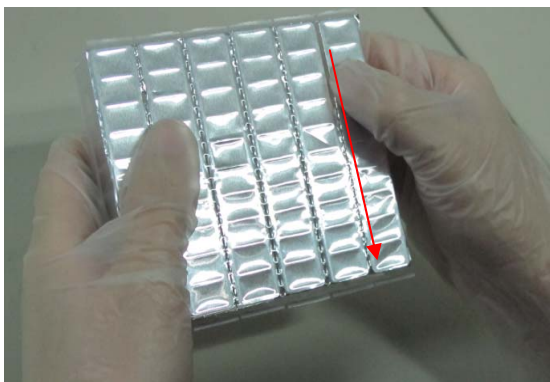
3. Open the sliding door and remove the sample rack from the instrument.
4. Load Reagent Cartridges, and all plastic disposables (Reaction Chamber, Tip Holder, Piercing Pin, and Filtered Tip)



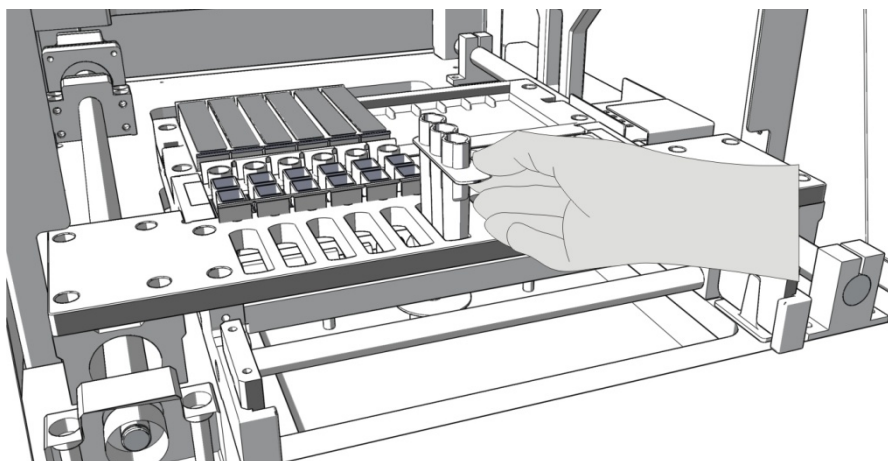
### Insert Reagent Cartridges

## ■ How to pull apart the reagent cartridges

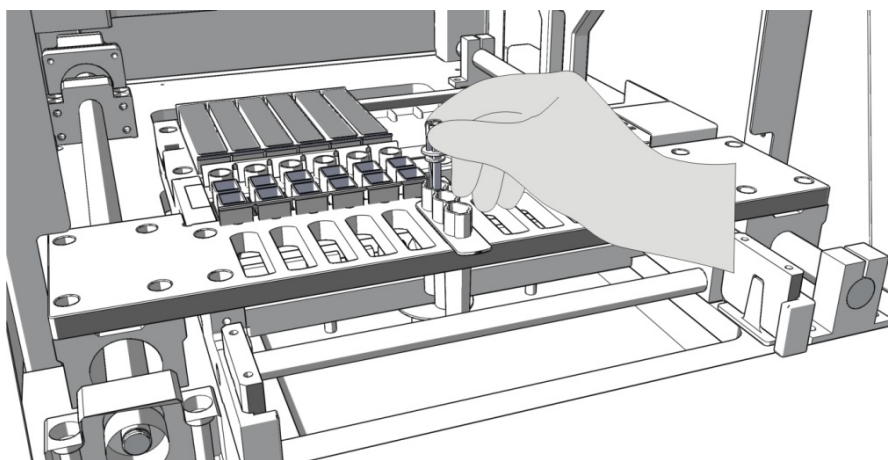
- Cut foil with a finger nail along the dotted line and then snap it apart with a little bit of force.



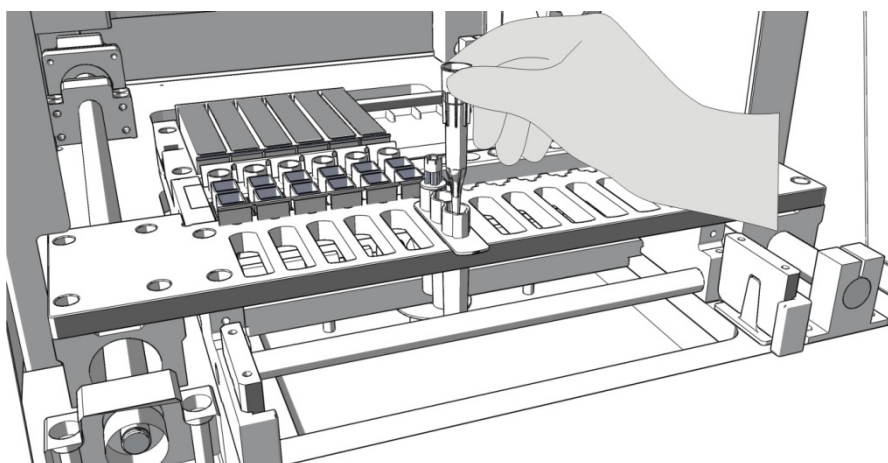
**Insert Reaction Chambers**



**Insert Tip Holder**



**Insert Piercing Pins**



**Insert Filtered Tips**

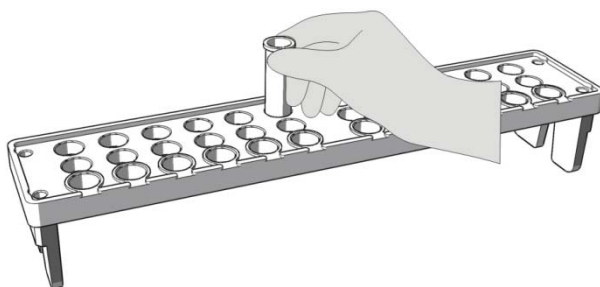
**Note:**

Load one Reagent Cartridge and one set of plastic disposables per sample.

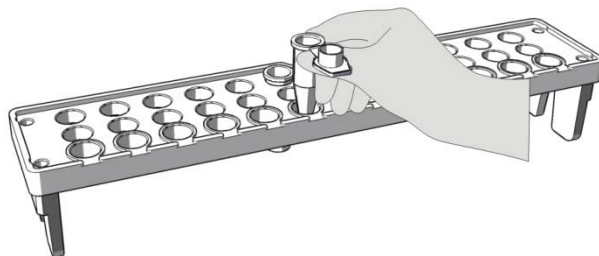
**Important:**

- Set Reagent Cartridges in the order of the number from left to right.
- Make sure that Cartridges are inserted in to the Cartridge Tray tightly.
- You can load 1-12 cartridges on the tray depending on the number of samples that you wish to process.

5. Load Sample Tube and Elute Tube to Sample Rack on the bench

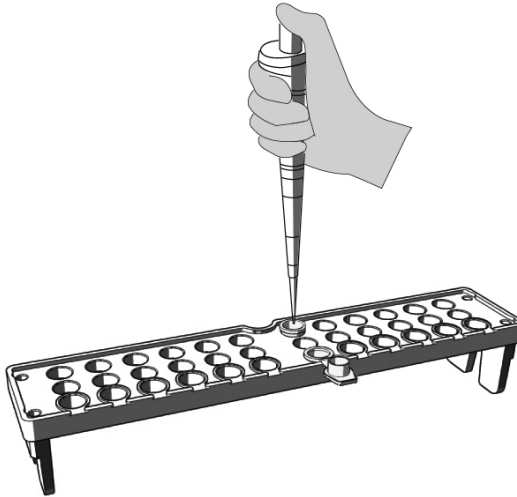


**Insert Sample Tube into the Sample Rack**



**Insert Elute Tube into the Sample Rack**

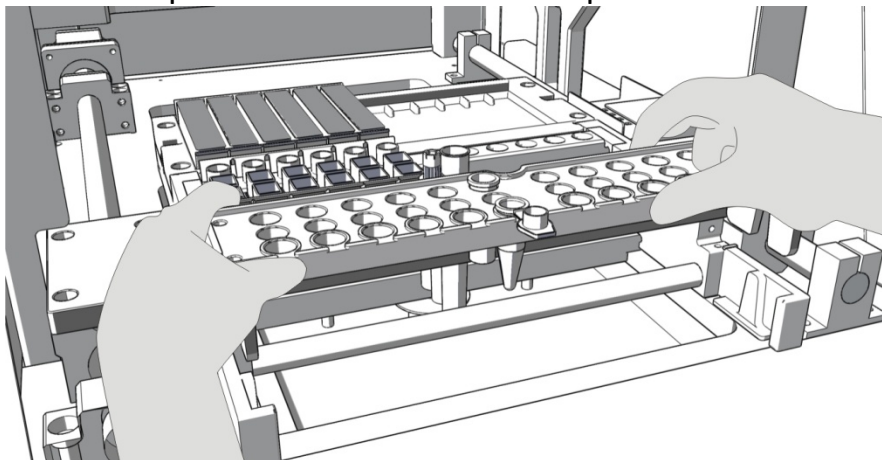
6. Load the sample(s) to Sample Tube.



**Note:**

- Pretreatments are essential for some sample types before loading to Sample Tube. Please refer to the handbook of reagent kits for details.
- Make sure the caps of Elute Tube are open as the figure shown above.

7. Place Sample Rack on the instrument platform



**Note:**

- Use two hands to handle the Sample Tray.
- Make sure the Sample Tray is placed correctly in the instrument.

8. Close the door.
9. Scan the protocol barcodes to select purification protocol, sample volume and elute volume.



**Note:**

- There is one protocol barcode paper enclosed in each reagent kit box.
  - The protocol's name, sample volume and elution volume will be shown on LCD screen after the protocol barcodes are scanned.
10. Follow the instructions displayed on the LCD screen to double check the operation steps to be completed prior to running the program.
  11. Press "Enter" to confirm. The instrument will start running the protocol program automatically and will terminate once all processes are completed.

**Note:**

- It takes 30 to 45 minutes to complete the extraction process and may vary according to reagent types.
12. At the end of the run, the instrument beeps briefly while the LCD screen displays "Protocol Completed".
  13. Open the instrument door.
  14. Remove the elute tubes containing the purified nucleic acids.  
Note: Store the purified nucleic acids at 4°C for short-term storage or store at -70°C for long-term storage.
  15. Discard the used cartridges and all plastic consumables into the biohazard waste. Do not reuse the cartridges
  16. If you're not using the instrument, place the Sample Rack back

into the AnaPrep, close the instrument door and press the “Start” button for 2 seconds to enter into “sleep mode”. If the instrument will not be used for a longer period of time turn-off the power switch.



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