

[Product Name] MagPure Universal DNA Precast Kit (96 channel machine)

**[Product Specification]** 96 Preps/Kit

#### [Intended Use]

This product is suitable for extracting high-purity DNA from various clinical samples (blood, tissue, exfoliated cells, FFPE samples, serum, plasma, blood plaques, oral swabs, saliva, and cultured bacteria). The extracted products can be used for in vitro clinical testing.

### [Principle]

This product is based on the purification method of high binding magnetic particles. The sample is lysed and digested by lysis buffer and Protease K Solution. DNA is released into the lysate. After adding magnetic particles and binding solution, DNA will be adsorbed on the surface of magnetic particles, and impurities such as proteins will be removed without adsorption. The adsorbed particles were washed with washing solution to remove proteins and impurities, washed with ethanol to remove salts, and finally DNA was eluted by Elution Buff or Nuclease Free Water.

#### [Main Composition]

Cat.No	Precast Reagent	IVD3102-F-96
RNase A		20 mg
Proteinase K		50 mg
Protease Dissolve Buffer		6 ml
Buffer ATL		30 ml
Buffer AL		30 ml
96-Tip		1
Sample Plate (DW Plate)	450µl Buffer BD (Ethanol Added)	1
Wash 1 Plate (DW Plate)	600µl Buffer BW1 (Ethanol Added)	1
Wash 2 Plate (DW Plate)	600µl Buffer BW1 (Ethanol Added)	1
Wash 3 Plate (DW Plate)	750µl Buffer GW2, 20µl MagPure Particle	1
Elute Plate (DW Plate)	80µl Elution Buffer	1

# 【Storage conditions and validity】

This kit is shipped and stored at room temperature and is valid for 18 months.

### [Applicable Instrument]

96 channel Nucleic Acid Extraction Machine such as Kingfisher Flex, MagMix 96 (Magen), Auto Pure 96 (Allsheng), EXM6000 (Zybio) or similar.

### [Preparation before Use]

- Add 2.5ml Protease Dissolve Buffer into the bottle of Proteinase K, mix well and stored at 20~8℃.
- Add 1.4ml Protease Dissolve Buffer into the bottle of RNase A, mix well and stored at 20~8℃.

#### [Part 1: Sample Preparation]

### A. Liquid samples (such as blood, serum, plasma, buffy coat, cell suspension, etc.)

Add  $20\mu$ l Proteinase K, (optional)  $5\mu$ l RNase A and  $200\mu$ l samples (such as blood, yellow layer, plasma, serum, cell suspension) to a 1.5ml centrifuge tube. Add  $220\mu$ l Buffer AL, vortex for 15 seconds, incubate with shaking at  $70^{\circ}$ C for 10 minutes. Follow the Part 2 operation.

#### B. Dry blood spot or Seminal Stain

Transfer  $3\sim5$  slices of 3mm diameter blood spots into a 2.0ml centrifuge tube. Add  $300\mu$  Buffer ATL and  $20\mu$ l Proteinase K to the sample, incubate at  $55^{\circ}$ C with shaking ( $900\sim1200$ rpm) for 60 minutes. For seminal stain samples, add  $10\mu$ l DTT (1M) to the lysate. Then add  $150\mu$ l Buffer AL, incubate at  $70^{\circ}$ C with shaking ( $900\sim1200$ rpm) for 15 minutes. Centrifuge at  $13,000\times g$  for 2 minutes. Follow the Part 2 operation.

# C. Dry swab

Transfer swab into a 2.0ml centrifuge tube. Add  $20\mu$ l Proteinase K and  $500\mu$ l Buffer ATL, incubate at  $55^{\circ}$ C with shaking (900~1200rpm) for 30 minutes. Centrifuge at 13,000 x g for 2 minutes. Follow the Part 2 operation.

# D. Wet swab (including cell preservation solution)

Centrifuge at  $10,000 \times g$  for 1 minute to collect exfoliated cells, discard excess storage solution, and leave the remaining  $300\mu l$  preservation solution and swab. Add  $100\mu l$  Buffer ATL, (Optional)  $5\mu l$  RNase A and  $20\mu l$  Proteinase K, incubate at  $55^{\circ}$ C with shaking ( $900\sim1200$ rpm) for 30 minutes. Follow the Part 2 operation.

# E. Saliva sample (including preservation solution)

Transfer  $450\mu l$  swab into a 2.0m l centrifuge tube. Add  $20\mu l$  Proteinase K and (Optional)  $5\mu l$  RNase A, incubate with shaking at  $55\sim65^{\circ}C$  for  $30\sim90$  minutes. Follow the Part 2 operation.

### F. Tissue sample (<20mg tissue sample)

Transfer <20mg tissue to a 1.5ml centrifuge tube. Add  $20\mu l$  Proteinase K and  $200\mu l$  Buffer ATL, incubate with shaking at  $55^{\circ}$ C for  $30^{\circ}120$  minutes or until the sample is completely digested. Add  $5\mu l$  RNase A, mix well and let sit for 10 minutes. Add  $200\mu l$  Buffer AL, incubate with shaking at  $70^{\circ}$ C for 10 minutes. Centrifuge at  $13,000 \times g$  for 2 minutes. Follow the Part 2 operation.

### G. Cultured cells (< 5x10<sup>6</sup> cells), exfoliated cells

Take an appropriate amount of liquid samples such as culture medium, urine, amniotic fluid or ascites into a centrifuge tube, centrifuge at  $2,000 \times g$  for 10 minutes to collect cells or exfoliated cells. Remove the culture medium and leave the remaining  $100\mu$ l culture medium or body fluid, vortex to resuspend cells. Add  $100\mu$ l Buffer ATL, (Optional)  $5\mu$ l RNase A and  $20\mu$ l proteinase K, incubate at  $55^{\circ}$ C with shaking (900-1200rpm) for 15-30 minutes. Add  $200\mu$ l Buffer AL, vortex for 15 seconds. Follow the Part 2 operation.

### H: FFPE sample

Transfer paraffin embedded tissue slices into a 1.5ml centrifuge tube and remove the paraffin with xylene or a substitute (such as dewaxing solution DPS). Add  $20\mu\text{l}$  Proteinase K and  $220\mu\text{l}$  Buffer ATL, mix well and incubate at  $55^{\circ}\text{C}$  for 60-90 minutes. Incubate at  $90^{\circ}\text{C}$  for 60 minutes. Add  $5\mu\text{l}$  RNase A, mix well and let sit for 10 minutes. Add  $220\mu\text{l}$  Buffer AL, vortex for 15 seconds. Centrifuge at  $10,000 \times \text{g}$  for 3 min to remove the undigested particles Follow the Part 2 operation

#### 【Part 2: Auto Pure 96 nucleic acid extractor operation】

- 1. Take out the required components of the kit.
- 2. Inverting the Wash 3 Plate several times to re-suspend the magnetic beads.
- 3. Remove the sealing bag and sealing film.
- 4. Place a 96 tip comb for deep well magnets on Wash 3 Plate.
- 5. Take out the sample plate, add  $400\sim450\mu l$  digestion solution to each hole
- 6. Turn on the machine and start the IVD3102-F-96 protocol
- 7. After the extraction complete, ~30 minutes.
- 8. Remove the 96 well plate and store the purified DNA at -20 $\sim$ 8  $^{\circ}$ C

#### [Basic Information]

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