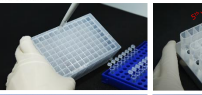
■ End of Run







Once extraction is completed, take tube rack out of the equipment and collect Genomic DNA from 5th (11th) well depending on where samples were dispensed

Transfer eluent into 1.5 ml or PCR tube.

Amount of eluted buffer will be around 80 µl and it is ready to use.

Regular UV sterilization eliminates bacteria and virus, nucleic acid, hence it minimizes internal pollution within the system.

Before/after the extraction, touch UV Lamp button. The sterilization process will continue for the time vou set.

Troubleshooting Guide

Problem	Causes	Comments and Suggestions				
	Upside down during transportation may cause beads to stick with sealing film.	Spin down by hand the plate or strip by hand before open it.				
Physical damage of the kit	Sealing film is detached and reagent is spilled to other wells due to improper storage temperature.	Spin down by hand and measure reagent volume with eyes. If reagent volumes are insufficient, extraction efficiency may decrease. Do not use it and contact customer service immediately.				
Magnetic rod function failure	Stain on the magnetic rod	Ensure the magnetic rod covers are inserted properly before extraction. Clean magnetic rods using 70% concentration of ethanol and clean with cloth.				
	System is not working	Make sure system is plugged. Refer to user manual of GENTI™ Advanced for further details.				
Extractor malfunction	Liquid spilled and adhered to system	Use UV light for sterilization and then clean with 70% concentration of ethanol.				
	Collision	Improperly attached plate or strip may cause collision (between plate & strip, plate & system component and strip & system component). Turn off the device and make sure plate and strip are properly attached.				

Storage Conditions

· Temperature : Room Temperature (15~25°C)

· Humidity : 20~80%



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Symbol Used for Symbol Used for LOT Batch number Manufacturer REF Catalogue number Do not reuse Consult Instructions For Date of w Use Manufacture Caution CE-Mark Representative in EC REP Temperature limitation EC In-vitro diagnostic IVD Expiry date medical device

2023.08

Ver. 1.2





GeneAll

GENTi TM 32 ADVANCED

Genomic DNA Extraction Kit (Single Tube Type/Plate Type)

Description

GENTi™ Advanced Genomic DNA Extraction Kit utilize magnetic bead-based equipment, enables highly efficient nucleic acid extraction from a wide range of samples.

Maximum 32 and 16 samples (901-048A/901-096A) are performed in deep well conical shaped bottom plate (tube), which enables precise fit with heating block and deliver the highest possible recovery of nucleic acids.

The purified Genomic DNA is of excellent quality and can be directly compatible with sensitive downstream detection methods such as PCR, qPCR and other molecular diagnostic testing.

Kit Contents

Components	Quantity				
Components	901-096A	901-048A			
Number of Preparation	96 tests/kit	48 tests/kit			
Pre-filled with reagents	6 plate	48 Tube			
Magnetic rod cover (6 pcs/pk)	2 pk	4 pk			
Proteinase K 24 mg/ml *	2 tubes	1 tubes			
PK storage buffer 1.5 ml	2 tubes	1 tubes			
RNase A (20 mg/ml) 500 μl	2 tubes	1 tubes			

* Reconstitute the Proteinase K by adding 1.2 ml of PK storage buffer (provided) before use.

Brief Workflow



- 3. Washing 4. Washing + Magnetic beads
- 6. Washing

- 96 Deep-well plate pre-filled with reagents





- 8 Strip deep-well tube pre-filled with reagents



- Disposable magnetic rod cover



- GENTi™ Advanced instrument heat block





- · Conical shape of the plate, magnetic rod cover and heating block
- · Heating block combined with the close fit of conical shaped material for fast and efficient heat transfer
- 6 consequent wells are aligned horizontally and each well contains specific reagents for extraction.
- 1st (7th) well contains lysis buffer which destroy cell membranes and bind Target DNA/RNA with magnetic beads.
- Magnetic bead is located at 4th (10th) well until the extraction begins and moves to first well by device once initiated.
- 2nd, 3rd, 4th, 6th (8th, 9th, 10th, 12th) well contain washing buffer in order to remove unwanted cell component and buffers.
- Elution buffer in 5th (11th) well detaches Target DNA/RNA from magnetic bead and complete extraction process.
- If particles are not visible in well 4th (10th) shake down the cartridge to dislodge particles that may have adhered to the seal material before removing the seal.

Protocol

Protocol	Ex) Condition
Fast 17′ 46″	Time-saving, high-speed extraction for diagnostics PCR-ready nucleic acid Double check the accuracy of sample confirmed positive
Normal 36′	Standard procedure of nucleic acid extraction Optimized for nucleic acid extraction from a variety of clinical sample
High 41' 10"	High quality nucleic acid extraction (High yield & purity) Accommodate complex clinical samples. ex) tissue, NGS-grade sample, etc

Protocol Normal

Step	Step 1	Step 2	Step 3	Step 4	Step 5	Step 6	Step 7	Step 8	Step 9	Step 10			
Well	4	1	1	2	3	4	6	6	5	4			
Name	Bead TF	Lysis	Lysis	Wash 1	Wash 2	Wash 3	Wash 4	Dry	Elution	Reclaim			
Waiting	00:00	00:00	00:00	00:00	00:00	00:00	00:00	01:00	00:00	00:00			
Mixing	00:00	05:00	07:00	02:00	02:00	03:00	03:00	00:00	05:00	00:20			
Magnet	00:20 x 2	00:00	00:30 x 3	00:20 x 2	00:20 x 2	00:40	00:40	00:00	00:15 x 5	00:00	Heat Block	Lysis	Elution
Volume	700	1000	1000	600	600	600	600	100	100	50	Block Tm	60°C	85°C
Speed	Slow	Fast	Fast	Fast	Fast	Fast	Fast	Slow	Fast	Fast	Start step	-	Step 3
Collet	Cycle	Cycle	Cycle	Cycle	Strong	Strong	Strong	Strong	Cycle	Strong	Stop step	Step 3	-

equipment Run













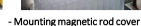
- 1. Turn on the **power switch** located on the right rear of the equipment.
- 2. Touch 'RUN' button when the home screen panels appear.
- 3. Select 'Self Test' at the File Browser screen and then touch 'RUN' button to run a self-test.
- 4. After self test completes, select 'Normal' protocol and then touch 'RUN' button for the operation. (Optimal protocol of the three options, 1)Fast <17min>, 2) Normal <36min>, 3) High <41min>)
- * Note: If 'Warning' screen appears, check system and touch 'RUN' button again.

Precautions for Use of equipment









When installing the magnetic rod

equipment.

cover, push it to the end of

- Mounting Tube rack

Automatic start self-test when Be careful when mounting the tube rack on the equipment the instrument is turned on and gently mount it.

Preparation of 8 Strip deep-well tube



(8 Strip deep-well tube) Cut the tube as much as necessary and install it in the GENTi™ Advanced tube rack.

■ Preparation of 96 Deep-well Plate









- * Protocol (Liquid sample)
- 1. Peel back the seal of pre-filled with reagents plate. (Tube)
- 2. Dispense 20 µl of dissolved Proteinase K to 1st (7th) well.
- * Note: To obtain a working solution of 20 mg/ml, add 1.2 ml of PK storage buffer to the tube containing 20 mg/ml of Proteinase K. Dissolve the Proteinase K thoroughly, divide it into conveniently sized aliquots, and store at -20°C, Do not freeze-thaw the aliquots of Proteinase K more than 3 times.
- 3. Dispense 10 µl of RNase A to 3rd (9th) well.
- 4. Dispense 200 μl of samples to 1st (7th) well.
- 5. Load plate on the tray of GENTi™ Advanced equipment system.
- * Note: Make it sure that diagonally cut edge of tube rack faces the top left of the heating block and check if the tube rack is placed evenly.
- 6. Insert magnetic rod cover to the end to strip bracket.
- * Note : Ensure that magnetic rod cover is in the correct position.

Proteinase K (20 ml/ml) at first use

This kit is provided with Proteinase K which is provided in freeze-dried format. Thus, it should be reconstituted thoroughly with PK storage buffer. To obtain a solution of 20 mg/ml of Proteinase K, add appropriate amount of the PK storage buffer. Reconstituted enzyme should be stored at 4°C for its stability. But for long-term strorage, storage at -20°C is recommended.