



TANBead® Nucleic Acid Extraction Kit

Rice DNA Auto Plate

(For use with the SLA-16 / 32 / E13200 series)

RUO

619A56

(For Research Use Only) V3

1. Intended Use

This product is designed for isolating nucleic acid from various samples, which can be performed by using TANBead® Nucleic Acid Extractor and is intended for research use only.

2. Purpose

TANBead® Nucleic Acid Extraction Kit (619A56) with TANBead® Nucleic Acid Extractor (SLA-16 / 32, SLA-E13200 Series) simplifies the process of isolating rice DNA from grinding grain and cell lysis to DNA purification completely by automated system. In addition, because TANBead® has its own nano-beads for nucleic acid binding with high efficiency, it takes only one single grain of rice for nucleic acid extraction and is ideal for applications such as germplasm identification and laboratories of academic research. TANBead® Nucleic Acid Extractor can handle 32 rice samples simultaneously and the automated process can save labor and time by enhancing the extraction efficiency, consistency and reproducibility.

3. The basic principle

The silicon dioxide layer coated on the magnetic beads can adsorb the negatively charged molecules to purify nucleic acids from samples.

4. Specification

Starting Materials	One single grain of rice
Elution Volume	90~130 µL

5. Component Supplied with the Kit



Auto Plate	60	Auto Plate with reagent buffers
Proteinase K	1.0 mL x 20	Proteinase K
Elution Buffer	50 mL	Nuclease-Free Water
Lysis Buffer 1	100 mL x 5	Tris buffer, surfactant, pH 8.0
Lysis Buffer 2	100 mL x 16	Guanidine salt, Tris buffer, surfactant
Strip	120	8-channel strip
Protocol	1	Instruction guide for user

6. Auto Plate Content

Well	Buffer	Volume (µL)
1 / 7	Elution Buffer	130
2 / 8	Washing Buffer 3	800
3 / 9	Magnetic Beads	800
4 / 10	Washing Buffer 1	800
5 / 11	-	-
6 / 12	-	-

7. Kit Storage and Shelf Life

- Components under room temperature (15~35°C) can be stored until the expiration date labeled on the box.
- The proteinase K is transported at room temperature. Upon received, please store proteinase K at 2~8°C.

8. Precautions

- For research use only.
- Avoid using expired reagents.
- When the temperature is below 20°C, place the Auto Plates in an oven (preheated 42~60°C) 5 to 10 minutes.
- Avoid vigorous shaking, in order to avoid excessive formation of foam.
- Carefully remove aluminum foil to avoid splashing.
- Do not expose the opened reagents or Auto Plates to air. The evaporation would lead to pH change, or effect on the extraction effectiveness.
- Please check the integrity of the Auto Plates and remember to insert the strips into the appropriate position of the suitable instrument before operating them.
- Please wear a mask and disposable gloves when handling.
- Use sterile consumables to avoid nuclease contamination.

- Reagent solution contains guanidine salt, avoid using bleach containing detergent.
- Avoid eyes, skin, and clothing contact with reagents. In case of any contact, flush with flowing water.
- If any serious incident occurs, please report to the manufacturer and the competent authority of the member state in which the user and / or the patient is established.

9. Materials required, Not Supplied

- TANBead® Nucleic Acid Extraction System
Model: SLA-16 / 32 / E13200 series (non-sterile)
- Disposable gloves
- Scissors, utility knives
- Micropipette, disposable tips (10 µL / 200 µL / 1000 µL)
- 1.5 mL microcentrifuge tube
- 15 mL / 50 mL conical tube

10. Nucleic Acids Extraction Protocol

Before operating, turn on the warm-up system of TANBead® Nucleic Acid Extractor, if it is equipped with temperature controller, please setting at 70°C.

- Carefully remove the aluminum foil on the Auto Plates.
- Use micropipette to load 400 µL Lysis Buffer 1 and 20 µL Proteinase K into column #6 / #12 of Auto Plate.
- Put one naked rice seed into column #6 / #12 of Auto Plate.
- Push Auto Plate combined with conducting plate which is attached to column #6 / #12 completely to the bottom of plate rack. Make sure that chamfer of the plate is at the lower left.
- Push strips completely to the bottom of strip rack frame.
- Close the door panel.
- Select the program "DNA-RICE-AUTO". The parameters are given in following section.
- The program will stop temporarily and the buzzer alarm after one hour.
- If Nucleic Acid Extractor is equipped with temp. controller, please set at 45°C.
- Carefully pull out Auto Plate and pay attention to high temperature of conducting plate.
- Use micropipette to transfer half of the lysate from column #6 / #12 to #5 / #11 and load 750 µL Lysis Buffer 2 into #5 / #6 and #11 / #12.
- Push Auto Plate completely to the bottom of plate rack, make sure that chamfer of the plate is at the lower left and press "Pause" to go on program.
- Once the program has ended, buzzer shall alarm. Take out Auto Plate carefully
- Use micropipette to transfer the purified nucleic acid from column #1 / #7 to a clean tube.
- Put the used Auto Plate and strips into the waste recovery can.

11. Program

■ SLA-16 / 32 series

Program Name: DNA-RICE-AUTO					Model: SLA-16 / 32 series			
Step	Well	Mixing (M)	Collect (S)	Rod	Mixing speed	Volume (μL)	Pause	Vapor (M)
1	6	20	0	Off	Medium	600	Off	0
2~26	6	1	0	Off	Medium	600	Off	0
27	6	10	0	Off	Medium	600	Off	0
28	6	5	0	Off	Medium	600	On	0
29	6	5	0	Off	Medium	900	Off	0
30	5	5	0	Off	Medium	900	Off	0
31	3	1	60	On	Medium	800	Off	0
32	4	1	60	On	Medium	800	Off	0
33	6	5	100	On	Medium	900	Off	0
34	5	5	100	On	Medium	900	Off	0
35	4	3	100	On	Medium	800	Off	0
36	3	2	60	On	Medium	800	Off	0
37	2	2	60	On	Medium	800	Off	10
38	1	5	100	On	Medium	150	Off	0
39	2	1	0	Off	Medium	800	Off	0
40	0	0	0	Off	Medium	0	Off	0

■ SLA-E13200 series

Program Name: DNA-RICE-AUTO						Model: SLA-E13200 series			
Step	Well	Temp (°C)	Mixing (M)	Collect (S)	Rod	Mixing speed	Volume (μL)	Pause	Vapor (M)
1	6	70	20	0	Off	Medium	600	Off	0
2~26	6	70	1	0	Off	Medium	600	Off	0
27	6	70	10	0	Off	Medium	600	Off	0
28	6	45	5	0	Off	Medium	600	On	0
29	6	45	5	0	Off	Medium	900	Off	0
30	5	45	5	0	Off	Medium	900	Off	0
31	3	45	1	60	On	Medium	800	Off	0
32	4	45	1	60	On	Medium	800	Off	0
33	6	45	5	100	On	Medium	900	Off	0
34	5	45	5	100	On	Medium	900	Off	0
35	4	45	3	100	On	Medium	800	Off	0
36	3	45	2	60	On	Medium	800	Off	0
37	2	45	2	60	On	Medium	800	Off	10
38	1	45	5	100	On	Medium	150	Off	0
39	2	N / A	1	0	Off	Medium	800	Off	0
40	0	N / A	0	0	Off	Medium	0	Off	0

12. Reagent performance

■ The stability of extracted DNA

Storage Conditions	DNA stability
-80°C	Over 90 days
-20°C	28 days
4°C	14 days
25°C	2 days
Freeze-thaw	10 times

13. Explanation of Symbols

	Manufacturer		Consult instructions for use
	Temperature limit		Contains sufficient for test
	Catalogue number		Caution
	Batch code		Non-sterile
	Do not re-use		Keep away from sunlight
	Date of manufacture		Use-by date
	For research use only		