

Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)

User Guide



Version 1.0

In-Vitro Diagnostics / For use with Automatic nucleic acid extractor compatible with Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)



T821H



Xi'an Tianlong Science and Technology Co., Ltd.

No.4266, Shanglin Road, Weiyang District, Xi'an, 710021, Shaanxi, P.R. China

Contents

Intended Use	
Product Performance Indicators	
Special Notes	1
Testing Principle	
Content of the Kit	2
Materials Required but not Provided	3
Warnings and Precautions	3
Precautions for Safe Handling	4
Reagent Storage and Handling	4
Sample Handling and Storage	4
Operation Guide	5
1. Automated Extraction Process	5
2. Operation Steps of Automated Extraction	5
2.1 Automatic Nucleic Acid Extractor (model:Npex 192)	5
Troubleshooting Guide	7
Quality Control	8
Limitations of Test Methods	8
Safety Symbols and Signs	8
Contact Information.	c



Intended Use

The *Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)* is intended for rapidly extracting genomic DNA from the samples of plant tissues (e.g. blade, pulp, seed, and tuber etc.). The extracted genomic DNA is of high purity and stability and can be used in a variety of routine operations, including enzyme digestion, Polymerase Chain Reaction (PCR), DNA library constructions, Southern hybridization and blotting and other experiments.

The *Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)* is intended to be used by professionals, such as biotechnologists, microbiologists, clinical technicians, and physicians who are trained in molecular and biological techniques.

Product Performance Indicators

The extraction kit can efficiently extract nucleic acids from plant tissue sample.

The coefficient of variation (CV) of intra-assay and inter-assay for the extraction kit is less than 15%.

Special Notes

The *Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)* is worked with TIANLONG® automatic nucleic acid extractor (Npex 192) that has been disinfected by UV light before use. After an experiment, wipe the inside of the extractor with 75% ethanol and disinfect it with UV light for 15 mins. An automatic nucleic acid extractor automates the entire purification process and can process 1-192 samples in a single run.

The **Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)** is particularly used for targeted genomic DNA isolation. Therefore, all of experiment supplies, such as pipettes, tubes, tips, must be processed by autoclave. Operator should wear gloves and masks and protective coveralls.

The kit has magnetic beads with a unique separation function and buffer system to extract, separate and purify high-quality nucleic acids from the samples of plant tissues .

Magnetic beads enable the purification of high-quality nucleic acids that are free of protein, nuclease, and other impurities. Purified nucleic acids can be widely used in the fields of diagnostics, genomics research, disease detection, food safety and forensic identification, etc.

Please carefully read the manual of instructions before attempting to install or use the product for the first time. To consider all possible consequences of incorrect operation or non-recommended functions, pay special attention to the possible consequences.

Testing Principle

The *Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)* is worked with TIANLONG® automatic nucleic acid extractor (Npex 192). During the nucleic acid extraction process, magnetic beads are adsorbed, transferred and released by special magnetic rods based on the principle of magnetic bead adsorption. The extraction process enables the conduction of nucleic acid extraction and final adsorption of highly pure nucleic acids with the transfer of magnetic beads and nucleic acids.

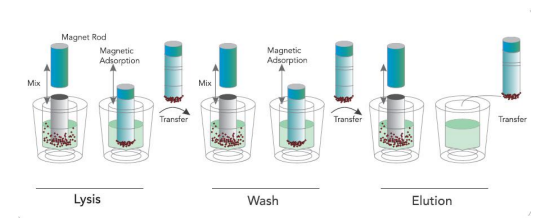


Figure 1. Schematic Diagram of Automatic Nucleic Acid Extractor

An automatic nucleic acid extractor performs the following steps on a sample which contains magnetic particles:

A magnetic rod protected by the mixing sleeve inserts into a well which contains sample. The mixing sleeve stirs rapidly and repeatedly in the liquid to ensure complete mixing of the liquid and magnetic beads. After cell lysis, nucleic acid adsorption, washing and elution, highly pure nucleic acid is obtained.

Npex 192 is equipped with an array of 192 magnetic rods, allowing it to process up to 192 samples simultaneously.

Content of the Kit

	Short Code	T821H	
ame of Component		ΙοΖΙΠ	
	Size	192 T/Box (Pre-filled)	
REAG1	Component	Pre-filled 96-deep well plate	
	Quantity	2	
	Component Specification	96 Tests	
	Size	192 T/Box (Pre-filled)	
REAG2	Component	Pre-filled 96-deep well plate	
	Quantity	2	
	Component Specification	96 Tests	
REAG3	Size	192 T/Box (Pre-filled)	
	Component	Pre-filled 96-deep well plate	
	Quantity	2	
	Component Specification	96 Tests	
	Size	192 T/Box (Pre-filled)	
REAG4	Component	Pre-filled 96-deep well plate	
	Quantity	2	
	Component Specification	96 Tests	
	Component	pretreatment reagent	
REAG5	Component Specification	76.8 mL	
	Quantity	1	
REAG6	Component	nucleic releaser	
	Component Specification	11.52 mL	
	Quantity	1	
Inst	tructions for Use	1 Copy	



*Note: Ribonuclease A and Ribonuclease A Diluent should be purchased separately

Materials Required but not Provided

When working in a laboratory, make sure to wear a proper lab coat, powder-free disposable gloves and protective goggles. For more information, please consult the Safety Data Sheet (SDS) available from the product supplier.

- Pipettor: 200 μL or 1000 μL
- Tip: 200 μL or 1000 μL
- Vortex Mixer
- High-speed centrifuge
- Water bath or metal bath
- Sample Holder
- 75% Ethanol
- Ribonuclease A and Ribonuclease A Diluent can be purchased from Tianlong
- Extractor

Warnings and Precautions

Please be sure to read the precautions before using the kit.

The Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192) is particularly used for targeted genomic DNA isolation; therefore, all of experiment supplies, such as pipettes, tubes, tips, must be processed by autoclave. Operator should wear gloves and masks.

Please read the manual carefully before using the kit, and strictly follow the manual throughout operation. The samples should be collected on a clean bench or in a bio-safety cabin.

Before using TIANLONG® automatic nucleic acid extractor (Npex 192), it must be disinfected by UV light. After an experiment, wipe the inside of the extractor with 75% ethanol and disinfect it with UV light for 15 minutes.

Due to the possibility of residual magnetic beads in the eluate following extraction, every possible effort should be made to avoid suctioning of any magnetic beads during eluate absorption.

Do not mix reagents from different batches and use the kit within expiry date.

Dispose of all samples and reagent materials used in an experiment, and thoroughly clean and disinfect the experimental workbench.

The Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192) is intended for in vitro diagnosis use.

When using kit, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate Material Safety Data Sheets (MSDSs). These documents are available online in a convenient and compact PDF format at

https://www.medtl.net/resources/download/catalogue-all/catalogue, where the operator can find, view and print the appropriate MSDSs.



Caution: Do not add any bleach or acidic solution directly to the pre-filled reagent.

The pre-filled reagent contains guanidinium salts, which, when combined with bleach can form highly reactive compounds. If any of buffers are spilled, clean immediately with a suitable laboratory detergent and water. If the spilled liquid contains potentially infectious agents, clean the affected area first with laboratory detergent and water first. Then clean with sodium hypochlorite at a concentration of 1% (v/v).

The Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192) comes with the following warnings and precautions

Name of Co	Component Hazard pictograms (CLP)		Classification under CLP:	H- and P-statements
			Acute toxicity (oral),	Hazard statements (CLP)
			Category 4	H302: Harmful if swallowed.
			Skin corrosion/irritation,	H315: Causes skin irritation.
			Category 2	H319: Causes serious eye irritation.
			Serious eye damage/eye	Precautionary statements (CLP)

			irritation,	P264: Wash hands, forearms and face thoroughly
			Category 2	after handling.
				P280: Wear protective gloves/protective
REAG 1	Lysis solution			clothing/eye protection/face protection/hearing
				protection.
				P321: Specific treatment (see supplemental first
				aid instruction on this label).
				P337+P313: If eye irritation persists: Get medical
				advice/attention.
ı				P501: Dispose of contents/container to hazardous
				or special waste collection point, in accordance
				with local, regional, national and/or international
				regulation.
REAG 2	Washing A	None	None	None
REAG 3	Washing B	None	None	None
REAG 4	Elution	None	None	None
REAG 5	pretreatment reagent	None	None	None
				Hazard statements (CLP)
		nucleic releaser	Serious eye damage/eye irritation, Category 2	H319 - Causes serious eye irritation.
REAG 6				Precautionary statements (CLP)
	nucleic releaser			P280 - Wear protective gloves/protective
	Tracicio reicaser			clothing/eye protection/face protection/hearing
				protection.
				P337+P313 - If eye irritation persists: Get medical
				advice/attention.

Please see MSDS for more details.

Precautions for Safe Handling

Do not dispose of the preparations or the packaging waste in drains leading to the sewage system or in the drainage system for waste not produced by industrial processing/analysis waste.

Any material in contact with reagents should be treated as a biological contaminant and treated in accordance with relevant local regulations.

Reagent Storage and Handling

The *Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)* should be stored at room temperature in a cool, dry and well-ventilated area. All components of the kit can be adequately stored for 12 months.

The kit should be used in a well-ventilated area, keep away from the source of heat, sparks, open flames, and smoking.

To avoid evaporation, the pre-filled reagent should be used immediately after opening and should not be placed for a long period of time.

Avoid exposure to UV light (e.g., for decontamination), which may result in accelerated aging.

Sample Handling and Storage

Avoid foam inside or on the samples. Depending on the starting material, sample pre-treatment may be required. Samples should be stored at room temperature (15~25 °C) before starting the experiment.

Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192) (T821H) –User Guide

Page 4 of 9



For detailed information on sample pretreatment, please refer to 2.1.3.

Operation Guide

1. Automated Extraction Process

Automatic nucleic acid extractor (Npex 192) enables nucleic acid extraction by magnetic beads. It uses magnetic rods to move the beads adsorbed with nucleic acid into different reagent wells. Magnetic rod protected by the mixing sleeve which stirs rapidly and repeatedly in the liquid to ensure complete mixing of the liquid and magnetic beads. After cell lysis, nucleic acid adsorption, washing, and elution, the highly pure nucleic acids are obtained. Automatic nucleic acid extractors are characterized by high automation, rapid extraction speed, stable results, and ease of operation.

The user needs to load samples and magnetic bead nucleic acid extraction reagents into the reaction consumables. The nucleic acid extractors are going to perform all nucleic acid extraction operations according to the experimental procedures. Please refer to the user manual provided with an instrument for operating instructions.

2. Operation Steps of Automated Extraction

2.1 Automatic Nucleic Acid Extractor (model: Npex 192)

2.1.1 Edit Experiment Program

The extraction procedure of Npex 192 Nucleic Acid Extractor is as follows:

Step	Name	Plate	Stir (min: s)	Magnetic (min: s)	Wait (min: s)	Speed	Volume	Temp (°C)	Heating State
1	Remove bead	W2	01:00	01:00	00:00	8	625 μL	0	OFF
2	Lysis	L	10:00	00:45	00:00	8	600 μL	90	ON
3	Washing A	W1	03:00	00:30	00:00	8	600 μL	80	ON
4	Washing B	W2	01:00	01:10	01:00	8	625 μL	80	ON
5	Elution	Е	05:00	00:40	00:00	8	100 μL	80	ON

2.1.2 Reagent Preparation

96-deep well plate:

Open the kit and take out the REAG1\REAG2\REAG3\REAG4 from the plastic package, slowly invert it several times to resuspend the magnetic beads. Gently shake the 96-well plate so that the reagent and magnetic beads are concentrated on the bottom of the 96-well plate (A 96-well plate horizontal centrifuge can also be used for centrifugation at 500 rpm for 1 min). Carefully tear off the aluminum foil sealing film before use to avoid liquid splashing.

2.1.3 Adding Sample to the Reagent

Ribonuclease A solution:

Dilute Ribonuclease A with Ribonuclease A dilution to final concentration of 20 mg/mL, it can be used after completely dissolved. The dissolved solution should storage at -20 °C, and repeated freezing and thawing shall not exceed 5 times.

Sample Pretreatment

Take out appropriate amount of plant tissue from the fresh tissue 100 mg or dry weight 50 mg, different samples please refer to the table as below. Then placed it in the pre-cooled mortar, adding liquid nitrogen, fully crushed (In order to avoid thawing, please add liquid nitrogen constantly); after that transfer it to 1.5 mL sterile centrifugal tube.

Samples	Amount of samples
Fresh Plant Blade	50~100 mg
Dried Leaves	50~100 mg
Soybean Seed	100~200 mg

Wheat Seed	50~100 mg
Corn Seed	100~200 mg

Add 400 μ L REAG5, and 60 μ L REAG6 into 1.5 mL sterile centrifuge tube with plant tissue. Mix thoroughly 5-6 times upside down during the water bath (water bath: 65°C for 30 mins). For dry weight tissue, water bath time should be appropriately extended; when the sample is seed, grind with a powder grind, then treat it with liquid nitrogen.

Remove the 1.5 mL sterile centrifuge tubes from the incubate, centrifuged 5 mins at 12000 rpm and the supernate is taken to be used for next step.

96 deep well plate: Add 300 μ L supernate to the REAG1 plate of the pre-filled reagent and add 20 μ L Ribonuclease A solution to the REAG2 plate of the pre-filled reagent. (Be aware of the effective wells).

Caution: When pipetting the sample, avoid having substance than liquid adhere to the tip of the sample injector; do not add the sample too quickly to avoid contaminating the upper portion of the well wall; and do not splash air bubbles to avoid contaminating adjacent wells.

Note: The following points should be taken into consideration when determining whether a sample is suitable for the *Nucleic Acid Extraction Kit*.

a. Type of sample: As stated in the intended use.

2.1.4 Loading in Deep Well Plate

Place the 96-deep well plates in the Automatic Nucleic Acid Extractor correctly, REAG1 at the L position, REAG2 at the W1 position, REAG3 at the W2 position and REAG4 at the E position, ensuring that the marked notch of the plate is on the left.

Note: The user should ensure the marked notch of the plate faces is on the left, which is shown in Figure 2.

Note: Place the 96-deep well plate docking into the experiment cabin and push the mixing sleeves into the right position. Check the position of the mixing sleeves. Otherwise, instrument dysfunction or malfunction may occur and affect the experiment results.

Caution: The user must ensure that the rotatory mixing sleeve is placed properly. Otherwise, the instrument may operate abnormally, or the magnetic rods may be contaminated.

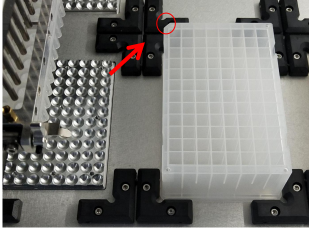


Figure 2. 96-deep well plate

2.1.5 Experimental Procedure Run

For special operations please see 2.1.1. After the procedure is completed, the instrument will notice the user the experiment has been completed. Transfer the extracted product from the REAG4 plate to clean centrifuge tubes which are free of nuclease.

• Note: If the user does not analyse the extracted product immediately, please seal and store in a refrigerator at -20 °C.

Caution: Any used deep well plate and mixing sleeve should be considered as biological contaminants and disposed of in accordance with relevant regulations.

Caution: Using expired reagents or those that are not compatible with this instrument does not guarantee expected results.

Follow the Cleaning and Maintenance of the Instrument in accordance with the user manual provided with the equipment. Ensure that the experimental cabin is cleaned regularly to minimize the risk of cross-contamination.

Troubleshooting Guide

This troubleshooting guide should assist you in resolving any problems that arise during the experimental process. For more information, please visit our Technical Support Centre and Frequently Asked Questions, page at http://www.medtl.net. The scientists in our Tianlong company's Technical Services Department are always available to answer any questions you may have about the information and protocols contained in the manual, as well as sample and assay technologies (for contact information is included on the back cover or at http://www.medtl.net).

When an exception or error occurs during the experiment, the current run step is terminated/stopped. After resolving the error or exception, restart the run from the beginning. The troubleshooting guide is shown in the following table.

No.	Fault Symptom	Fault Cause	Handling Method
1	The well plate vibrates and the liquid splashes when tearing off the aluminum foil sealing film.	When tearing the film, please press the well plate to prevent it from rocking.	The reagent for this plate shall be scrapped, and re-extraction shall be performed.
2	Add the sample to unexpected wells.	Please read this manual carefully before adding samples.	The reagent for this plate shall be scrapped, and re-extraction shall be performed.
3	The amount of liquid in the reagent wells is insufficient	/	Contact the after-sales service of Tianlong.
4	Reuse of pre-filled components	Please read the precautions in this manual before using the kit.	Perform re-extraction of nucleic acid.
F	Abnormal noise from the instrument during extraction	The 96-deep well plate may be placed incorrectly.	Reposition the deep well plate.
5		The mixing sleeve may not be inserted in place.	Reinsert the mixing sleeve.
		Please follow the operation requirements in the manual	Contact the after-sales service of Tianlong.
6	6 Poor extraction performance	The temperature control components of the instrument may be abnormal.	Contact the after-sales service of Tianlong.
		Other	Contact the after-sales service of Tianlong.

^{*} Ensure that the reagents have been preserved and used according to the manufacturer's instructions.

Quality Control

In accordance with Tianlong Company's ISO-certified Quality Management, each lot of **Nucleic Acid Extraction Kit (For Plant Tissues Genomic DNA-Npex192)** is tested against predetermined specifications to ensure consistent product quality.

Limitations of Test Methods

The system performance has been established through performance evaluation studies using the samples of plant tissues to extract genomic DNA.

It is the user's responsibility to validate system performance for any procedures used in their laboratory that are not covered by the performance evaluation studies of Xi'an Tianlong Science and Technology Co., Ltd.

The extraction kit is intended for clinical diagnostics, health system and scientific research only, whose usage can act as an ancillary step for molecular detection and should be matched with other molecular detection methods. The concentration and purity of its extraction product are affected by instruments and operators. Any generated diagnostic results must be interpreted in conjunction with the other clinical or laboratory findings.

Safety Symbols and Signs

No.	Symbol	Implication
1	REF	Catalogue number
2	LOT	Batch code
3	Σ <n></n>	Contains sufficient for <n> tests</n>
4	\square	Use by date
5	\triangle	Caution
6	¥	Temperature limit
7	(!)	Reminder
8		Manufacturer
9	②	Do not re-use
10	CONT	Content of the kit
11	REAG1	Lysis solution
12	REAG2	Washing 1
13	REAG3	Washing 2
14	REAG4	Elution
15	REAG5	Pretreatment Reagent
16	REAG6	Nucleic Acid Releaser



17		Warning
18	PAP	PAP21: Not-corrugated cardboard

Contact Information

For technical assistance and more information, please contact our Technical Support Center at +86-29-826 82132 (Tel), +86-29-82216680 (Fax), inquiry@medtl.com or contact your local distributor.

For up-to-date licensing information or product-specific disclaimers, please see the respective User Guide. Tianlong User Guides are available at www.medtl.net or can be requested from Tianlong Technical Services or the local distributor.

IFU T821H EN © 2024 Xi'an Tianlong Science and Technology Co., Ltd., all rights reserved.