

MagPurix[®] Plant DNA Extraction Kit (ZP02014)

Instructions for Use (Handbook)



Version: 1.8



48



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Intended Use

The MagPurix[®] Plant DNA Extraction Kit provides a complete set of reagents and consumables for the automated purification of genomic DNA from plant tissue and yeast with MagPurix system.

The product is intended to be used by professional users, such as technicians and physicians who are trained in molecular biology techniques.

Introduction

Product Name	MagPurix [®] Plant DNA Extraction Kit
Catalogue Number	ZP02014
Product Overview	The MagPurix [®] Plant DNA Extraction Kit is designed to extract genomic DNA from plant tissue and yeast. The kit uses unique magnetic ZiBeads [®] technology and in combination with MagPurix [®] series automatic instruments, achieved superior product quality, consistent and high product yield and reproducible results. The final product is suitable for a wide range of diagnostic and research applications, such as sequencing, genotyping, qPCR, ddPCR and NGS assays.
Applicable Instrument Model	All MagPurix [®] Instrument
Display Protocol Name on The Instrument	2014 PLANT DNA
Applicable Instrument Firmware	Check and download the latest firmware from www.zinexts.com
Processing Time	MagPurix [®] 12 series 45-60 minutes MagPurix [®] 24 series 50-65 minutes MagPurix [®] EVO series 40-45 minutes

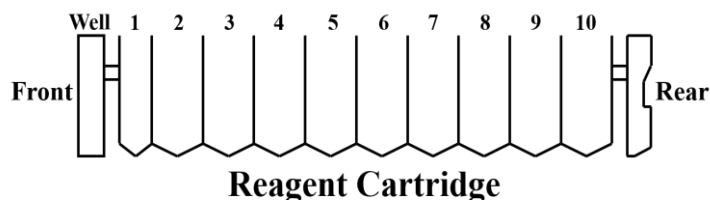
Kit Contents and Storage

Shipping and Storage	The kit is shipped at room temperature. Upon receipt, store the kit at room temperature. All kit components are stable when stored properly until the expiration date shown on the kit box.	
Kit Content	The components supplied in the kit are listed below. Sufficient reagents are supplied to perform 48 purifications.	
	Contents	Amount
	1 Reagent Cartridge	48 pcs (6x8)
	2 Reaction Chamber	48 pcs (6x8)
	3 Tip Holder	48 pcs (6x8)
	4 Piercing Pin	50 pcs
	5 Filter tip	50 pcs
	6 Sample Tube (2 ml)	50 pcs
	7 Elution Tube (1.5 ml)	50 pcs
	Filter Column	50 pcs
	Collection Tube	50 pcs
	RNase A, 10 mg/mL (0.5 ml)	1 pc
	PLA Buffer (25 ml)	1 pc
	PLB Buffer (25 ml)	1 pc
	Barcode sticker (EVO only)	50 pcs

Reagent
Cartridge
Contents

Each Reagent Cartridge has 10 positions with 10 sealed well.
Positions 1-10 contain wells filled reagents for this protocol.

Reagent	Well No.
Empty	1
Lysis Buffer 2	2
Binding Buffer 1	3
Magnetic Bead Solution	4
Washing Buffer 1B	5
Washing Buffer A	6
Washing Buffer B	7
Elution Buffer 1	8
Elution Buffer 2	9
Empty	10



Materials Required But Not Provided

The following general laboratory equipment and consumables are required to perform the extraction. All laboratory equipment should be installed, calibrated, operated, and maintained according to the manufacturer's recommendations. The following tables display required and special equipment along with the list of consumables.

Item
MagPurix [®] series instrument
1.5 or 2.0 ml microcentrifuge tubes
Pipettes and filter tips
Phosphate-buffered saline (PBS, may be required for diluting samples)
Optional: Plastic consumables, DNase-free RNase A (to minimize RNA content)

Warnings and Precautions

For research use only. Read all the instructions carefully before using the kit. Use of this product should be limited to trained personnel in the techniques of DNA purification. Strict compliance with the user manual is required for optimal results. Attention should be paid to expiration dates printed on the box and labels of all components. Do not use a kit after its expiration date.

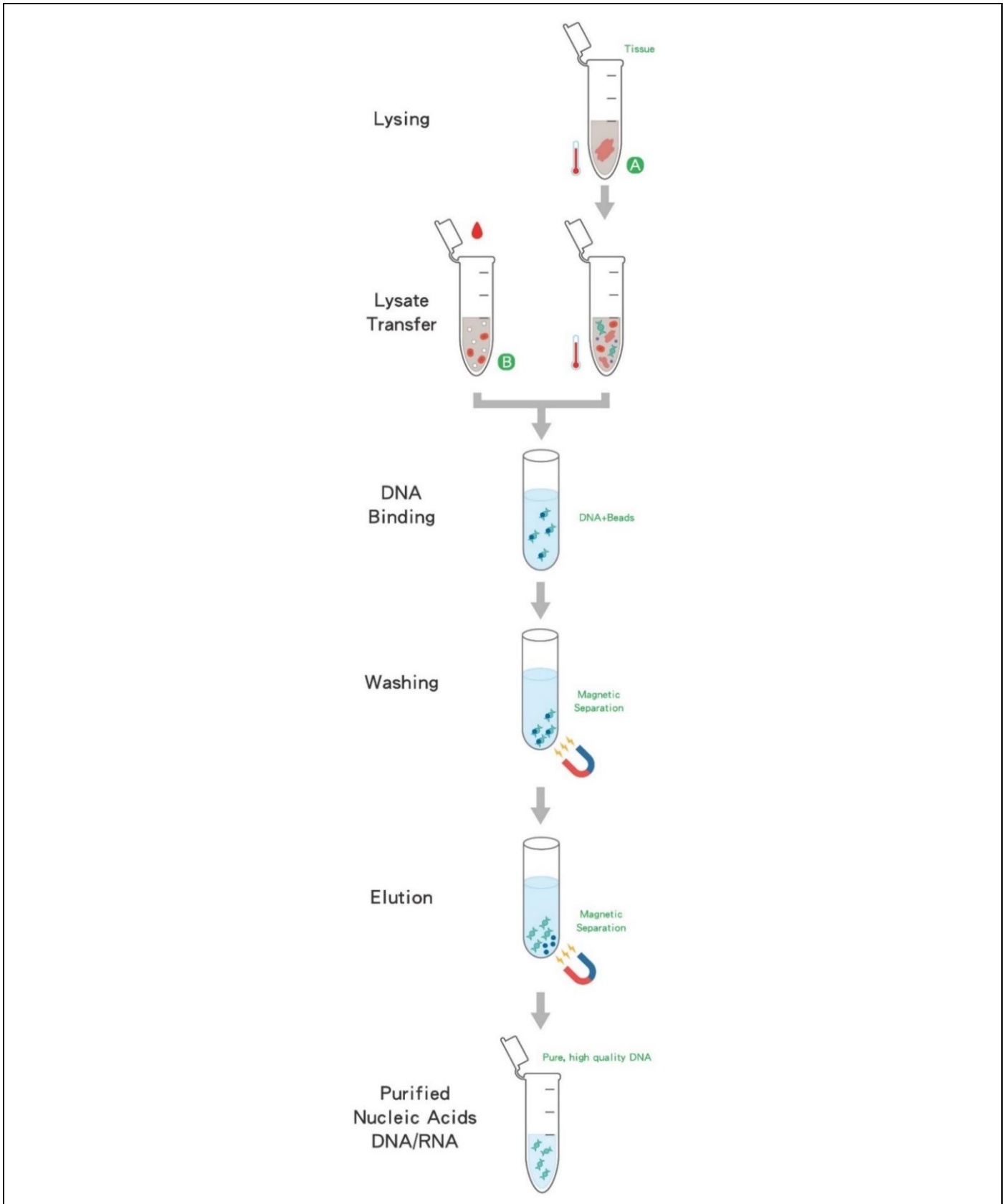
When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate safety data sheets (SDSs). These are available online in convenient and compact PDF format at

<http://www.zinexts.com/download.php?lang=en&tb=1&cid=7> where you can find, view, and print the SDSs for each kit and kit component.



CAUTION: DO NOT add bleach or acidic solutions directly to the sample preparation waste.

Purification Principle



A Perform certain pretreatment process before extraction.

B Transfer sample to extraction directly.

Before Starting

Preparation of sample materials

The purification procedure is optimized for the use of appropriate samples as below table.

Plant tissue	<ol style="list-style-type: none">a. Perform homogenization by using proper homogenizer.b. Add 440 μl lysis buffer to the sample.c. Vortex vigorously.d. Incubate the mixture at 65°C for 10 minutes in a thermomixer (set at 1000 rpm) or vortex several times during incubation in the heat block or water bath.e. Pre-filter the digested lysate using a Filter Column to remove residual debris.f. Short spin at 6.000 x <i>g</i> to collect the clear flow-through in Collection Tube.g. Add 10 μl RNase A, mix well, incubate for 10 minutes at room temperature.h. Transfer 100-400 μl into each Sample Tube.
Yeast/Suspension culture	<ol style="list-style-type: none">a. Centrifuge at 6.000 x <i>g</i>, 3 minutes.b. Remove supernatant.c. Add 440 μl lysis buffer, vortex for 30 seconds.d. Incubate the mixture at 65°C for 10 minutes in a thermomixer (set at 1000 rpm or vortex several times during incubation in the heat block or water bath.e. Pre-filter the digested lysate using a Filter Column to remove residual debris.f. Short spin at 6.000 x <i>g</i> to collect the clear flow-through in Collection Tube.g. Transfer 100-400 μl into each the Sample Tube.
Yeast/Colony	<ol style="list-style-type: none">a. Take 1-3 colony from culture plate with an inoculation loop and suspend in 440 μl of lysis buffer by stirring vigorously.b. Incubate the mixture at 65°C for 10 minutes in a thermomixer (set at 1000 rpm) or vortex several times during incubation in the heat block or water bath.c. Pre-filter the digested lysate using a Filter Column to remove residual debris.d. Short spin at 6.000 x <i>g</i> to collect the clear flow-through in Collection Tube.

e. Transfer 100-400 μ l suspension into each Sample Tube.

Note:

After harvesting plant tissues, it should be frozen in liquid nitrogen if not be used immediately. It can then be stored at -80°C . Alternatively, tissue can be dried or lyophilized after harvesting to allow storage at room temperature ($15\text{-}25^{\circ}\text{C}$). To ensure DNA quality, samples should be completely dried within 24 hours of collection. If possible, it is preferable to collect young materials (e.g., leaves, needles) since they contain more cells per weight and therefore result in higher yields.

When working with fungi, harvest mycelium directly from a culture dish or from liquid culture. For liquid culture, first pellet cells by centrifugation. Remove the supernatant completely before disruption and lysis. Fresh, frozen, or freeze-dried fungal material can be used.

The disruption method may require optimization to ensure maximum DNA yield and quality. Complete and quick disruption of starting material is essential to ensure high DNA yields and to avoid DNA degradation.

Before DNA extraction, plant material should be first mechanically disrupted with lysis buffer*. After homogenization, remove the debris and other precipitations by using Filter Column. Collect the clear flow-through and incubate with RNase A to digest the RNA in the sample before DNA extraction.

* We provide two lysis buffers: **PLA Buffer** and **PLB Buffer** for dealing with different tissue types. Before extraction of a new tissue type, try both the lysis buffer to get the optimized lysis procedure and a better DNA yield. If the precipitation formed in the lysis buffer, warm it at 65°C before using.

The suggested starting material and elution volume range for each nucleic acid extraction.

Sample type	Starting material per sample	Concentration (ng/ μ l)	Elution Volume
Plant tissue			
Soybean	100-400 μ l/100 mg seed	5-12 (PLA)/50-80 (PLB)	50-300 μ l (EVO 50-200 μ l)
Rice	100-400 μ l/20 mg seed	5-8 (PLA)/15-25 (PLB)	
<i>Arabidopsis</i>	100-400 μ l/100 mg leaf	2-5 (PLA)/5-7 (PLB)	
Tomato		20-40 (PLA)	
Corn		10-15 (PLA)/25-60 (PLB)	
<i>Tectaria</i>		5-10 (PLA)	
<i>Aspidistra</i>		3-6 (PLA)	
<i>Pharius</i>		20-25 (PLA)/50-100 (PLB)	
<i>Zingiber</i> -		3-8 (PLA)/20 -25 (PLB)	

Yeast			
Suspension Cultures	100-400 µl		50-300 µl (EVO 50-200 µl)
Colony	1-3 colony		

Procedure of MagPurix System Procedure

Workflow of MagPurix operation

Place the cartridge and plastic consumables on the MagPurix instrument



Select the protocol and setup the condition



Follow onscreen message for worktable setup



Start the protocol



Collect elution product *



UV decontamination

* Output the bench record (option)

Note: Perform all steps at room temperature (20-25°C) unless otherwise notified.

Purification Protocol- MagPurix[®] series


1	Turn on the Instrument	a. Turn on the power switch and wait for the screen to turn on.
2	Load new Consumable(s) and Cartridge(s)	a. Open the door and remove the Sample Rack from the instrument. b. Load 1 Reagent Cartridge, and all plastic disposables (2 Reaction Chamber, 3 Tip Holder, 4 Piercing Pins, 5 Filtered Tips and other components presented in the kit intended to use). c. Place 6 Sample Tubes and 7 Elution Tubes into the Sample Rack.
3	Load the Samples	a. Transfer appropriate volume of sample into each Sample Tube on the Sample Rack. b. Put the Sample Rack back into the instrument and close the door.


- | | | |
|----------|---------------------------|--|
| 4 | Program Set up | <ul style="list-style-type: none"> a. Scan the protocol barcodes to select the purification protocol, sample volume and elution volume. |
| 5 | Start Extraction | <ul style="list-style-type: none"> a. Follow the instructions displayed on the screen to double-check the operating steps being completed before program running. b. Press “ENTER” to start the experiment. Instrument will run the protocol program automatically until the whole process is completed. c. At the end of the run (approximately 12 series 45-60 minutes, 24 series 50-65 minutes), instrument alarms briefly. |
| 6 | Collect the Elution tubes | <ul style="list-style-type: none"> a. Open the instrument door. b. Collect the elution tubes containing the purified nucleic acids. c. The purified nucleic acids are ready for immediate use. Store the purified nucleic acids at 4°C (short-term, less than 10 days) or aliquot and store at -70°C (long-term) before performing downstream analysis. d. Discard the used cartridges and all plastic consumables into biohazard waste. *Do not reuse the cartridges. e. If you are not using the instrument immediately, place the Sample Rack back to the workplace, close the instrument door and press “Start” button for 2 seconds to enter sleep mode. Moreover, if the instrument will not be used in a long time, please turn off the power switch. |

Purification Protocol-MagPurix[®] EVO series

- | | | |
|----------|---|---|
| 1 | Turn on the Instrument | <ul style="list-style-type: none"> a. Turn on the power switch and wait for the screen to turn on. a. Login the instrument and enter the Home Page. |
| 2 | Load new Consumable(s) and Cartridge(s) | <ul style="list-style-type: none"> a. Open the door and remove the Sample Rack from the instrument. b. Open the Tip-Holder Lid. c. Load 1 Reagent Cartridge and all plastic disposables (2 Reaction Chamber, 3 Tip Holder, 4 Piercing Pins, 5 Filtered Tips and other components presented in the kit intended to use). d. Close the Tip-Holder Lid. e. Paste the barcode stickers on Elution Tubes. a. Place 6 Sample Tubes and 7 Elution Tubes into the Sample Rack. |
| 3 | Load the Samples | <ul style="list-style-type: none"> a. Transfer appropriate volume of sample into each Sample Tube on the Sample Rack. a. Put the Sample Rack back into the instrument and close the door. |

- 4** Program Set up
- a. Select the appropriate protocol program on the instrument. Press **NEXT**.
 - b. Select the appropriate Sample Volume and Elution Volume and press **NEXT**.
 - c. Press the number button to select the right Sample Numbers.
 - d. Scan/Edit each primary Sample ID directly. After finished, press **NEXT**.
 - e. Scan/Edit each Elution Tube ID directly. After finished, press **NEXT**.
 - f. Scan Reagent Cartridge Barcode. Press **NEXT**.
***If the cartridge is expired, the next step cannot be performed.**
 - a. Follow the instructions on the screen to double-check the operating steps being completed before running the program. Press **NEXT**.
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- 5** Start Extraction
- a. Check "**PROGRAM CONFIRMATION**" on the screen.
 - b. Press "**START**" to start the experiment. Instrument will run the protocol program automatically until the whole process is completed.
 - c. At the end of the run (approximately **40-45 minutes**), instrument alarms briefly and the screen indicates "**PROGRAM FINISH**".
 - d. If you want to perform the same protocol, press "**RERUN**" to perform the same experiment. If you do not need to re-run the experiment, press the function button " **HOME**" to exist the experiment mode.
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- 6** Collect the Elution tubes
- a. Open the instrument door.
 - b. Collect the elution tubes containing the purified nucleic acids.
 - c. The purified nucleic acids are ready for immediate use. Store the purified nucleic acids at 4°C (short-term, less than 10 days) or aliquot and store at -70°C (long-term) before performing downstream analysis.
 - d. Discard the used cartridges and all plastic consumables into biohazard waste. ***Do not reuse the cartridges.**
 - e. If you are not using the instrument immediately, please put the Sample Rack back into the instrument, close the instrument door, and press the " **POWER**" function button to enter sleep mode. If the instrument will not be used in a long time, please turn off the power switch.
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Troubleshooting

***This table is helpful for solving common problem. If you need other technical support, please contact Zinexts team (<http://www.zinexts.com/index.php?lang=en>) or your distributor.**

Problem	Possible Cause	Comments and suggestions
Poor DNA quality or yield	Deterioration or contamination of reagents.	Please ensure that the kit reagents are still in the effective using period before use. Discard any kit reagent that shows discoloration or evidence of microbial contamination.
	Kit stored under non-optimal conditions	Store kit at 15-25°C at all time after arrival. If either reagent or buffer precipitate upon shipping in cold weather or during long-term storage, dissolve precipitates by gently warming and stirring the solution. Please do not freeze the Reagent Cartridges.
	Insufficient sample input	DNA yield depends on the sample type and the number of nucleated cells in the sample. Please proportionally adjust the total input amount of sample to increase the DNA yield.
	Too much of elution buffer was used	The elution volume can be reduced proportionally.
	The eluate of final product(s) is not enough.	Please collect issue information and provide it to your Support Representative/Technical Support as soon as possible.
Clogging issue	Too much sample material was used.	Decrease the input amount of sample material or dilute your sample.
No results in downstream analysis	No signal/The PCR was inhibited.	Using appropriate controls for analysis. Check the positive control, negative control, water (NTC) and internal control to clarify the possible causes.
Instrument malfunction/abnormal sound	Abnormal consumables: 1. Deformed Filtered Tips 2. Deformed Reaction	Please replace the batch with normal consumables.

	Chamber 3. Deformed Tip Holder	
	Abnormal action of instrument: 1. Inaccurate correction value 2. Spare parts or components damaged	Please collect issue information (videos and pictures) and provide it to your Support Representative/Technical Support as soon as possible to calibrate or replace any other damaged or worn parts.

Related Products

Product Name	Cat. no.
MagPurix [®] Blood DNA Extraction Kit 200	ZP02001
MagPurix [®] Blood DNA Extraction Kit 1200	ZP02002
MagPurix [®] Viral NA Extraction Kit	ZP02003
MagPurix [®] Tissue DNA Extraction Kit	ZP02004
MagPurix [®] Cultured Cell DNA Extraction Kit	ZP02005
MagPurix [®] Bacterial DNA Extraction Kit	ZP02006
MagPurix [®] HPV DNA Extraction Kit	ZP02007
MagPurix [®] TB DNA Extraction Kit	ZP02008
MagPurix [®] FFPE DNA Extraction Kit	ZP02009
MagPurix [®] Forensic DNA Extraction Kit	ZP02010
MagPurix [®] Viral Pathogen DNA Extraction Kit A	ZP02011
MagPurix [®] Viral Pathogen DNA Extraction Kit B	ZP02012
MagPurix [®] Viral RNA Extraction Kit	ZP02013
MagPurix [®] Plant DNA Extraction Kit	ZP02014
MagPurix [®] Total RNA Extraction Kit	ZP02015
MagPurix [®] Viral NA Extraction Kit LV	ZP02016
MagPurix [®] CFC DNA Extraction Kit	ZP02017
MagPurix [®] cfDNA Extraction Kit Plus	ZP02024
MagPurix [®] cfDNA Extraction Kit LV	ZP02025
MagPurix [®] Coronavirus RNA Extraction Kit	ZP02027

References

- Tan SC *et al.* J Biomed Biotechnol. (2009)

Limited Product Warranty

Zinexts Life Science is committed to provide 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 any question or concerns, contact our Technical Support Representatives.

Zinexts Life Science guarantees the performance of all products according to the specifications stated in our product literature. The purchasers/users must determine the suitability of the product for their particular use. We reserve the right to change, alter, or modify any product to enhance its performance and design.

This warranty limits Zinexts Life Science Corporation'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 and used in accordance with instructions.

Revision History

Version	Date	Description
1.0	20 Feb. 2020	New document release
1.2	05 Jun. 2020	Update information
1.3	31 Jul. 2020	Update information
1.4	05 Mar. 2021	Revise the REP information
1.5	07 Jul. 2021	Revise the REP information
1.6	20 Aug. 2021	Revise the address
1.7	17 Sep. 2021	Revise the address
1.8	16 Feb. 2022	Revise the address