

# Ribospin™ vRD

VIRAL RNA/DNA PURIFICATION HANDBOOK

REF

302-150/302-103



HB3200



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GeneAll

## **Customer & Technical Support**

Should you have any further questions, do not hesitate to contact us.  
We appreciate your comments and advice.

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This protocol handbook is included in :

GeneAll® Ribospin™ vRD (302-150, 302-103)

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# Used symbols and Markings

<b>REF</b>	Catalog number	<b>IVD</b>	In vitro diagnostic medical device
<b>LOT</b>	Batch number	<b>H B</b>	Handbook code
	Use by		Consult instruction for use
	Manufacturer information		Contains sufficient for <N> tests
	Do not reuse		Temperature limitation
	Production date	<b>EC REP</b>	European Authorized Representative
	Important note	<b>CONC</b>	Contains the concentrated solution. Additional material must be added before use
	Write down the current date after adding ethanol to the bottle	<b>EtOH ?</b> <input checked="" type="checkbox"/>	Mark up after adding ethanol

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## Kit Contents

Components	Quantity		Storage
Cat. No.	302-150	302-103	
No. of preparation	50	300	
Buffer VL	30 ml	170 ml	
Buffer RB I (concentrate) *	8 ml	48 ml	
Buffer RBW (concentrate) *	13 ml	77 ml	
Buffer RNW (concentrate) * †	6 ml	34 ml	
Nuclease-free water	15 ml	20 ml	Room temperature (15~25°C)
Column Type V (mini) (with collection tube)	50	300	
1.5 ml microcentrifuge tube	50	300	
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\* Before first use, add absolute ethanol (ACS grade or better) into Buffer RB I, RBW and RNW as indicated on the bottle.

† Contains sodium azide as a preservative

## Product Specifications

Ribospin™ vRD	
Type	Using spin column
Maximum volume of starting samples	300 µl/prep
Preparation time	20 min
Maximum loading volume	800 µl
Minimum elution volume	30 µl

## Quality Control

All components of Ribospin™ vRD are manufactured in strictly clean conditions, and their degree of cleanliness is monitored periodically.

To maintain consistency, a quality control process is carried out thoroughly from lot to lot and only the qualified kits are approved for delivery according to ISO 9001:2008 and EN ISO 13485:2012.

## Storage Conditions

All components of Ribospin™ vRD should be stored at room temperature (15~25°C).

During shipment or storage under cool ambient condition, a precipitate can form in Buffer VL. In such a case, heat the bottle to 56°C to dissolve completely. Ribospin™ vRD is guaranteed until the expiration date printed on the product box.

## Safety Information

The buffers included in the Ribospin™ vRD contain irritants which are harmful when in contact with skin or eyes, or when inhaled or swallowed. Care should be taken when handling such materials. Always wear gloves and eye protection, and follow standard safety precautions.



Buffer VL, RB I, and RBW contain chaotropes, which can form highly reactive compounds when combined with bleach. Do NOT add bleach or acidic solutions directly to the sample-preparation waste.

## Preventing RNase contamination

RNase can be introduced accidentally during RNA purification. Wear disposable gloves always, because skin often contains bacteria and molds that can be a source of RNase contamination. Use sterile, disposable plastic wares and automatic pipettes to prevent cross-contamination of RNase from shared equipment.

## Product Description

Ribospin<sup>TM</sup> vRD provides a convenient method for isolation of RNA and DNA from cell-free fluid, cell-culture medium, plasma, serum, swab, urine and virus-infected samples.

Ribospin<sup>TM</sup> vRD utilizes the glass fiber membrane technology for the fastest and the most convenient nucleic acid isolation as a sufficient level for downstream application instead of conventional alcohol precipitation or phenol/chloroform extraction.

The buffer system of Ribospin<sup>TM</sup> vRD provides the effective binding condition of RNA and DNA to glass fiber membrane and the impurities on the membrane are washed away by two different wash buffers. At least, pure RNA and DNA are eluted in Nuclease-free water. The whole procedure may take only 15 minutes at room temperature and the eluate is suitable for PCR, RT-PCR, or any downstream application without further manipulation. The purified nucleic acid should be treated with care because RNA is very sensitive to contaminants such as RNases, often found on general lab ware and dust. To ensure RNA-stability after extraction, it is recommended to store at 4°C for immediate analysis or to freeze at -70°C for long-term storage.

# PROTOCOL FOR

## Ribospin™ vRD

### Equipment and reagents to be supplied by user

- \* Ethanol (>99%, ACS grade or better)
- \* 1.5 ml microcentrifuge tubes
- \* Micropipettes and sterile pipet tips
- \* Centrifuge capable of attaining 10,000 x g
- \* Vortex mixer



- Ethanol (>99%, ACS grade or better) must be added before the first use of Buffer RB1, RBW and RNW. Please refer to the information on the label of each bottle.
- If a precipitate is formed in Buffer VL, heat to 56°C to dissolve completely before use.

### **1. Transfer up to 300 µl sample (swab-storage media, cell-free fluid, cell-culture supernatant, plasma, serum, urine) in 1.5 ml microcentrifuge tube.**

### **2. Add 500 µl Buffer VL to the tube and lyse the sample by pipetting or vortexing.**

The volume of Buffer VL can be adjusted in proportion to the volume of sample.

For proper lysis, the complete mixing of sample and Buffer VL is essential.

### **3. Incubate the lysate for 10 min at room temperature.**

After this step, briefly centrifuge the tube to remove drops from the inside of the lid.

### **4. Add 700 µl Buffer RB1 to the lysate and mix thoroughly by inverting or vortexing.**

The volume of Buffer RB1 can be adjusted in proportion to the volume of lysate.

\* Do NOT centrifuge at this step.

### **5. Transfer up to 750 µl of the mixture to a Column Type V (mini).**

**6. Centrifuge at  $\geq 10,000 \times g$  for 30 sec at room temperature.**

Discard the pass-through and reinsert the mini column back into the same tube.

**7. Repeat step 5~6 with the remainder of the sample.**

Discard the pass-through and reinsert the mini column back into the same tube.

**8. Add 500  $\mu l$  Buffer RBW to the mini column.**

**9. Centrifuge at  $\geq 10,000 \times g$  for 30 sec at room temperature.**

Discard the pass-through and reinsert the mini column back into the same tube.

**10. Add 500  $\mu l$  Buffer RNW to the mini column.**

**11. Centrifuge at  $\geq 10,000 \times g$  for 30 sec at room temperature.**

Discard the pass-through and reinsert the mini column back into the same tube.

**12. Centrifuge at  $\geq 10,000 \times g$  for an additional 1 min at room temperature to remove residual wash buffer. Transfer the mini column to a new 1.5 ml microcentrifuge tube (provided).**

Residual ethanol may interfere with downstream reactions. Care must be taken at this step for eliminating the carry-over Buffer RNW.

If the carry-over Buffer RNW still occurs, centrifuge again for 1 min at full speed before transferring the column to the new 1.5 ml microcentrifuge tube.

**13. Add 30~50  $\mu l$  of Nuclease-free water to the center of the membrane in the mini column.**

**Let it stand for 1 min.**

**14. Centrifuge at  $\geq 10,000 \times g$  for 1 min at room temperature.**

Purified nucleic acid can be stored at 4°C for immediate analysis or at -70°C for long-term storage.

## Troubleshooting Guide

Facts	Possible Causes	Suggestions
<b>Low yield</b>	<b>Poor quality of starting material</b>	Fresh sample or well-conserved sample should be used for good result. Repeated freezing and thawing the sample should be avoided.
	<b>Low concentration of viral particle in the starting sample</b>	Use more starting sample. If the amount of sample is more than 300 $\mu$ l, concentrate the volume to 300 $\mu$ l using a micro-concentrator.
	<b>Inefficient or insufficient lysis</b>	Be sure to incubate for 10 minutes at room temperature after adding Buffer VL. For proper lysis, the complete mixing of the sample and Buffer VL is essential.
	<b>Improper elution</b>	Add Nuclease-free water to the center of the mini column membrane and perform incubation for 1 minute before centrifugation.
	<b>Precipitate in Buffer VL</b>	A precipitate can be formed in Buffer VL at cool ambient temperature. It is because the Buffer VL is saturated and its solubility would be reduced at low temperature. Before experiment, any precipitate in the Buffer VL should be dissolved completely by heating the buffer at 56°C or above until it disappears.
	<b>Degradation of RNA</b>	RNase can be introduced during purification of nucleic acid. Be certain not to introduce any RNases during the procedure of later handling. Keep tubes closed whenever possible during the extraction and use RNase-free products with sterile and disposable plastic ware.
	<b>Buffer RBI, RBW, or RNW was prepared incorrectly</b>	Check that the concentrated Buffer RBI, RBW, and RNW were diluted with the correct volume of absolute ethanol.

## Troubleshooting Guide

Facts	Possible Causes	Suggestions
<b>Purified nucleic acid does not perform well in down-stream application</b>	<b>Residual ethanol from Buffer RNW remains in eluate</b>	Care must be taken for eliminating the carry-over Buffer RNW before elution step. The membrane of mini column should be kept completely dry via additional centrifugation (Step 12, page 9) or air-drying.
	<b>Incorrect order of Buffer RBW and RNW</b>	Ensure that Buffer RBW and RNW are used in the correct order during extraction. If used in the wrong order, perform the last washing step with Buffer RNW.

# Ordering Information

Products	Scale	Size	Cat. No.	Type	Products	Scale	Size	Cat. No.	Type
<b>GeneAll® Hybrid-Q™</b> for rapid preparation of plasmid DNA									
Plasmid Rapidprep	mini	50 200	100-150 100-102	spin	Blood SV	mini	100 250	105-101 105-152	spin / vacuum
<b>GeneAll® Exprep™</b> for preparation of plasmid DNA									
Plasmid SV	mini	50 200	101-150 101-102	spin / vacuum	Midi	26 100	105-226 105-201	spin / vacuum	
		26	101-226		MAXI	10 26	105-310 105-326	spin / vacuum	
	Midi	50 100	101-250 101-201	spin / vacuum	Cell SV	mini	100 250	106-101 106-152	spin / vacuum
<b>GeneAll® Exfection™</b> for preparation of transfection-grade plasmid DNA									
Plasmid LE (Low Endotoxin)	mini	50 200	111-150 111-102	spin / vacuum	MAXI	10 26	106-310 106-326	spin / vacuum	
	Midi	26 100	111-226 111-201	spin / vacuum	Clinic SV	mini	100 250	108-101 108-152	spin / vacuum
Plasmid EF (Endotoxin Free)	Midi	20 100	121-220 121-201	spin	Midi	26 100	108-226 108-201	spin / vacuum	
<b>GeneAll® Expi™</b> for purification of fragment DNA									
Gel SV	mini	50 200	102-150 102-102	spin / vacuum	MAXI	10 26	108-310 108-326	spin / vacuum	
PCR SV	mini	50 200	103-150 103-102	spin / vacuum	Genomic DNA micro	50	118-050	spin	
CleanUp SV	mini	50 200	113-150 113-102	spin / vacuum	mini	100 250	117-101 117-152	spin / vacuum	
Combo GP	mini	50 200	112-150 112-102	spin / vacuum	Midi	26 100	117-226 117-201	spin / vacuum	
<b>GeneAll® Exgene™</b> for isolation of total DNA									
Tissue SV	mini	100 250	104-101 104-152	spin / vacuum	MAXI	10 26	117-310 117-326	spin / vacuum	
	Midi	26 100	104-226 104-201	spin / vacuum	Soil DNA mini	mini	50	114-150	spin
	MAXI	10 26	104-310 104-326	spin / vacuum	Stool DNA mini	mini	50	115-150	spin
	mini	100 250	109-101 109-152	spin / vacuum	Stool-Bead DNA mini	mini	50	115-151	spin
Tissue Plus SV	Midi	26 100	109-226 109-201	spin / vacuum	Viral DNA/RNA	mini	50 50	128-150 138-150	spin
	MAXI	10 26	109-310 109-326	spin / vacuum	FFPE Tissue DNA	mini	250	138-152	spin
<b>GeneAll® GenEx™</b> for isolation of total DNA without spin column									
GenEx™ Blood	Sx	100 500	220-101 220-105	solution	GenEx™ Cell	Sx	100 500	221-101 221-105	solution
	Lx	100	220-301			Lx	100	221-301	solution
GenEx™ Tissue	Sx	100 500	222-101 222-105	solution		Lx	100	222-301	solution

Products	Scale	Size	Cat. No.	Type
<b>GeneAll® GenEx™</b> for isolation of total DNA without spin column				
GenEx™ Plant	Sx	100	227-101	solution
	Mx	100	227-201	
	Lx	100	227-301	
GenEx™ Plant Plus	Sx	100	228-101	solution
	Mx	50	228-250	
	Lx	20	228-320	

Products	Scale	Size	Cat. No.	Type
<b>GeneAll® DirEx™ series</b> for preperation of PCR-template without extraction				
DirEx™		100	250-101	solution
DirEx™ Fast-Tissue		96 T	260-011	solution
DirEx™ Fast-Cultured cell		96 T	260-021	solution
DirEx™ Fast-Whole blood		96 T	260-031	solution
DirEx™ Fast-Blood stain		96 T	260-041	solution
DirEx™ Fast-Hair		96 T	260-051	solution
DirEx™ Fast-Buccal swab		96 T	260-061	solution
DirEx™ Fast-Cigarette		96 T	260-071	solution

Products	Scale	Size	Cat. No.	Type
<b>GeneAll® RNA series</b> for preperation of total RNA				
RiboEx™	mini	100	301-001	solution
		200	301-002	
Hybrid-R™	mini	100	305-101	spin
Hybrid-R™ Blood RNA	mini	50	315-150	spin
Hybrid-R™ miRNA	mini	50	325-150	spin
RiboEx™ LS	mini	100	302-001	solution
		200	302-002	
Riboclear™	mini	50	303-150	spin
Riboclear™ Plus	mini	50	313-150	spin
Ribospin™	mini	50	304-150	spin
Ribospin™ II	mini	50	314-150	spin
		300	314-103	
Ribospin™ vRD	mini	50	302-150	spin
Ribospin™ vRD Plus	mini	50	312-150	spin
Ribospin™ vRD II	mini	50	322-150	spin
Ribospin™ Plant	mini	50	307-150	spin
Ribospin™ Seed/Fruit	mini	50	317-150	spin
Ribospin™ Pathogen/TNA	mini	50	314-150	spin
		250	314-152	
Allspin™	mini	50	306-150	spin
RiboSaver™	mini	100	351-001	solution

Products	Scale	Size	Cat. No.	Type
<b>GeneAll® AmpONE™</b> for PCR amplification				
Taq DNA polymerase			250 U	501-025
			500 U	501-050 (2.5 U/ $\mu$ l)
			1,000 U	501-100
Taq Premix		20 $\mu$ l x 96 tubes	526-200	solution
		50 $\mu$ l x 96 tubes	526-500	

Products	Scale	Size	Cat. No.	Type
Taq Master mix		0.5 ml x 2 tubes	541-010	solution
		0.5 ml x 10 tubes	541-050	

Products	Scale	Size	Cat. No.	Type
Reverse Transcriptase		10,000 U	601-100	solution
RT Master mix		0.5 ml x 2 tubes	601-710	solution
One-step RT-PCR Master mix		0.5 ml x 2 tubes	602-110	solution
One-step RT-PCR Premix		20 $\mu$ l x 96 tubes	602-102	solution

Products	Scale	Size	Cat. No.	Type
SYBR qPCR Master mix (2X, Low ROX)		200 rxn	2 ml	solution
		500 rxn	5 ml	
SYBR qPCR Master mix (2X, High ROX)		200 rxn	2 ml	solution
		500 rxn	5 ml	

Products	Scale	Size	Cat. No.	Type
ProteinEx™		100 ml	701-001	solution
Animal cell/tissue				
PAGESTA™				
Reducing		1 ml x 10 tubes	751-001	solution
5X SDS-PAGE Sample Buffer				

Products	Size	Cat. No.	Type
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**GeneAll® GENTi™ 32** Newly designed automated extraction system

Automatic extraction equipment		GTI032A	system
Genomic DNA	48	901-048A	tube
	96	901-096A	plate
Viral DNA/RNA	48	902-048A	tube
	96	902-096A	plate
Blood DNA	48	903-048A	tube
	96	903-096A	plate
Plant DNA/RNA	48	904-048A	tube
	96	904-096A	plate
LMO	48	906-048A	tube
	96	906-096A	plate
Fecal DNA/RNA	48	913-048A	tube
	96	913-096A	plate

**GeneAll® ALLEX® 64** Compact yet Comprehensive automated extraction system

Automatic extraction equipment		AEX064	system
Genomic DNA	48	931-048A	tube
	96	931-096A	plate
Viral DNA/RNA	48	934-048A	tube
	96	934-096A	plate
Blood DNA	48	935-048A	tube
	96	935-096A	plate
Plant DNA/RNA	48	937-048A	tube
	96	937-096A	plate
Fecal DNA/RNA	48	948-048A	tube
	96	948-096A	plate

## **NOTE**

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