Cat. No. 313-150

Size: 50 prep



GeneAll® RiboclearTM plus!

Kit Contents

Components	Quantity	Storage
DNase I	55 ul	-20 ℃
Buffer MS	30 ml	
Buffer RNW	60 ml	
RNase-free water	15 ml	
DNase I buffer (10 X)	1 ml	Room temperature
Micro column type S (with collection tube)	50	
1.5 ml collection tube	50	

Product Specifications

Riboclear [™] plus! Specifications				
Туре	Spin			
Maximum amount of starting samples	~ 100 ul			
RNA recovery rate	~ 95%			
Preparation time	~ 17 minutes			
Maximum loading volume	~ 800 ul			
Minimum elution volume	20 ul			
Binding capacity	~ 100 ug			

Quality Control

RiboclearTM plus! is manufactured in strictly clean condition, and its degree of cleanness is monitored periodically. For consistency of product, the quality certification process is carried out from lot to lot thoroughly and only the qualified is approved to be delivered.

Storage Conditions

Riboclear™ plus! should be stored at room temperature. But prolonged storage at high temperature over 30°C can reduce the performance of the kit. All components are stable for 1 year. Keep out of direct sunlight.

Precautions

The buffers included in Riboclear™ plus! contain irritant which is harmful when in contact with skin or eyes, or when inhaled or swallowed. Care should be taken during handling. Always wear gloves and eye protector, and follow standard safety precautions. In case of contact, wash immediately with plenty of water and seek medical advice

Buffer MS contains chaotropes. It 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 into a RNA preparation. Wear disposable gloves always, because skin often contains bacteria that can be a source of RNase. Use sterile, disposable plasticwares and automatic pipettes reserved for RNA work to prevent cross-contamination with RNase on shared equipment.

GeneAll® Product Information							
GeneAll [®] Hybrid-Q ™ fo	or rapid preparation	n of plasmid DNA		GeneAll [®] GenEx [™] for iso	lation of total DNA		
Products	Format	Size	Cat. No	Products	Format	Size	Cat. No
Plasmid Rapidprep	mini	50	100-150	GenEx™ C	Sx†	100	221-101
Plasmid Rapidprep	mini	200	100-102	GenEx™ C	Sx [†]	500	221-105
GeneAll [®] Exprep [™] for	nrenaration of plas	mid DNA		GenEx™ C	Lx ^{††}	100	221-301
Plasmid SV	mini	50	101-150	GenEx™ T	Sx [†]	100	222-101
Plasmid SV	mini	200	101-102	GenEx™ T	Sx†	500	222-105
Plasmid SV	mini	1,000	101-111	GenEx™ T	Lx ^{††}	100	222-301
Plasmid SV	Midi	26	101-226	GeneAll [®] DirEx [™] Single to	ube DNA extraction	n buffer for PCR	
Plasmid SV	Midi	50	101-250	DirEx™	Solution	50	250-050
Plasmid SV	Midi	100	101-201	GeneAll® RNA Series for	r preparation of F	'NA	
GeneAll [®] Exfection™	for preparation of h	nighly pure plasmi	d DNA	Hybrid-R™	spin	100	305-101
Plasmid LE	mini	50	111-150	•			
Plasmid LE	mini	200	111-102	Hybrid-R™ Blood RNA	spin	50	315-150
Plasmid LE	Midi	26	111-226	Hybrid-R™ miRNA	spin	50	325-150
Plasmid LE	Midi	100	111-201	RiboEx™	solution	100	301-001
Plasmid EF	Midi	20	121-220	RiboEx™	solution	200	301-002
Plasmid EF	Midi	100	121-201	RiboEx™ LS	solution	100	302-001
GeneAll [®] Expin [™] for pu	rification of fragme	ent DNA		RiboEx™ LS	solution	200	302-002
•				Riboclear™	spin	50	303-150
Gel SV Gel SV	mini mini	50 200	102-150 102-102	Riboclear™ plus!	spin	50	313-150
				Ribospin™	spin	50	304-150
PCR SV	mini	50	103-150	Ribospin™ vRD		50	302-150
PCR SV	mini	200	103-102		spin		
CleanUp SV	mini	50	113-150	Ribospin™ vRD plus!	spin	50	312-150
CleanUp SV	mini	200	113-102	Ribospin [™] Plant	spin	50	307-150
Combo GP Combo GP	mini	50	112-150	Allspin™	spin	50	306-150
	mini	200	112-102	GeneAll® AmpONE™ for	PCR amplification	1	
GeneAll [®] Exgene [™] for	isolation of total D	NA		Tag DNA polymerase	(2.5 U/µℓ)	250 U	501-025
Tissue SV (plus!)*	mini	100	104(9)-101	Taq DNA polymerase	(2.5 U/µℓ)	500 U	501-025
Tissue SV (plus!)*	mini	250	104(9)-152	Taq DNA polymerase	(2.5 U/µℓ)	1000 U	501-100
Tissue SV (plus!)**	Midi	26	104(9)-226	α-Taq DNA polymerase	(2.5 U/µℓ)	250 U	502-025
Tissue SV (plus!)**	Midi	100	104(9)-201		(2.5 U/μℓ)	500 U	502-050
Tissue SV (plus!)**	MAXI	10	104(9)-310		(2.5 U/µℓ)	1000 U	502-100
Tissue SV (plus!)**	MAXI	26	104(9)-326	Pfu DNA polymerase	(2.5 U/µℓ)	250 U	503-025
Blood SV	mini	100	105-101	Pfu DNA polymerase	(2.5 U/µℓ)	500 U	503-050
Blood SV	mini	250	105-152	Pfu DNA polymerase	(2.5 U/µℓ)	1000 U	503-100
Blood SV Blood SV	Midi Midi	26 100	105-226 105-201	HS-Taq DNA polymerase	(2.5 U/μℓ)	250 U	531-025
Blood SV	MAXI	10	105-310	HS-Taq DNA polymerase	(2.5 U/µℓ)	500 U	531-050
Blood SV	MAXI	26	105-326	HS-Taq DNA polymerase	(2.5 U/µℓ)	1000 U	531-100
Cell SV	mini	100	106-101	Clean Taq DNA polymerase	(2.5 U/µℓ)	250 U	551-025
Cell SV	mini	250	106-152	Clean Taq DNA polymerase	(2.5 U/µℓ)	500 U	551-050
Cell SV	MAXI	10	106-310	Clean Taq DNA polymerase	(2.5 U/µℓ)	1000 U	551-100
Cell SV	MAXI	26	106-326	Clean &-Taq DNA polymeras	e (2.5 U/µℓ)	250 U	552-025
Clinic SV	mini	100	108-101	Clean α -Taq DNA polymeras		500 U	552-050
Clinic SV	mini	250	108-152	Clean \(\alpha\)-Taq DNA polymeras		1000 U	552-100
Clinic SV	Midi	26	108-226	Tag Premix	96 tubes	20 μℓ	521-200
Clinic SV	Midi	100	108-201	Tag Premix	96 tubes	50 μℓ	521-500
Clinic SV	MAXI	10	108-310	α-Taq Premix	96 tubes	20 μℓ	522-200
Clinic SV	MAXI	26	108-326	α-Tag Premix	96 tubes	50 μl	522-500
Genomic DNA micro	micro	50	118-050	HS-Tag Premix	96 tubes	20 με	525-200
Plant SV	mini	100	117-101	HS-Tag Premix	96 tubes	20 με 20 με	525-500
Plant SV	mini	250	117-152				
Plant SV	Midi	26	117-226		.5 ml x 2 tubes .5 ml x 10 tubes	1 ml	541-010 541-050
Plant SV	Midi	100	117-201	·		5 ml	
Plant SV Plant SV	MAXI MAXI	10 26	117-310 117-326		1.5 ml x 2 tubes	1 ml	542-010
					5 ml x 10 tubes	5 ml	542-050
GMO SV	mini mini	50 200	107-150		.5 ml x 2 tubes	1 ml	545-010
GMO SV	mini	200	107-102		5 ml x 10 tubes	5 ml	545-050
GeneAll [®] GenEx [™] for is	solation of total DN	Α		Taq Premix (w/o dye)	96 tubes	50 μl	524-200
<i>GenEx</i> ™ B	Sx [†]	100	220-101	dNTP mix	2.5 mM each	500 µl	509-020
GenEx™ B	Sx†	500	220-105	dNTP set	100 mM	1 ml x 4 tube	509-040
GenEx™ B	Lx ^{††}	100	220-301	(set of dATP, sCTP,dGTP and dTTP)	1		
* GeneAll® Tissue SV mini, Midi, and MAXI plusl kit provide the additional methods for the purification from animal whole blood. ** GeneAll® SV Midi / MAXI kits require the centrifuge which has a swinging-bucket rotor and ability of 4,000 ~ 5,000 xg.							

† On the basis of DNA purification from 300 $\mu\ell$ whole blood. †† On the basis of DNA purification from 10 ml whole blood

■ Product description

Riboclear[™] plus! provides a convenient method for DNase I treatment and clean-up of total RNA. Riboclear[™] plus! procedures employed the glassfiber membrane technology for the clean-up of total RNA, instead of conventional alcohol precipitation.

In this Riboclear $^{\text{TM}}$ plus! kit, especially, contaminated DNA in extracted total RNA can be removed by DNase I treatment prior to starting the procedure of RNA clean-up.

After the step for removal of contaminated DNA, RNA-containing samples mixed with buffer MS are applied to a micro spin column, followed by centrifugation. RNA binds to silica membrane while most of impurities pass through. The membrane is washed by buffer RNW for removal of some molecules bound nonspecifically. At last, pure RNA is eluted by RNase-free water. RiboclearTM *plus!* procedure should be performed at room temperature. The eluate 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, it is recommended to store at 4°C for immediate analysis or to freeze at -70°C for long-term storage.

■ Protocol of Riboclear[™] plus!

<The procedure for removal of contaminated DNA>

1. Prepare the mixture as below in a 1.5 ml tube.

50 ul RNA eluate

5 ul DNase | buffer (10 X)

1 ul DNase I

If the volume of your sample is more than 50 ul, adjust DNase I buffer to the volume of RNA eluate proportionally.

Also, DNase $\, \, | \, \,$ is sensitive to physical damage.

Therefore, Do NOT vortex vigorously.

- 2. Incubate the mixture at room temperature for 10 minutes.
- Continue with "The procedure of RNA clean-up and concentration"

<The procedure of RNA clean-up and concentration>

 Add 5 volumes of buffer MS to 1 volume of the sample and mix thoroughly.

For 50 ul reaction, add 250 ul of Buffer MS.

- * Do not centrifuge.
- 2. Transfer the mixture to a micro spin column.
- 3. Centrifuge at ≥ 10,000 xg for 30 seconds.

Discard the pass-through and reinsert the micro spin column back into the collection tube.

If the mixture volume is more than 700 ul, apply the mixture twice; apply 700 ul of the mixture, spin down, discard the pass-through, re-insert empty collection tube, and repeat the step again until all of the mixture has been applied to the micro spin column.

- 4. Apply 700 ul of buffer RNW.
- 5. Centrifuge at ≥ 10,000 xg for 30 seconds.

Discard the pass-through and reinsert the micro spin column back into the collection tube.

 Centrifuge at ≥ 10,000 xg for an additional 1 minute to remove residual wash buffer.

Residual ethanol may interfere with downstream reactions. Care must be taken at this step for eliminating the carryover of buffer RNW.

- Transfer the micro spin column to a new 1.5 ml tube (provided).
- Apply 50 ul of RNase-free water to the center of the membrane in the micro spin column. Let it stand for 1 minute, and centrifuge at ≥ 10,000 xg for 1 minute.

To obtain more concentrated RNA solution, apply 20 ul of RNase-free water. The yield can be significantly decreased if the volume of eluent is lower than 20 ul. Purified RNA can be stored at $4\,\mathrm{^{\circ}C}$ for immediate analysis and stored at $-70\,\mathrm{^{\circ}C}$ for long term storage.

■ Troubleshooting Guide

Problem	Possible cause	Suggested solution
Poor quality and yield of RNA	Incorrect procedure	Buffer MS and samples should be mixed completely. Do not centrifuge after mix.
	Improper storage of kit	Store kit components at room temperature. Storage at low temperature may cause salt precipitation. Keep bottles tightly closed in order to avoid evaporation or contamination.
	RNase-free water applied incorrectly	Ensure that RNase-free water is applied to the center of membrane.
	Too much volume of RNase-free water	Reduce the volume of eluent.
Degradation of RNA	Contamination of RNase	RNase can be introduced during use. Be certain not to introduce any RNases during the procedure or later handling. Keep tubes closed whenever possible during the preparation.
	Improper storage of RNA	Store isolated RNA at -70°C, Do not store at -20°C.
Genomic DNA contamination	Starting sample has high DNA mass	At step 1, 1 ul of DNase I can be used for upto 25 ug of DNA contaminants. Increase the DNase I upto 2 ul or decrease the starting sample down to 50 ul.
	DNase I not active	For prolonged activity, aliquot the DNase I into small portion. Do not freezing and thawing the aliquots several times.
RNA does not perform well in downstream application	Residual ethanol remains in eluate	To remove any residual ethanol included in buffer RNW from micro spin column membrane, centrifuge again for complete removal of ethanol (step 6).

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