

Handbook for

PLASMID LE MINI PLASMID LE MIDI PLASMID EF MIDI

Exfection[™] Plasmid Kits Protocol Handbook

For purification of advanced transfection-grade plasmid DNA



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 :

 $GeneAII^{\circledR} \ Exfection^{\intercal M} \ Plasmid \ LE \ mini \ (\ | \ I \ I \ - \ I \ 50, \ | \ I \ I \ - \ I \ 02)$

 $\mathsf{GeneAll}^{\texttt{®}} \; \mathsf{Exfection}^{\mathsf{TM}} \; \mathsf{Plasmid} \; \mathsf{LE} \; \mathsf{Midi} \; (\mathsf{III-226}, \; \mathsf{III-20I})$

GeneAll® Exfection™ Plasmid EF Midi (121-220, 121-220)

Visit www.geneall.com for FAQ, Q&A and more information.

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Kit Contents

	Exfection	™ LE mini	Exfection	™ LE Midi
Cat. No.	111-150	111-102	111-226	111-201
Size	mini	mini	Midi	Midi
No. of preparation	50	200	26	
Column Type Qe (with collection tube)	50	200	-	
Column Type E (clear ring)	_	_	26	
(with collection tube)				
EzClear TM Filter (blue ring)			0.4	
(with collection tube)	-	-	26	
Collection Tube	-	-	26	
Buffer P1	20 ml	60 ml	80 ml	111-226
Buffer P2	20 ml	60 ml	80 ml	× 4
Buffer G3	25 ml	90 ml	110 ml	
Buffer EW1	40 ml	150 ml	150 ml x 2	
Buffer EW2 (concentrate) * †	12 ml	50 ml	40 ml x 2	
Buffer EF **	15 ml	30 ml	60 ml	
RNase A (20 mg/ml)	100 µl	300 µl	400 µl	
Mix Vu [™]	-	-	110 µl	
Protocol Handbook	1	1	1	

	Exfection	M EF Midi
Cat. No.	121-220	121-201
Size	Midi	Midi
No. of preparation	20	
Column Type E (clear ring) (with collection tube)	20	
EzClear TM Filter (blue ring) (with collection tube)	20	
15 ml collection tube	20	
50 ml collection tube	20	
Buffer P1	100 ml	
Buffer P2	100 ml	
Buffer P3	100 ml	121-220
Buffer ER	15 ml	× 5
Buffer EG	150 ml	
Buffer EW 1	250 ml	
Buffer EW2 (concentrate) * †	40 ml x 2	
Buffer EF **	60 ml	
RNase A (20 mg/ml)	500 µl	
Mix Vu [™]	110 µl	
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^{*} Before using for the first time, add absolute ethanol (ACS grade or better) into Buffer EW2 as indicated on the bottle.

[†] Contains sodium azide as a preservative ** 10 mM TrisCl, pH 8.5 endotoxin-free

Precautions and Disclaimer

Exfection™ Plasmid Kits are for research use only, and should not be used for drug, household or other unintended uses. All due care and attention should be taken in every procedure in this handbook. Please consult the Material Safety Data Sheet for information regarding hazards and safe handling practices.

Chemical Hazard

The buffers included in Exfection™ Plasmid Kits contain irritant which is harmful when in contact with skin, or when inhaled or swallowed. Care should be taken during handling. Always wear gloves and eye protector, and follow standard safety precautions.

Some buffers including Buffer G3 and EG contain chaotropic salts. It can form highly reactive compounds when combined with bleach. Do NOT add bleach or acidic solutions directly to the sample-preparation waste.

Quality Control

All components in Exfection™ Plasmid Kits are manufactured in strictly clean condition, and its degree of cleanness is monitored periodically. Restriction enzyme assay, gene cloning, PCR amplification assay, automated sequencing analysis and LAL test as quality control are carried out from lot to lot thoroughly, and only the qualified is delivered.

Storage Condition

Exfection™ Plasmid Kits are shipped at room temperature. All components are stable at room temperature until the date of expiration that is printed on the product label. After the addition of RNase A, Buffer P1 is stable for 1 year when stored at 4°C.

In cold ambient conditions, Buffer P2 and G3 may exhibit salt precipitation and this causes reduction of DNA recover-yields. If so, heat the bottle with occasional swirling in 37°C water bath until completely dissolved.

Buffer P3 can be stored at 4°C. Pre-chilled Buffer P3 may lead to better result.

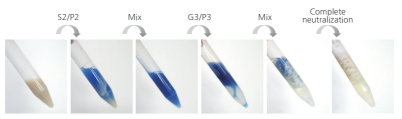
pH Indicator

Mix Vu^{T} is a visual pH indicator that monitors pH shifts during alkaline lysis, ensuring proper mixing to prevent errors. This helps achieve complete neutralization and consistent yield.

Mix Vu^{TM} should be added at a 1:1000 ratio (e.g., $I \mu I$ per I ml suspension). For Exprep Midi or Exfection LE Midi, use 2.5 μI per prep, and for Exfection EF Midi, use 4 μI per prep. Vortex thoroughly after adding Mix Vu^{TM} .

Alternatively, Mix Vu^{TM} can be pre-mixed with SI(PI) Buffer at a 1:1000 ratio (e.g., $80 \,\mu$ l Mix Vu^{TM} into $80 \,\text{ml}$ SI(PI) Buffer or $100 \,\mu$ l into $100 \,\text{ml}$ PI Buffer). Since Mix Vu^{TM} does not fully dissolve in SI(PI) Buffer, shake the buffer before use to resuspend any precipitated particles.

The solution turns blue upon adding S2(P2) Buffer, indicating complete alkalization. Mix thoroughly until the color is uniform. After adding G3(P3) Buffer, the solution becomes colorless, confirming full neutralization. If any color remains, mix further before proceeding.



Product Specifications

	Exfection™ LE	Exfection™ LE	Exfection™ EF
	mini	Midi*	Midi*
Format	Spin/Vacuum	Spin/Vacuum	Spin
Recommended sample volume	5 ml	50 ml	100 ml
Maximum sample volume	10 ml	100 ml	150 ml
Clearing of lysate	Centrifuge	EzClear™	EzClear™
Preparation time	<30 min	<50 min	<70 min
Maximum loading volume	800 µl	15 ml	15 ml
Binding capacity	30 µg	300 µg	300 µg
The level of endotoxins	<10 EU/µg	<10 EU/µg	<0.1 EU/µg
Recovery rate	80~95%	85~95%	75~90%
Minimum elution volume	50 μΙ	500 μl	500 µl

^{*} Exfection™ Plasmid Midi procedure requires a conventional centrifuge which has a swinging-out bucket and ability of 4,000~5,000 x g.

Exfection™ LE and Exfection™ EF GeneAll® I **Plasmid DNA Purification Kit**

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Plasmid DNA Purification Kit Exfection™ LE and Exfection™ EF Plasm

Introduction

ExfectionTM Plasmid Kits provide a simple and fast method for the purification of plasmid DNA with low endotoxin contaminants. Endotoxins (also known as lipopolysaccharides, LPS) are present in the cell membranes of gram-negative bacteria, such as *Escherichia coli*. It is a common contaminant in plasmid preparations and can significantly reduce transfection efficiencies, if not removed during DNA preparations. Endotoxins can be removed with ExfectionTM procedures by two major technologies: advanced phase separation and endotoxin removal washing. The contamination of endotoxin can be reduced to 0.1 EU/ μ g with ExfectionTM Plasmid EF Midi. ExfectionTM Plasmid LE Kits use a new developed wash-buffer which can remove both endotoxins and endonucleases, and these kits can be used for the transfection of most cell-lines.

These kits can be used to isolate and purify any plasmid, but works most efficiently when the plasmid is less than 20,000 bp in size. All process to prepare pure plasmid DNA takes less time than other anion-exchange resin based-kit and simultaneous processing of multiple samples can be easily performed. Up to 30 μ g (300 μ g for Midi) of supercoiled plasmid can be purified using ExfectionTM Plasmid Kits and this pure plasmid DNA is ready for PCR, cloning, fluorescent sequencing, synthesis of labeled hybridization probes, cell transfection, electroporation, enzymatic restriction analysis, and other sensitive applications without further manipulation.

Principle of Method

ExfectionTM Plasmid Kits utilize glass microfiber membrane based on the advanced alkaline lysis method. Alkaline lysis releases plasmid DNA from bacterial cells and degrades RNA, and RNase A removes any survived RNA in the lysate. Cell debris and salt precipitates are removed by centrifugation for mini kit and by EzClearTM Filter for Midi kits. Endotoxins are removed by advanced phase separation technology.

In the presence of high salt, plasmid DNA in cleared lysate binds selectively to glass microfiber membrane in Exfection™ spin column. Bound plasmid DNA is purified in a series of washing steps to eliminate contamination of other bacterial components, such as endotoxins and endonuclease. Finally elution by low salt buffer or deionized water releases plasmid DNA from the glass microfiber membrane. This simple method not only eliminates the need for organic solvent extractions and alcohol precipitation, but also reduce the experiment time of tedious cesium-chloride method or anion-exchange chromatography.

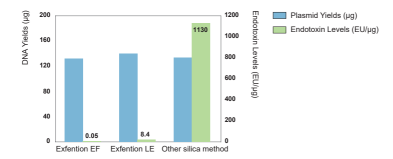
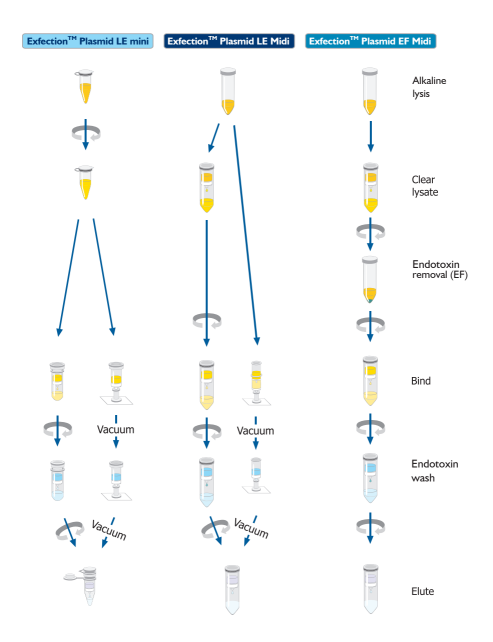


Fig. 1 Comparison of average yields and endotoxin levels in pEGFP-N3 prepared each from 50 ml overnight culture of DH10B by the methods indicated. Endotoxin levels were reduced by more than 10,000 fold using Exfection™ Plasmid EF procedure, compared to other silicabased kit.

Exfection™ Plasmid Procedures



General Considerations

Starting materials

The yield and quality of plasmid DNA depends on several factors such as plasmid copy number, bacterial strain, antibiotics, inoculation and type of culture medium.

Wherever possible, plasmids should be purified from bacterial cultures that have been inoculated with a single transformed colony picked from an agar plate. Usually, the colony is transferred to a small starter culture, which is grown to late log phase. Aliquots of this culture can be used to prepare small amounts of the plasmid DNA for analysis and/or as the innoculum for a large-scale culture. The conditions of growth of the large-scale culture depend chiefly on the copy number of the plasmid and whether it replicates in a stringent or relaxed fashion. At all times, the transformed bacteria should be grown in selective conditions, i.e., in the presence of the appropriate antibiotics.

The copy number of a plasmid is defined as the average number of plasmids per bacterial cells under normal growth conditions. Plasmids have own copy number per cell, depending on their origin of replication (replicon) and the size of plasmid DNA. A plasmid replicon can be defined as the smallest piece of plasmid DNA that is able to replicate autonomously and maintain normal copy number by determining whether they are under relaxed or stringent control. More than 30 different replicons have been identified in plasmids. However, almost all plasmids used routinely in molecular cloning carry a replicon derived from pMB1. pUC plasmids contain a modified pMB1 replicon, have relaxed control, and replicate to a very

Table I. Replicons carried by various plasmid vectors

Plasmid	Size in bp	Copy number	Replicon
pUC series	2,686	500~700	рМВ1
pBluescript series	~3,000	300~500	CoIE1
pGEM series	~3,000	300~400	pMB1
pMK16 and derivatives	~4,500	>1 5	CoIE1
pBR322 and derivatives	4,362	15~20	pMB1
pACYC and derivatives	~4,000	18~22	p15A
pSC101 and derivatives	9,263	~5	pSC101
pRK353 and derivatives	~11,100	~15	R6K

high copy number, otherwise pSCI01 has stringent control and maintain low-copy number. Generally, high-copy number plasmid will result in higher yield. Very large plasmids are often maintained at very low copy numbers per cell.

ExfectionTM Plasmid Kits procedure is optimized to high-copy number plasmid, so larger starting sample may be needed if low-copy number plasmids are used.

Most E.coli strains can be used to propagate and isolate plasmid DNA. Host strains such as DH5 α and XLI-Blue yield DNA of very high-quality. But some strains, particularly those derived from HB101 (e.g. TG I and the IM series), release relatively large amount of carbohydrates when they are lysed. Carbohydrates can inhibit the activity of many restriction enzymes and polymerases, if not completely removed.

Many endA+ strains produce endonuclease I which is encoded in endA and cleaves double-strand DNA. If endonuclease I is not completely removed during DNA preparations, the plasmid DNA in eluate is degraded during subsequent incubation in the presence of Mg²⁺

(e.g. during incubation with restriction enzyme). The wash with Buffer EWI efficiently removes endonucleases from column membrane. Otherwise, this problem can be avoided by use of endA- strains (denoted as endA1) such as DH5 α and XL1-Blue.

Exfection™ Plasmid Kit series is optimized to Luria-Bertani (LB) broth which is the most widely used culture medium for propagation of E.coli. Use of $5 \sim 10$ ml LB culture for mini or $50 \sim 150$ ml LB culture for Midi generally results in good plasmid yields and endotoxin levels. However, the optimal volume of culture to use depends upon the strain, the plasmid, and the density of the culture since the number of bacterial cells can vary greatly between cultures. Too few cells (low cell mass) will result in low DNA yields and may cause a co-filtration of fine precipitates with cleared lysates when clearing of lysate using EzClear™ Filter. Conversely, with too many cells (high cell mass) the bacteria may not lyse efficiently and it causes poor release of the plasmid DNA resulting in a lower yield.

Use of rich broth such as Terrific Broth (TB) or 2xYT will lead to very high cell density. If these media were used, starting sample volume should be reduced not to overload Exfection™ Plasmid Kits column and buffer system. Alternatively, the volume of buffers including P1, P2, P3, and G3 should be increased for efficient lysis. Overnight culture in TB or 2xYT may yield 2~5 times the number of cells compared to cultures grown in LB broth. TB or 2xYT can be used to obtain more yield of plasmid DNA, in case of lowcopy number plasmid.

Harvested bacterial culture is resuspended by Buffer P1 in the presence of RNase A. Exposure of bacterial suspensions to the strongly anionic detergent at high

Alkaline lysis

pH (Buffer P2, SDS/NaOH) opens the cell wall, denatures chromosomal DNA and proteins, and releases plasmid DNA into the supernatant. Although Buffer P2, the alkaline solution, completely disrupts base pairing, the strands of closed circular plasmid DNA are unable to separate from each other because they are topologically intertwined. As long as the intensity and duration of exposure to high pH (OH-) is not too great, the two strands of plasmid DNA fall once again into register when the pH is returned to neutral. However, prolonged exposure to denaturing conditions causes closed circular DNA to enter an irreversibly denatured state. The resulting collapsed coil, which can not be cleaved with restriction enzymes, migrates through agarose gels at about twice the rate of native superhelical closed circular DNA and stains poorly with intercalating dyes.

During lysis, bacterial proteins, broken cell walls, and denatured chromosomal DNA become enmeshed in large complexes that are coated with dodecyl sulfate. These complexes are efficiently precipitated from solution by addition of Buffer G3/P3 which replaces sodium ions by potassium ions.

Vigorous handling of lysate may cause denatured chromosomal DNA to shear, followed by contamination of genomic DNA. It is important for good result that the solution is gently but thoroughly mixed to ensure complete precipitation.

Filtration of lysate with EzClear™ Filter

After alkaline lysis, the cellular debris and precipitates should be removed completely not to clog ExfectionTM Plasmid column in subsequent binding.

New patented EzClearTM Filter facilitates the clearance of the lysate by filtration instead of tedious centrifugation which has been used widely in traditional methods. EzClearTM Filter can be assembled with ExfectionTM Plasmid binding column, and this column stack makes it one-step the clearance of lysate and the binding of plasmid DNA to SV column membrane when using vacuum as driving force. EzClearTM Filter is included only in ExfectionTM Plasmid Midi series.

Removal of endotoxins

Endotoxins (known as lipopolysaccharides, LPS) can be efficiently removed during plasmid preparations by two steps; advanced phase separation and endotoxinremoval washing. After the addition of Buffer ER to the lysate, the mixture is incubated on ice and then at 37°C, resulting in phase-separation of clear upper phase and blue bottom phase. While endotoxins transfer to the lower phase, plasmid DNA locates the upper phase. Because the phase can be dispersed after phase-separating, the mixture should be handled gently during transfer of the upper phase by pipetting. Repetition of this procedure will reduce not only the contamination of endotoxins, but also the yield of plasmid DNA. Generally, the level of endotoxin may be less than 0.1 EU/ μ g after phase-separating and endotoxin-washing.

Washing

When working with endA⁺ strains, endonucleases can be efficiently removed by first washing with Buffer EWI to ensure that plasmid DNA is not degraded during storage or enzyme reactions. Because Buffer EWI not only removes endonucleases but also enhances the quality of plasmid DNA by removal of residual proteins, it is essential especially when working with low-copy plasmids which are generally used with larger culture volume. Buffer EW2 removes the salt and other cellular components bound nonspecifically to column membrane.

Elution

Purified DNA can be eluted in low salt buffer or deionized water as need for downstream applications. Buffer EF contains 10 mM TrisCl, pH 8.5 and is endotoxin-free. When using water as eluent, make sure that it is endotoxin-free and that the pH value is within 7.0 and 8.5. Because plasmid in water is susceptible to hydrolysis and lacks a buffering agent, it is recommended to store at -20°C.

The elution volume can be adjusted as necessity, but it has to be over the minimum requirement to soak completely the column membrane. For higher concentration of DNA, decrease the volume of elution buffer. For higher overall yield, increase the volume of elution buffer and repeat the elution step once again. The concentration and yield as the elution volume is shown below.

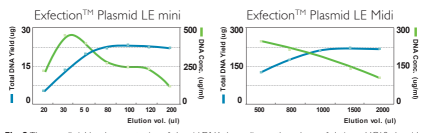


Fig. 2 The overall yield and concentration of plasmid DNA depending on the volume of elution. pUC18 plasmid DNA was purified from 3 ml (mini) and 40 ml (Midi) of overnight cultured DH5 α using ExfectionTM Plasmid LE protocol. Plasmid DNA was eluted with the indicated volume of Buffer EF.

Centrifuge for Midi Kits

Exfection TM Plasmid Midi procedures require the conventional centrifuge which has a swinging-bucket rotor and ability of $4,000 \sim 5,000 \times g$. Use of fixed-angle rotor will cause inconsistent contact of column membrane with sample mixtures and buffers, and lead to unsatisfactory yield. Lower g-force may lead to not only uncomplete removal of ethanol, but also fail of eluting DNA from the membrane of spin column. Available centrifuges and rotors were listed below, but you can employ any equivalent.

Company	Centrifuge	Rotor
Beckman CoulterInc.	Allegra X-15R	Sx4750
(California, USA)	Allegra 25R	Sx4750A
		TS-5.1-500
Eppendorf AG	5804/5804R	A-4-44
(Hamburg, Germany)	5810/5810R	A-1-11
EYELA Inc.	5800	RS-410
(Tokyo, Japan)	5900	RS-410M
Hanil Science	Union 5KR	R-WS1000-6B
Industrial Inc.	Union 55R	W-WS750-6B
(Incheon, Korea)	MF-550	HSR-4S
	HA1000-6	WHSR-4S
	HA1000-3	
Hettich AG	Rotina 35	1717
(Kirchlengern,	Rotanta 460	1724
Germany)	Rotixa 50S	5624

○ Exfection[™] Plasmid LE mini

Before Experiment

Prepare

Before using for the first time, add absolute ethanol (ACS grade or better) into Buffer EW2 as indicated on the bottle.

Add RNase A to Buffer P1 and store it at 4°C.

Prepare 1.5 ml microcentrifuge tubes.

Temperature

All experiment should be performed at room temperature.

Centrifugation

Unless otherwise noted, all centrifugation steps should be performed at full speed ($> 12,000 \times g$ or 12,000 rpm) in a microcentrifuge at room temperature.

Check

Buffer P2 and G3 may precipitate at cold ambient conditions. If precipitate appears, dissolve it in 37°C water bath.

Preparation of Cleared Lysate

I. Pellet the bacterial culture by centrifugation for 5 min at $10,000 \times g$ in a tabletop centrifuge. Discard the supernatant as much as possible.

Use the appropriate volume of bacterial cultures; up to 5 ml for high copy number plasmid, or up to 10 ml for low copy number plasmid. Bacterial culture should be grown for 16 to 24 hours in LB media containing selective antibiotics. Use of other rich broth, such as TB or 2xYT, and/or higher culture volume can cause reduction of lysis efficiency or overload of a SV column, resulting in unsatisfactory yields.

Alternatively, bacterial cells can be pelleted repeatedly in 1.5 ml or 2 ml microcentrifuge tube, by centrifugation for 1 min at full speed.

2. Resuspend the pelleted bacterial cells thoroughly in 250 μ I of Buffer PI. Transfer the suspension to a new I.5 ml microcentrifuge tube.

It is essential to thoroughly resuspend the cell pellet.

* Add RNase A to Buffer P1 before first use.

3. Add 250 μ I of Buffer P2 and mix by inverting the tube 4 times (DO NOT VORTEX).

Incubate until the cell suspension becomes clear and viscous, but DO NOT incubate for more than 5 min. It is important to proceed to next step immediately after the lysate becomes clear without any cloudy clumps.

If precipitated material has formed in Buffer P2 before use, heat to dissolve at 37°C (or above). Precipitated Buffer P2 may cause significant decrease in DNA recover yield.

4. Add 350 μ I of Buffer G3 and immediately mix by inverting the tube 4~6 times (DO NOT VORTEX).

For better precipitation, mix the lysate gently but completely and immediately after addition of Buffer G3.

5. Centrifuge for 10 min at full speed.

Isolation and Purification of Plasmid DNA

6. Transfer carefully the supernatant to a SV column by decanting or pipetting. Centrifuge for 30 sec at full speed. Remove the SV column, discard the pass-through, and re-insert the SV column to the collection tube.

Avoid the white precipitate co-transferring into the SV column.

7. Apply 600 μ l of Buffer EW1 and centrifuge for 30 sec at full speed. Remove the SV column, discard the pass-through, and re-insert the SV column to the collection tube.

This step will remove any traces of lipopolyssacharides (endotoxin), endonucleases, proteins, carbohydrates and other cellular components bound nonspecifically to the column membrane.

- 8. Apply 700 µl of Buffer EW2 and centrifuge for 30 sec at full speed. Remove the SV column, discard the pass-through, and re-insert the SV column to the collection tube.
- 9. Centrifuge for an additional I min to remove residual wash buffer. Transfer the SV column to a new 1.5 ml microcentrifuge tube (not provided).

This step removes residual ethanol from SV column membrane. Residual ethanol in eluate may inhibit subsequent enzymatic reaction. If carryover of Buffer EW2 occurs, centrifuge again for I min before proceeding to next step.

10. Add 50 μ l of Buffer EF or endotoxin-free water, let stand for 1 min, and centrifuge for I min.

Ensure that the Buffer EF or endotoxin-free water is dispensed directly onto the center of SV column membrane for optimal elution of DNA. For the preparation from larger sample volume, elution volume can be increased to 200 μ l maximum. It will increase the total yields of plasmid but decrease the concentration of eluate. For long-term storage, eluting in Buffer EF (10 mM TrisCl, pH 8.5) and storing at -20°C is recommended.

02 Exfection[™] Plasmid LE Midi

Before Experiment

Prepare

Before using for the first time, add absolute ethanol (ACS grade or better) into Buffer EW2 as indicated on the bottle.

Add RNase A to Buffer P1 and store it at 4°C.

Temperature

All experiment should be performed at room temperature.

Centrifugation

Unless otherwise noted, all centrifugation steps should be performed at room temperature in a conventional centrifuge capable of $4,000 \sim 5,000 \times g$, which has a swinging-bucket rotor.

Check

Buffer P2 and G3 may precipitate at cold ambient conditions. If precipitate appears, dissolve it in 37°C water bath.

Preparation of Cleared Lysate

1. Pellet the 50 ml of bacterial culture by centrifugation for 5 min at $10,000 \times g$ in a tabletop centrifuge. Discard the supernatant as much as possible.

Use the appropriate volume of bacterial cultures; up to 100 ml a bacterial culture can be used when $A_{600} < 1.8$. For the small sample less than 50 ml with an $A_{600} < 1.8$, decrease the volume of Buffer P1, P2 and G3 to 2, 2 and 2.8 ml, respectively. If the cell-mass of starting sample is very low, some precipitates can be co-filtered through EzClearTM Filter when clearing of lysate.

Bacterial culture should be grown for 16 to 24 hours in LB-broth containing a selective antibiotics. If other rich broth, such as TB or 2xYT, and/or higher culture volume than 100 ml is used, increase the volume of Buffer P1, P2 and G3 proportionally, since too high cell density of bacterial cells can cause the reduction of lysis efficiency, resulting in unsatisfactory yields.

2. Resuspend the pelleted bacterial cells thoroughly in 2.5 ml of Buffer P1.

It is essential to thoroughly resuspend the cell pellet.

* Add RNase A before first use of the Buffer P1.

3. Add 2.5 ml of Buffer P2 and mix by inverting the tube 4 times (DO NOT VORTEX).

Incubate until the cell suspension becomes clear and viscous, but DO NOT incubate for more than 5 min. It is important to proceed to next step immediately after the lysate becomes clear without any cloudy clumps.

If precipitated material has formed in Buffer P2, heat to dissolve at 37°C (or above). Precipitated Buffer P2 may cause significant decrease in DNA recover yield.

4. Add 3.5 ml of Buffer G3 and thoroughly but gently mix by inverting the tube $5\sim8$ times (DO NOT VORTEX).

For better precipitation and adjustment of binding condition, mix the solution gently but completely and immediately after the addition of Buffer G3.

Incubating on ice may help precipitate the denatured cell components more efficiently; and it may reduce the possibility of the contamination of chromosomal DNA.

Isolation and Purification of Plasmid DNA

In this procedure, one of the two methods can be chosen to purify plasmid DNA. Plasmid DNA can be purified using centrifugation to pull the cleared lysate through the SV column. Alternatively, vacuum can be used to force the cleared lysate through the SV column.

A. Centrifugation Protocol

Always close the cap of the tube at centrifugation.

5. Pour all of the lysate or the cleared lysate into EzClear™ Filter unit (blue ring) sitting on a 50 ml conical collection tube (provided). Incubate for 2 min and centrifuge for 2 min at 1,500 x g (2,800 rpm). Cellular debris will rise to the top during incubation, and this will assist the clearing of lysate through EzClear™ Filter unit. Failure to perform the incubation may lead to incomplete filtration of lysate. A small amount of liquid can remain trapped in the residual insoluble material, but this will not lead to noteworthy decrease in yield.

>> Optional centrifugation before this step

- If the cell mass of the bacterial culture is very dense (A₆₀₀>2.0) and the starting volume is larger than 50 ml, it may be necessary to centrifuge the alkaline lysate before transferring to EzClear[™] Filter. Because too high cell mass can cause the clogging of EzClear[™] Filter.
- This optional centrifugation can be done at 4,500 x g for 20 min on a swinging-bucket rotor or at 10,000 x g for 10 min on a fixed-angle rotor.
- After this optional centrifugation, transfer only the supernatant into EzClear™ Filter. Some debris can be co-transferred.
- 6. Decant carefully the pass-through fraction to SV Midi column (clear ring). Centrifuge for 2 min at 1,500 x g (2,800 rpm). Remove the SV column, discard the pass-through, and re-insert the SV column to the collection tube.

7. Apply 10 ml of Buffer EW1 and centrifuge for 2 min at 1,500 x g (2,800 rpm). Remove the SV column, discard the pass-through, and re-insert the SV column to the collection tube.

This step will remove any traces of lipopolyssacharides (endotoxin), endonucleases, proteins, carbohydrates, and other cellular components bound nonspecifically to the column membrane.

- 8. Apply 10 ml of Buffer EW2 and centrifuge for 2 min at 1,500 x g (2,800 rpm). Remove the SV column, discard the pass-through, and re-insert the SV column to the collection tube.
- 9. Apply 3 ml of Buffer EW2 and centrifuge for 15 min at 4,500 x g (5,000 rpm). Transfer the SV column to a new 50 ml conical tube (not provided).

Care must be taken at the removal of SV Midi column from the collection tube so the SV column does not come into contact with the pass-through fraction, as this will result in carryover of ethanol from Buffer EW2.

Residual ethanol in eluate may interfere with the subsequent reactions. If carryover of ethanol occurs, incubate the SV column for $10\sim20$ min at room temperature to evaporate residual ethanol.

10.Add 0.6 ml of Buffer EF directly onto the center of the SV column membrane. Incubate for 5 min at room temperature and centrifuge for 5 min at $4,500 \times g$ (5,000 rpm).

Ensure that the Buffer EF or distilled water is dispensed directly onto the center of SV column membrane for optimal elution of DNA.

The volume of eluent can be increased or decreased as necessity. Higher volume will decrease the concentration of the eluate but yield slightly more DNA.

The volume can be decreased but it should be 0.5 ml at least, because less volume is insufficient to soak the entire membrane.

For long-term storage, eluting in Buffer EF (10 mM TrisCl, pH 8.5) and storing at -20° C is recommended. Ensure that the pH of water is within 7.0~8.5, when using water for elution.

[]. (Optional:)

- A. For higher concentration of eluate; re-load the eluate from step 10 at page 24 into the SV column membrane, close the cap, incubate 5 min at room temperature, and centrifuge for 5 min at 4,500 x g (5,000 rpm).
- B. For more overall yield; add 0.6~ I ml of fresh Buffer EF into the SV column, close the cap, incubate 5 min at room temperature, and centrifuge for 5 min at $4,500 \times g (5,000 \text{ rpm})$.

The first and second eluates can be combined or collected separately as necessity.

B. Vacuum Protocol

The vacuum pressure should be in this range; 23~26 inHg, 580~660 mmHg, 770~880 mbar or 11~12.5 psi. Lower vacuum pressure may reduce DNA yield and purity, and too high vacuum pressure may cause to burst the column membrane.

5. Assemble a column stack by nesting EzClear™ Midi Filter unit (bluering) into the top of SV Midi column (clear ring). Attach the assembled column stack onto a port of the vacuum manifold tightly.

Most commercial vacuum manifold with luer connectors can be adopted to this protocol.

- 6. Decant all of the lysate to EzClear™ Midi Filter unit and incubate I~3 min to allow the cellular debris and precipitates to rise to the top.
- 7. Apply maximum vacuum to draw the solution through the column stack. When all liquid has been pulled through the SV Midi column at the bottom, slowly release the vacuum.

The lysate will pass through EzClear™ Filter unit and plasmid DNA will bind the membrane in Midi SV column.

If some of the lysate does not pass through the EzClearTM Filter unit, remove the filter unit, place it into a new 50 ml conical tube, and centrifuge for 3 min at 1,750 x g (3,000 rpm). Then apply the pass-through to the Midi SV column.

If the vacuum is released too quickly, the membrane may detach from the SV column. If the membrane becomes detached, tap it down gently with something sterile.

8. Discard the upper EzClear™ Filter unit (blue) and apply 10 ml of Buffer EWI to SV Midi column (clear). Switch on vacuum source to draw the solution through the SV Midi column and slowly release the vacuum.

This step will remove any traces of endotoxin, endonucleses, proteins, carbohydrates, and other cellular components bound nonspecifically to the SV column membrane.

- 9. Apply 14 ml of Buffer EW2 and switch on vacuum source. When all liquid has been pulled through the SV Midi column, slowly release the vacuum.
- 10. Transfer the SV Midi column to the collection tube (provided).
- []. Go to step 9 in 'Centrifugation Protocol' (page 24).

03 Exfection™ Plasmid EF Midi

Before Experiment

Prepare

Before using for the first time, add absolute ethanol (ACS grade or better) into Buffer EW2 as indicated on the bottle.

Add RNase A to Buffer P1 and store it at 4°C.

Chill Buffer P3 for better result. Buffer P3 can be stored at 2~8°C without any precipitation.

Temperature

Prepare ice.

Prepare 37°C water bath or incubator.

All experiment should be performed at room temperature.

Centrifugation

Unless otherwise noted, all centrifugation steps should be performed at room temperature in a conventional centrifuge capable of $4,000 \sim 5,000 \times g$, which has a swinging-bucket rotor.

Check

Buffer P2 and EG may precipitate at cold ambient conditions. If precipitate appears, dissolve it in 37°C water bath.

Preparation of Cleared Lysate

|. Pellet $50\sim100$ ml of bacterial culture by centrifugation for 5 min at $10,000 \times g$ in a tabletop centrifuge. Discard the supernatant as much as possible.

Use the appropriate volume of bacterial cultures; up to 150 ml of a bacterial culture can be used when the A_{600} is less than 1.8. For the bacterial culture with high cell density ($A_{600} > 2.0$), reduce the starting sample volume to 50 ml. Too high cell mass of starting sample can cause the reduction of lysis efficiency and the clogging of the columns, resulting in unsatisfactory yields.

Bacterial culture should be grown for 16 to 24 hours in LB-broth containing selective antibiotics.

2. Resuspend pelleted bacterial cells thoroughly in 4 ml of Buffer P1.

It is essential to thoroughly resuspend the cell pellet.

* Add RNase A before first use of Buffer P1.

3. Add 4 ml of Buffer P2 and mix by inverting the tube 5~6 times (DO NOT VORTEX). Incubate until the cell suspension becomes clear and viscous, but DO NOT incubate for more than 5 min.

It is important to proceed to next step immediately after the lysate becomes clear without any cloudy clumps.

If precipitated material has formed in the bottle of Buffer P2, heat to dissolve at 37°C (or above). Use of precipitated Buffer P2 may cause significant decrease in DNA recover yield.

Keep Buffer P2 capped tightly after use because it can be acidified gradually when in contact with air.

4. Add 4 ml of Buffer P3 and thoroughly but gently mix by inverting the tube 7~8 times (DO NOT VORTEX).

Mix the solution gently but completely and immediately after addition of Buffer P3 for optimal precipitation.

Use of prechilled Buffer P3 or Incubation on ice may help precipitate the denatured cell components more efficiently; and it may reduce the possibility of the contamination of chromosomal DNA.

5. Pour all of the lysate into EzClear™ Filter (blue ring) sitting on a 50 ml conical collection tube (provided). Let it stand for 2 min and centrifuge for 2 min at 1,500 x g (2,800 rpm).

Cellular debris will rise to the top during incubation, and this will assist the clearing of lysate through EzClear[™] Filter. Failure to perform the incubation may lead to incomplete filtration of lysate. A small amount of liquid can remain trapped in the residual insoluble material, but this will not lead to noteworthy decrease in yield.

>> Optional centrifugation before this step

- If the cell mass of the bacterial culture is very dense $(A_{600}>2.0)$ and the starting volume is larger than 100 ml, it may be necessary to centrifuge the lysate before transferring to EzClearTM Filter. Because too high cell mass can cause the clogging of EzClearTM Filter on next step.
- This optional centrifugation can be done at 4,500 x g for 20 min on a swinging-bucket rotor or at 10,000 x g for 10 min on a fixed-angle rotor.
- After this optional centrifugation, transfer the supernatant into EzClear[™] Filter unit (Some debris can be co-transferred).

Removal of Endotoxin

6. Apply 500 μ l of Buffer ER to the filtrate and close the cap of 50 ml conical tube.

The volume of filtrate may be about 10 ml.

7. Vortex to mix and incubate for 15 min on ice.

The mixture will be turbid with vortexing and then become clear during incubation on ice.

8. Incubate for 15 min at 37° C and centrifuge for 2 min at 1,500 x g (2,800 rpm).

After centrifugation, the mixture will be separated by two phases: the white upper phase and the blue (greenish) lower phase. Handle the tube gently not to break up the phase.

9. Transfer carefully the upper phase (clear) into a fresh 15 ml conical tube (provided) by pipetting.

Be careful not to co-transfer the lower phase (greenish blue) since it is liable to be dispersed.

If phase is broken up, repeat centrifugation again.

Isolation and Purification of Plasmid DNA

Always close the cap of the tube at centrifugation.

[0. Add 1/2 volume of Buffer EG to the transfer and invert several times to mix completely.

For 10 ml of the solution, 5 ml of Buffer EG should be added.

- []. Transfer all of the mixture to SV Midi column (clear ring) by decanting or pipetting. Centrifuge for 2 min at 1,500 x g (2,800 rpm). Remove the column, discard the pass-through, and re-insert the column to the collection tube.
- 12. Apply 10 ml of Buffer EW1 and centrifuge for 2 min at 1,500 x g (2,800 rpm). Remove the column, discard the pass-through, and re-insert the column to the collection tube.

This step will remove any traces of lipopolyssacharides (endotoxin), endonucleses, proteins, carbohydrates, and other cellular components bound nonspecifically to the column membrane.

13. Apply 10 ml of Buffer EW2 and centrifuge for 2 min at 1,500 x g (2,800 rpm). Remove the column, discard the pass-through, and re-insert the column to the collection tube.

14. Apply 3 ml of Buffer EW2 and centrifuge for 15 min at 4,500 x g (5,000 rpm). Transfer the column to a new 50 ml conical tube (provided).

Care must be taken at the removal of SV Midi column from the collection tube so the column does not come into contact with the pass-through fraction, as this will result in carryover of ethanol from Buffer EW2.

Residual ethanol in eluate may interfere with the subsequent reactions. If carryover of ethanol occurs, incubate the column for 10~20 min at room temperature to evaporate residual ethanol.

15. Add 0.6 ml of Buffer EF or endotoxin-free water directly onto the center of the column membrane and close the cap. Incubate for 5 min at room temperature and centrifuge for 5 min at $4,500 \times g$ (5,000 rpm).

Ensure that the elution buffer is dispensed directly onto the center of column membrane for optimal elution of DNA.

The volume of eluent can be increased or decreased as necessity. Higher volume will decrease the concentration of the eluate but yield slightly more DNA.

The volume can be decreased but it should be 0.5 ml at least, because less volume is insufficient to soak the entire membrane.

For long-term storage of DNA, other buffers, such as TE (I mM EDTA, I0 mM TrisCl, pH 8.0), can be used for elution. The buffer for elution should be neutral pH (7.0 < pH < 9.0) and low-salt condition.

Ensure that the solution and the plastic-ware for elution is endotoxin-free.

6. (Optional:)

- A. For higher concentration of eluate; re-load the eluate from step 15 into the column membrane, close the cap, incubate 5 min at room temperature, and centrifuge for 5 min at 4,500 x g (5,000 rpm).
- B. For more overall yield; add 0.6~1 ml of fresh Buffer EF into the column, close the cap, incubate 5 min at room temperature, and centrifuge for 5 min at $4,500 \times g (5,000 \text{ rpm})$.

The first and second eluates can be combined or collected separately as necessity.

Troubleshooting Guide

	- ".	
Facts	Possible Causes	Suggestions
Low or no yield of plasmid DNA	Too many cells in sample	Cultures should be grown for $16\sim24$ hours in proper media with antibiotics. Reduce the culture volume of sample. If rich broth such as Terrific Broth (TB) or $2xYT$ is used, starting sample volume must be reduced because these media have very high cell density ($2\sim5$ times to LB).
	Too few cells in sample	Confirm cell density by taking absorbance at 600 nm. Use more cells.
	Low-copy-number plasmid used	Low-copy-number plasmid may yield as little as 0.5 μ g of DNA from a 5 ml overnight culture. Increase the culture volume or use high-copy-number plasmid or rich broth, if possible.
	Too old sample	Streak a fresh plate from a freezer stock. Pick a single colony and prepare a new culture.
	Insufficient antibiotic activity	Confirm that the appropriate amount of fresh antibiotic was present during growth of culture. Most antibiotics are light sensitive and degrade during long-term storage at $2\sim8$ °C. Use fresh antibiotics.
	Poor resuspension of bacterial pellets in Buffer PI	Bacterial cell pellets must be thoroughly resuspended in Buffer P1.
	Buffer P2 precipitated	Redissolve Buffer P2 by warming to 37°C.
	Insufficient digestion with RNase A	Excessive RNA can interfere the binding of plasmid DNA with spin column membrane. If Buffer P1 containing RNase A is more than a year old, the activity of RNase A can be decreased. Add additional RNase A in such a case. (Working concentration = $100 \mu g/ml$)

Facts	Possible Causes	Suggestions
	Inadequate elution buffer	DNA can be eluted only by low salt condition. Buffer EF (10 mM TrisCl, pH 8.5) has the optimal elution efficiency, but other eluent can be engaged as user's need. Elution efficiency is dependent on pH and the maximum efficiency is achieved between 7.0 and 8.5. When using water for elution, make sure the pH value.
	Improper centrifuge (Midi)	Swinging-bucket rotor (capable of 4,000~5,000 x g) must be used. Use of fixed-angle rotor may lead to failure of proper contact between cleared lysate and a column membrane, resulting in poor and inconsistent yield of DNA.
Low purity	Contamination of precipitate when binding	When the cleared lysate is transferred to Exfection TM Plasmid Kits column, ensure that any precipitate does not contain to the transfer.
	Improper centrifuge (Midi)	Swinging-bucket rotor (capable of $4,000 \sim 5,000 \times g$) must be used. Do not use fixed-angle rotor.
Lysate is not clear after filtration (LE Midi)	Precipitation may have occurred	When working with culture volumes less than 50 ml, or with low cell-density cultures ($A_{600} < 1.8$), excessive SDS may be precipitated by substitution of sodium with potassium ions. This fine precipitate can be co-filtered with cleared lysate. When using cultures containing low-cell mass, decrease the buffers for lysis (Buffer P1, P2, G3) as described in the annotation of step 1 at page 21.
The phase is reversed after phase-separating or the color of upper phase is greenish-blue (EF Midi)	The biomass used is too much	Reduce the starting sample. The high density of cleared lysate will reverse the phase.

Facts	Possible Causes	Suggestions
Chromosomal DNA contamination	Mis-handling of the lysate after addition of Buffer G3/P3	Vigorous vortexing after the addition of Buffer G3/P3 can cause shearing of DNA followed by chromosomal DNA contamination. Handle gently the lysate after the addition of Buffer G3/P3. Inverting and rotating tube to cover walls with lysate is sufficient for mixing. But it should be thorough to neutralize the whole lysate.
Smearing of plasmid DNA; additional band behind or ahead of plasmid DNA	Too long lysis time	Too long lysis time in Buffer P2 can cause denaturation of supercoiled plasmid DNA. Proceed to next step immediately after no more clumps are visible in the lysate. Lysis time should not be over 5 minutes in any case. A small amount of this species of DNA is common and is suitable for downstream applications.
	Vigorous mixing in Buffer P2	Vigorous handling after the addition of Buffer P2 can lead to irreversible denaturation of plasmid DNA. Gentle inverting and rotating tube to cover walls with viscous lysate is sufficient for mixing.
RNA Contamination	RNase A omitted or old	RNase A should be added to Buffer P1 before first use. If Buffer P1 containing RNase A is more than a year old, the activity of RNase A can be decreased. Add additional RNase A (working concentration=100 µg/ml). Buffer P1 containing RNase A should be stored at 4°C.
	Too many cells in sample	Reduce the sample volume. Too many cells may not be subjected properly to RNase A digestion.
High salt concentration in eluate	Improper wash step	Ensure that washing steps are performed properly. The column membrane should be completely dried via additional centrifugation or air-drying for good result.

Facts	Possible Causes	Suggestions
DNA floats out of well while loading of agarose gel	Alcohol included in wash buffer is not completely removed during wash steps	Ensure the wash step in protocols. Supplementarily, incubate for 5 minutes at room temperature after applying Buffer EW2 in wash steps.
Enzymatic reaction is not performed well with purified DNA	High salt concentration in eluate	Ensure that washing steps have been carried out just in accordance with the protocols. Repeat of washing steps may help to remove high salt in eluate. Supplementarily, incubate for 5 minutes at room temperature after applying Buffer EW2 in wash steps.
	Prepared plasmid DNA is permanently denatured	Do not allow the lysis reaction in Buffer P2 to proceed longer than 5 minutes. Prolonged alkaline lysis may permanently denature plasmid DNA.
	Residual alcohol in eluate	Ensure that the washing steps have been performed properly. The column membrane should be completely dried via additional centrifugation or air-drying.

Ordering Information

Products	Scale	Size	Cat. No.	Туре	Products	Scale	Size	Cat. No.	Туре	
GeneAll® <i>Hybrid</i>	I-Q TM fo	r rapid p	reparation of	plasmid DNA	GeneAll® Exgene	TM for is	olation c	f total DNA		
S	. 50 100-150			mini	100	105-101	spin /			
Plasmid Rapidprep	mini	200	100-102	spin	_	TTHITH	250	105-152	vacuum	
					Blood SV	Midi	26	105-226	spin /	
GeneAll® Expre p	o TM for pi	reparatio	n of plasmid	DNA	BIOOU 3V	Fildi	100	105-201	vacuum	
		50	101-150	spin /		MAXI	10	105-310	spin /	
r	mini	200	101-102	vacuum		1 1/2/1	26	105-326	vacuum	
Plasmid SV		26	101-226	. ,		mini	100	106-101	spin /	
	Midi	50	101-250	spin / vacuum	Cell SV -		250	106-152	vacuum	
		100	101-201	vacuum	CCII 3V	MAXI	10	106-310	spin /	
GeneAll® <i>Exfec</i> t	ion TM					I I/V	26	106-326	vacuum	
for prepa	ration of	transfect	tion-grade pla	smid DNA		mini	100	108-101	spin /	
		50	111-150	spin /	_		250	108-152	vacuum	
Plasmid LE	mini	200	111-102	vacuum	Clinic SV	Midi	26	108-226	spin /	
(Low Endotoxin)	NA: J:	26	111-226	spin /	Cili lic 5V	1 IIGI	100	108-201	vacuum	
	Midi	100	111-201	vacuum		MAXI	10	108-310	spin /	
Plasmid EF	Midi	20	121-220	cnin		11/04	26	108-326	vacuum	
(Endotoxin Free)	Fildi	100	121-201	spin	Genomic DNA micro)	50	118-050	spin	
						mini	100	117-101	spin /	
GeneAll [®] Expin [™] for p	M for pur	ification	of fragment D	NA	_	11111111	250	117-152	vacuum	
0.107	C 10/		50	102-150	spin /	spin / Plant SV	Midi	26	117-226	spin /
Gel SV	mini	200	102-102	vacuum	Tidilic SV Tildi	1 IIGI	100	117-201	vacuum	
		50	103-150	spin /		MAXI	10	117-310	spin /	
PCR SV	mini	200	103-102	vacuum		1 1/-/	26	117-326	vacuum	
		50	113-150	spin /	Soil DNA mini	mini	50	114-150	spin	
CleanUp SV	mini	200	113-102	vacuum	Stool DNA mini	mini	50	115-150	spin	
		50	112-150	spin /	Stool-Bead DNA mini	mini	50	115-151	spin	
Combo GP	mini	200	112-102	vacuum	Viral DNA/RNA	mini	50	128-150	spin	
					FFPE Tissue DNA	mini	50	138-150	spin	
GeneAll® Exgen	e TM for is	olation o	f total DNA			11111111	250	138-152	shiii	
		100	104-101	spin /	Forensic	mini	100	122-101	- spin / vacuu	
	mini	250	104-152	vacuum	TOTETISIC	11111111	250	122-152	spii i / vacuu	
		26	104-226	spin /	cfDNA	mini	100	129-101	spin / vacuu	
Tissue SV	Midi	100	104-201	vacuum						
		10	104-310	spin /	Companie Cont. I			of total DNA		
	MAXI	26	104-326	vacuum	GeneAll® GenEx ^T	with	nout spin	column		
		100	109-101	spin /		Sx	100	220-101	- solution	
	mini	250	109-152	vacuum	GenEx [™] Blood		500	220-105	Solution	
-		26	109-226	spin /		Lx	100	220-301	solution	
	Midi	100	109-201	vacuum		Sx	100	221-101	- solution	
Tissue Plus SV		10	109-310	spin /	GenEx [™] Cell		500	221-105		
Tissue Plus SV				JUII /		Lx	100	221-301	solution	
Tissue Plus SV	MAXI		109-326				100	221-301	Joidtion	
Tissue Plus SV	MAXI	26	109-326	vacuum			100	222-101		
Tissue Plus SV	MAXI		109-326		GenEx TM Tissue	Sx			- solution	

Products	Scale	Size	Cat. No.	Туре
GeneAll® GenE x		solation out spin	of total DNA column	
	Sx	100	227-101	
GenEx [™] Plant	Mx	100	227-201	solution
	Lx	100	227-301	
	Sx	100	228-101	
GenEx [™] Plant Plus	Mx	50	228-250	solution
	Lx	20	228-320	

GeneAll® DirExTM series
for preperation of PCR-template without extraction

	for preperation of	y i cir-con	ipiace withou	it CAU de don
DirE	< [™]	100	250-101	solution
DirEx	x [™] Fast-Tissue	96 T	260-011	solution
DirEx	x [™] Fast-Cultured cell	96 T	260-021	solution
DirEx	x [™] Fast-Whole blood	96 T	260-03 I	solution
DirEx	x [™] Fast-Blood stain	96 T	260-041	solution
DirE	x [™] <i>Fast-</i> Hair	96 T	260-051	solution
DirEx	x [™] <i>Fast-</i> Buccal swab	96 T	260-061	solution
DirEx	[™] Fast-Cigarette	96 T	260-071	solution

GeneAll® RNA series for preperation of total RNA

		1 1	- 1	
RiboEx [™]	mini	100	301-001	solution
NIDOEX	TTHITH	200	301-002	SOIULION
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
NIDOEX L3	TTHITH	200	302-002	SOIULION
Riboclear™	mini	50	303-150	spin
Riboclear [™] Plus	mini	50	313-150	spin
Ribospin TM	mini	50	304-150	spin
Dile i TM II	mini	50	314-150	onin
Ribospin [™] II	TTHITH	300	314-103	spin
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 TM		50	314-150	onin
Pathogen/TNA	mini	250	314-152	spin
Allspin [™]	mini	50	306-150	spin
RiboSaver™	mini	100	351-001	solution

Products	Scale	Size	Cat. No.	Туре
GeneAll® A mp	ONE TM fo	r PCR ar	mplification	
		250 U	501-025	
Taq DNA polyme	rase	500 U	501-050	(2.5 U/µI)
		1,000 U	501-100	
	20 μl x 9	6 tubes	526-200	1
Taq Premix	50 11 × 9	6 tubes	526-500	solution

GeneAll® AmpMasterTM for PCR amplification

Tag Master mix	0.5 ml x 2 tubes	541-010	solution
iaq i iaster mix	0.5 ml x 10 tubes	541-050	solution

GeneAll® HyperScriptTM for Reverse Transcription

Reverse Transcripta	se 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 μl × 96 tubes	602-102	solution

GeneAll® RealAmp[™] for qPCR amplification

SYBR qPCR Master	200 rxn	2 ml	801-020	solution
mix (2X, Low ROX)	500 rxn	5 ml	801-050	SOIULION
SYBR qPCR Master	200 rxn	2 ml	801-021	and other
mix (2X, High ROX)	500 rxn	5 ml	801-051	solution

GeneAll® Protein series

ProtinEx [™] Animal cell/tissue	100 ml	701-001	solution
PAGESTA [™] Reducing 5X SDS-PAGE Sample Buffer	I ml x 10 tubes	751-001	solution

Products	Size	Cat. No.	Туре
GeneAll® GENTi	M 32 Ne aut	wly designed omated extracti	on system
Automatic extraction equipn	nent	GTI032A	system
Genomic DNA	48	901-048A	tube
Genomic DIVA	96	901-096A	plate
Versi DNIA (DNIA	48	902-048A	tube
Viral DNA/RNA	96	902-096A	plate
DI I DAIA	48	903-048A	tube
Blood DNA	96	903-096A	plate
DI DATA DATA	48	904-048A	tube
Plant DNA/RNA	96	904-096A	plate
1140	48	906-048A	tube
LMO	96	906-096A	plate
	48	913-048A	tube
Fecal DNA/RNA	96	913-096A	plate

GeneAll® AllEx ®64	npact yet Comp omated extracti	
Automatic extraction equipment	AEX064	system

Automatic extraction equipm	ent	AEXU64	system
Carania DNIA	48	931-048	tube
Genomic DNA	96	931-096	plate
\	48	934-048	tube
Viral DNA/RNA	96	934-096	plate
Pland DNIA	48	935-048	tube
Blood DNA	96	935-096	plate
Disast DNIA (DNIA	48	937-048	tube
Plant DNA/RNA	96	937-096	plate
Fecal DNA/RNA	48	948-048	tube
recai DINAYNINA	96	948-096	plate
Forensic	48	936-048	tube
Torchise	96	936-096	plate
Rice DNA	48	949-048	tube
MICE DINA	96	949-096	plate
Meat Genomic DNA	48	950-048	tube
irleat Genomic DINA	96	950-096	plate

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Products	Size	Cat. No.	Туре
Cell/Tissue Total RNA	48	951-048	tube
Celly 1133de Total 14 47	96	951-096	plate
dDNA	48	953-048	tube
CIDINA	96	953-096	plate
GeneAll® AllEx ® V	lini ^{Con}	npact yet Comp omated extracti	rehensive on system
	Con	ob act unt Comb	vahanai va
GeneAll® AllEx®M Automatic extraction equipr	III auto	npact yet Compormated extraction AEX012	on system
Automatic extraction equipr	ment	AEX012	on system system
	III auto	omated extracti	on system
Automatic extraction equipr	ment	AEX012	on system system
Automatic extraction equipr Genomic DNA	ment 48	AEX012 971-048	system system tube
Automatic extraction equipr Genomic DNA Viral DNA/RNA	ment 48	971-048	system tube tube
Automatic extraction equipr Genomic DNA Viral DNA/RNA Blood DNA	ment 48 48 48	971-048 973-048	system system tube tube tube
Automatic extraction equipr Genomic DNA Viral DNA/RNA Blood DNA Plant DNA/RNA	ment 48 48 48 48	971-048 973-048 973-048 974-048	system system tube tube tube tube
Automatic extraction equipr Genomic DNA Viral DNA/RNA Blood DNA Plant DNA/RNA Forensic DNA	48 48 48 48	971-048 972-048 973-048 974-048 975-048	system system tube tube tube tube tube





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