



pGEM[®]-T and pGEM[®]-T Easy Vector Systems

Technical Manual No. 042

INSTRUCTIONS FOR USE OF PRODUCTS A1360, A1380, A3600 and A3610.
PLEASE DISCARD PREVIOUS VERSIONS.
All Technical Literature is Available on the Internet at www.promega.com

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I. Introduction

The pGEM[®](a,b)-T and pGEM[®]-T Easy Vector Systems are convenient systems for the cloning of PCR^(c) products. The vectors are prepared by cutting Promega's pGEM[®]-5Zf(+)^(b) and pGEM[®]-T Easy Vectors with *EcoR* V and adding a 3' terminal thymidine to both ends. These single 3'-T overhangs at the insertion site greatly improve the efficiency of ligation of a PCR product into the plasmids by preventing recircularization of the vector and providing a compatible overhang for PCR products generated by certain thermostable polymerases (1,2). As summarized in Table 1, these polymerases often add a single deoxyadenosine, in a template-independent fashion, to the 3'-ends of the amplified fragments (3,4).

The high copy number pGEM[®]-T and pGEM[®]-T Easy Vectors contain T7 and SP6 RNA Polymerase promoters flanking a multiple cloning site (MCS) within the α -peptide coding region of the enzyme β -galactosidase. Insertional inactivation of the α -peptide allows recombinant clones to be directly identified by color screening on indicator plates. The multiple cloning region of the two vectors includes restriction sites conveniently arranged for use with Promega's Erase-a-Base[®] System (Cat.# E5750) for generating nested sets of deletions.

Both the pGEM[®]-T and pGEM[®]-T Easy Vector contain multiple restriction sites within the MCS. These restriction sites allow for the release of the insert by digestion with a single restriction enzyme. The pGEM[®]-T Easy Vector MCS is flanked by recognition sites for the restriction enzymes *EcoR* I, *BstZ* I and *Not* I, thus providing three single-enzyme digestions for release of the insert, while the pGEM[®]-T Vector cloning site is flanked by recognition sites for the enzyme *BstZ* I. Alternatively, a double-digestion may be used to release the insert from either vector.

The pGEM[®]-T and pGEM[®]-T Easy Vectors also contain the origin of replication of the filamentous phage f1 for the preparation of single-stranded DNA (ssDNA; see Section VII). The ssDNA molecule exported corresponds to the bottom strand shown in Figure 1A and 1B for the pGEM[®]-T and pGEM[®]-T Easy Vectors (nonrecombinant), respectively.

The pGEM[®]-T and pGEM[®]-T Easy Vector Systems now include a 2X Rapid Ligation Buffer for ligation of PCR products. Reactions using this buffer may be incubated for 1 hour at room temperature. The incubation period may be extended to increase the number of colonies after transformation. Generally, an overnight incubation at 4°C will produce the maximum number of transformants.

Table 1. Comparison of PCR Product Properties for Some Thermostable DNA Polymerases.

Characteristic	<i>Taq</i>	<i>Tfi</i>	<i>Tth</i>	<i>Vent</i> _R [®] / <i>Deep</i>	<i>Pfu</i>	<i>Pwo</i>
	<i>Ampli</i> <i>Taq</i> [®]			<i>(Tli)</i>		
Resulting DNA ends	3' A	3' A	3' A	>95% Blunt	>95% Blunt	Blunt N.A.
5'→3' exonuclease activity	Yes	Yes	Yes	No	No	No
3'→5' exonuclease activity	No	No	No	Yes	Yes	Yes

N.A.: not available

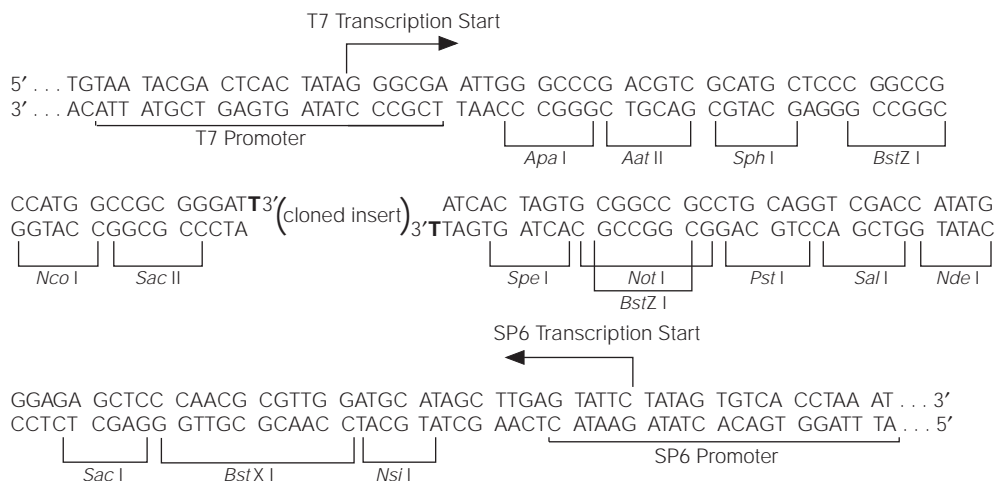
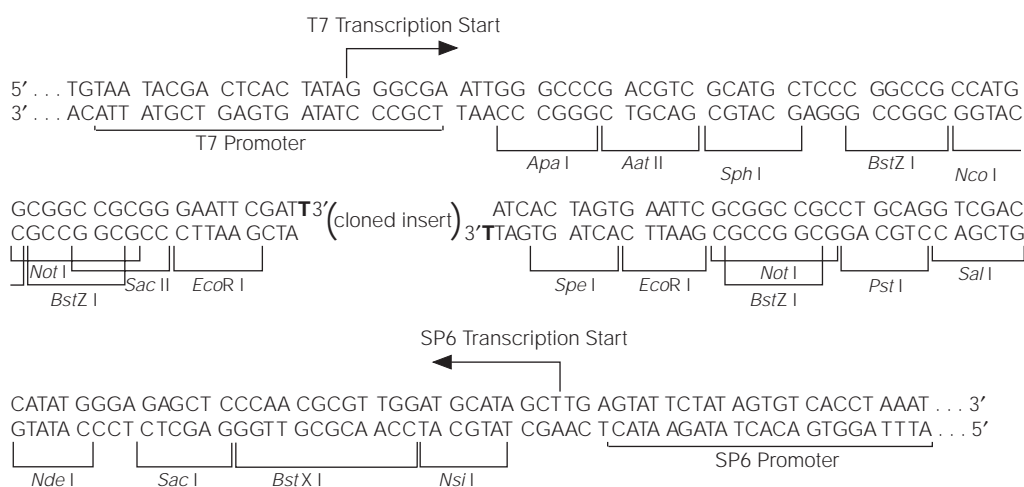
Panel A

Panel B


Figure 1. The promoter and multiple cloning sequence of the pGEM[®]-T (Panel A) and pGEM[®]-T Easy (Panel B) Vectors. The top strand of the sequence shown corresponds to the RNA synthesized by T7 RNA Polymerase. The bottom strand corresponds to the RNA synthesized by SP6 RNA Polymerase.

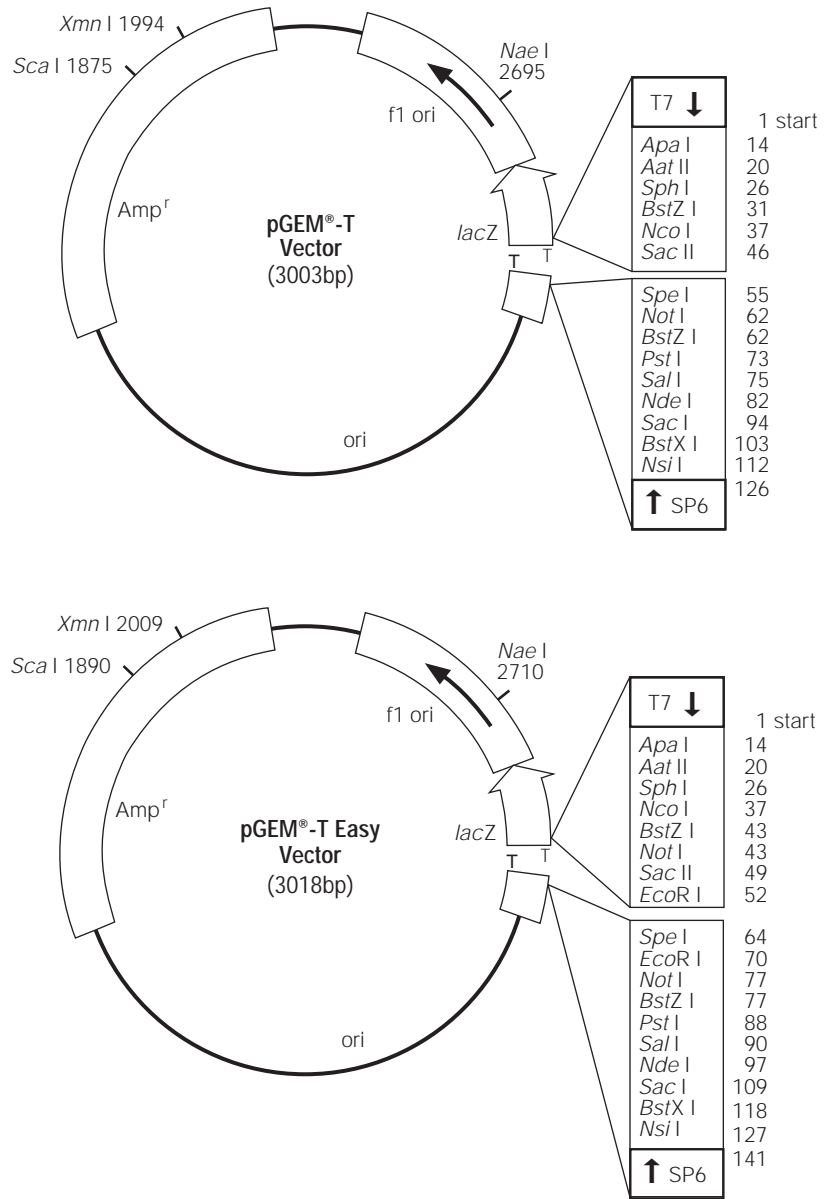


Figure 2. pGEM®-T and pGEM®-T Easy Vector circle maps.

Figure 2 Circle Map Notes:
pGEM®-T Vector Sequence reference points:

T7 RNA Polymerase transcription initiation site	1
SP6 RNA Polymerase transcription initiation site	126
T7 RNA Polymerase promoter	2987-6
SP6 RNA Polymerase promoter	121-143
multiple cloning site	10-113
<i>lacZ</i> start codon	165
<i>lac</i> operon sequences	2824-2984, 151-380
<i>lac</i> operator	185-201
β -lactamase coding region	1322-2182
phage f1 region	2368-2823
binding site of pUC/M13 Forward Sequencing Primer	2944-2960
binding site of pUC/M13 Reverse Sequencing Primer	161-177

pGEM®-T Easy Vector Sequence reference points:

T7 RNA Polymerase transcription initiation site	1
SP6 RNA Polymerase transcription initiation site	141
T7 RNA Polymerase promoter	3002-6
SP6 RNA Polymerase promoter	136-158
multiple cloning site	10-128
<i>lacZ</i> start codon	180
<i>lac</i> operon sequences	2839-2999, 166-395
<i>lac</i> operator	100-216
β -lactamase coding region	1337-2197
phage f1 region	2383-2838
binding site of pUC/M13 Forward Sequencing Primer	2959-2975
binding site of pUC/M13 Reverse Sequencing Primer	176-192

Specialized applications of the pGEM®-T and pGEM®-T Easy Vectors:

1. cloning PCR products
2. used with the Erase-a-Base® System for construction of unidirectional nested deletions
3. ssDNA production
4. blue/white screening for recombinants
5. transcription *in vitro* from dual opposed promoters (For protocol information, please request Promega's *Riboprobe*® *in vitro Transcription Systems*^(d) *Technical Manual #TM016*.)

Use the T7 Promoter Primer or the pUC/M13 Forward Primer to sequence ssDNA produced by the pGEM®-T and pGEM®-T Easy Vectors.

II. Product Components

Product	Size	Cat.#
pGEM [®] -T Vector System I	20 reactions	A3600

Includes:

- 1.2µg pGEM[®]-T Vector (50ng/µl)
- 12µl Control Insert DNA (4ng/µl)
- 100u T4 DNA Ligase
- 200µl 2X Rapid Ligation Buffer, T4 DNA Ligase
- 1 Protocol

Storage Conditions: Store all components at –20°C or –70°C. pGEM[®]-T Vector System I is guaranteed for at least 1 year from date of purchase when stored and handled properly.

Product	Size	Cat.#
pGEM [®] -T Vector System II	20 reactions	A3610

Includes:

- 1.2µg pGEM[®]-T Vector (50ng/µl)
- 12µl Control Insert DNA (4ng/µl)
- 100u T4 DNA Ligase
- 200µl 2X Rapid Ligation Buffer, T4 DNA Ligase
- 1.2ml JM109 Competent Cells, High Efficiency (6 x 200µl)
- 1 Protocol

Storage Conditions: Store the Competent Cells at –70°C. All other components can be stored at –20°C or –70°C. Components of pGEM[®]-T Vector System II are guaranteed for at least 1 year from date of purchase when stored and handled properly, except for JM109 Competent Cells, which are guaranteed until the expiration date that is printed on the tube label.

Product	Size	Cat.#
pGEM [®] -T Easy Vector System I	20 reactions	A1360

Includes:

- 1.2µg pGEM[®]-T Easy Vector (50ng/µl)
- 12µl Control Insert DNA (4ng/µl)
- 100u T4 DNA Ligase
- 200µl 2X Rapid Ligation Buffer, T4 DNA Ligase
- 1 Protocol

Storage Conditions: Store all components at –20°C or –70°C. pGEM[®]-T Easy Vector System I is guaranteed for at least 1 year from date of purchase when stored and handled properly.

Product	Size	Cat.#
pGEM [®] -T Easy Vector System II	20 reactions	A1380

Includes:

- 1.2µg pGEM[®]-T Easy Vector (50ng/µl)
- 12µl Control Insert DNA (4ng/µl)
- 100u T4 DNA Ligase
- 200µl 2X Rapid Ligation Buffer, T4 DNA Ligase
- 1.2ml JM109 Competent Cells, High Efficiency (6 x 200µl)
- 1 Protocol

Storage Conditions: Store the Competent Cells at –70°C. All other components can be stored at –20°C or –70°C. Components of pGEM[®]-T Easy Vector System II are guaranteed for at least 1 year from date of purchase when stored and handled properly, except for JM109 Competent Cells, which are guaranteed until the expiration date that is printed on the tube label.

III. General Considerations

A. PCR Product Purity

An aliquot of the PCR reaction should be analyzed on an agarose gel before using it in the ligation reaction. The PCR product to be ligated can be gel-purified or directly purified using the Wizard® PCR Preps DNA Purification System (Cat.# A7170) or used directly from the reaction. Exposure to shortwave ultraviolet light should be minimized in order to avoid the formation of pyrimidine dimers. If smearing of the PCR product or inappropriate banding is observed on the gel, excise the bands to be cloned from a low-melt agarose gel and purify the DNA with Wizard® PCR Preps or AgarACE® Agarose-Digesting Enzyme (Cat.# M1741). Even if distinct bands of expected size are observed, primer-dimers should be removed. Use Wizard® PCR Preps to purify the bands of interest directly from the reaction mix. Use of crude PCR product may also produce successful ligations in some cases; however, the number of white colonies containing the relevant insert may be reduced due to preferential incorporation of primer-dimers or other extraneous reaction products. Therefore, it may be necessary to screen numerous colonies in order to identify clones that contain the PCR product of interest.

B. Blunt End PCR Products

Thermostable DNA Polymerases with proofreading activity, such as *Pfu* DNA Polymerase^(e) (Cat.# M7741), *Pwo* DNA polymerase and *Tli* DNA Polymerase^(e) (Cat.# M7101) generate blunt-end fragments during PCR amplification. Nevertheless, PCR fragments generated using these polymerases can be modified and ligated to the pGEM®-T and pGEM®-T Easy Vectors (5). Greater than 80% recombinants were obtained (Table 2) when the fragments were purified using the Wizard® PCR Preps DNA Purification System (Cat.# A7170) and A-tailed with *Taq* DNA Polymerase.

The standard A-tailing protocol outlined in Figure 3 requires purification of the PCR product. When working with a large number of different amplified fragments this tailing procedure may prove cumbersome. To determine whether the protocol can be abbreviated in any manner, we have performed several A-tailing reactions using the alternative procedure (Figure 3). In this shortened protocol, the PCR product is not purified and *Taq* DNA Polymerase and dATP are added directly to the 50µl PCR mix after the amplification profile is complete.

Using this alternative procedure, 20-30% recombinant colonies were obtained when *Pfu* DNA polymerase was used in the amplification reaction (Table 2). Recombinants were identified by blue/white screening and were confirmed by restriction digestion analysis. Few positives were observed in the control reaction, in which the PCR fragment was not tailed. These control results confirm that the majority of the pGEM®-T Easy Vector used contains 3'-terminal deoxythymidine. Although the efficiency of this shortened protocol is reduced compared to the standard tailing procedure, the low background exhibited by the pGEM®-T Vectors means that this procedure is a useful way to obtain clones when processing large numbers of samples.

The alternative tailing protocol outlined in Figure 3 cannot be used with all thermostable DNA polymerases. The DNA polymerase used in the amplification is a critical factor. The results in Table 2 show that no recombinants were obtained using the alternative tailing procedure when *Tli* DNA Polymerase was used in the amplification reaction. This may be due to the greater thermostability of *Tli* DNA Polymerase compared with *Pfu* DNA Polymerase (6). However, greater than 90% recombinant colonies were obtained using the standard tailing procedure in conjunction with a *Tli* DNA Polymerase-generated amplification fragment.

Table 2. Comparison of A-Tailing Procedures Used With Different DNA Polymerases.

Polymerase	A-Tailing Procedure	
	Standard	Alternative
<i>Pfu</i> DNA Polymerase	85-90%	20-30%
<i>Tli</i> DNA Polymerase	80-90%	0%

Table 2 lists percent recombinants obtained following PCR amplification with *Pfu* DNA Polymerase and *Tli* DNA Polymerase in combination with either a standard or alternative A-tailing procedure.

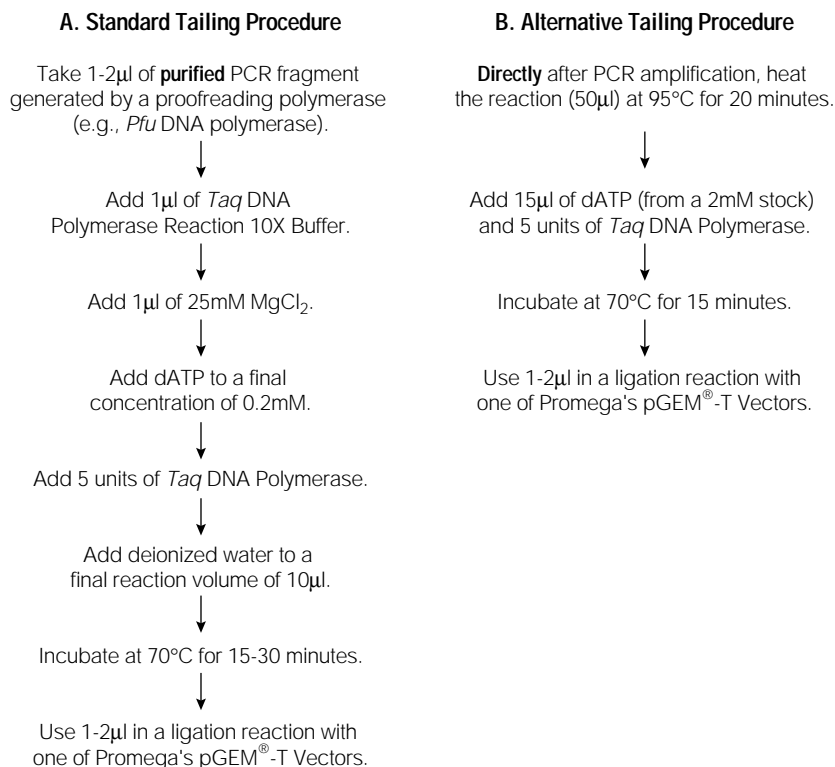


Figure 3. A-Tailing procedures. Panel A: Standard tailing procedure for blunt-end PCR fragments purified with the Wizard[®] PCR Preps DNA Purification System (Cat.# A7170). **Panel B:** Alternative tailing procedure for blunt-end PCR fragments.

C. Optimizing Insert:Vector Molar Ratios

The pGEM[®]-T and pGEM[®]-T Easy Vector Systems have been optimized using a 1:1 molar ratio of the Control Insert DNA to the Vectors. However, ratios of 8:1 to 1:8 have been successfully used. If initial experiments with your PCR product are suboptimal, ratio optimization may be necessary. Ratios from 3:1 to 1:3 provide good initial parameters. The concentration of PCR product should be estimated by comparison to DNA mass standards on a gel or by using a fluorescent assay (5). The pGEM[®]-T and pGEM[®]-T Easy Vectors are approximately 3kb and are supplied at 50ng/ml. To calculate the appropriate amount of PCR product (insert) to include in the ligation reaction, use the following equation.

$$\frac{\text{ng of vector} \times \text{kb size of insert}}{\text{kb size of vector}} \times \text{insert:vector molar ratio} = \text{ng of insert}$$

Sufficient pGEM[®]-T or pGEM[®]-T Easy Vector is provided to vary insert:vector ratios as recommended and to perform control reactions.

Example of insert:vector ratio calculation:

How much 0.5kb PCR product should be added to a ligation in which 50ng of 3.0kb vector will be used if a 3:1 insert:vector molar ratio is desired?

$$\frac{50\text{ng vector} \times 0.5\text{kb insert}}{3.0\text{kb vector}} \times \frac{3}{1} = 25\text{ng insert}$$

Note: Using the same parameters for a 1:1 insert:vector molar ratio, 8.3ng of a 0.5kb insert would be required.

D. Screening Transformants for Inserts

Successful cloning of an insert in the pGEM[®]-T and pGEM[®]-T Easy Vectors interrupts the coding sequence of β -galactosidase; recombinant clones can usually be identified by color screening on indicator plates. However, the characteristics of PCR products cloned into the pGEM[®]-T and pGEM[®]-T Easy Vectors can significantly affect the ratio of blue:white colonies obtained following transformation of competent cells. Clones that contain PCR products, in most cases, produce white colonies, but blue colonies can result from PCR fragments that are cloned in-frame with the *lacZ* gene. Such fragments are usually a multiple of 3 base pairs long (including the 3'-A overhangs), which do not contain in-frame stop codons. It has been reported in the literature that DNA fragments up to 2kb have been cloned in-frame and produced blue colonies.

Even if your PCR product is not a multiple of 3 bases long, the amplification process can introduce mutations (e.g., deletions or point mutations) that may result in blue colonies when competent cells are transformed with the fragment inserted into the pGEM[®]-T or pGEM[®]-T Easy Vectors.

The Control Insert DNA supplied with the pGEM[®]-T and pGEM[®]-T Easy Systems is a 542bp fragment from Promega's pGEM[®]-*luc*^(f) DNA. This sequence has been mutated to contain multiple stop codons in all six reading frames, which ensures a low background of blue colonies for the control reaction. Results obtained with the Control Insert DNA may not be representative of those achieved with your PCR product.

E. Experimental Controls

Promega strongly recommends performing the controls detailed below. These are necessary to accurately assess the performance of the pGEM[®]-T and pGEM[®]-T Easy Vector Systems.

Positive Control

Set up a ligation reaction with the Control Insert DNA as described in the protocol (Section IV.A) and use it for transformations as described in Section V. This control will allow you to determine whether the ligation is proceeding efficiently. Typically, approximately 100 colonies should be observed, 10-30% of which are blue, when competent cells that have a transformation efficiency of 1×10^8 cfu/ μ g DNA are transformed. Greater than 60% of the colonies should be white, and therefore recombinant since the Control Insert DNA is designed to reduce the number of background blue colonies (discussed in Section III.D).

Background blue colonies arise from non-T-tailed or undigested pGEM[®]-T or pGEM[®]-T Easy Vector. These blue colonies are a useful internal transformation control; if no colonies are obtained, the transformation has failed. If blue colonies are obtained, but no whites, the result suggests that the ligation reaction failed. If <50% white colonies are seen in this positive control reaction, then the ligation conditions were probably suboptimal.

The concentration of the Control Insert DNA is such that 2 μ l (4ng/ μ l) can be used in a 10 μ l ligation reaction to achieve a 1:1 molar ratio with 50ng of the pGEM[®]-T or pGEM[®]-T Easy Vectors.

Background Control

Set up a ligation reaction with 50ng of pGEM[®]-T or pGEM[®]-T Easy Vector and no insert as described in the protocol (Section IV.A) and use it for transformations as described in Section V. This ligation will allow determination of the number of background blue colonies resulting from non-T-tailed or undigested pGEM[®]-T or pGEM[®]-T Easy Vector alone. If the recommendations in Section V are followed closely, 10-30 blue colonies will, typically, be observed if the transformation efficiency of the competent cells is 1×10^8 cfu/ μ g DNA. (Under these conditions, cells that are 1×10^7 cfu/ μ g DNA would yield 1-3 blue colonies and cells with a transformation efficiency of 1×10^9 cfu/ μ g DNA would yield 100-300 blue colonies). Compare the number of blue colonies obtained with this background control to the number of blue colonies obtained in the standard reaction using the PCR product. If ligation of the PCR product yields dramatically more blue colonies than the background control reaction, then recombinants are probably among these blue colonies (see Section III.D).

Transformation Control

Check the transformation efficiency of the competent cells by transforming them with an uncut plasmid and calculating cfu/ μ g DNA. If the transformation efficiency is lower than 1×10^8 cfu/ μ g DNA, prepare fresh cells. (Competent cells are available from Promega. See Section X.B.) If you are not using JM109 High Efficiency Competent Cells (provided with pGEM[®]-T and pGEM[®]-T Easy Vector Systems II; Cat.# A3610 and A1380, respectively), be sure the cells are compatible with blue/white screening and standard ampicillin selection, and are at least 1×10^8 cfu/ μ g DNA.

Example of Transformation Efficiency Calculation:

After 100µl competent cells are transformed with 0.1ng uncut plasmid DNA, the transformation reaction is added to 900µl of SOC medium (0.1ng DNA/ml). From that volume, a 1:10 dilution with SOC medium (0.01ng DNA/ml) is made and 100µl plated on two plates (0.001ng DNA/100µl). If 200 colonies are obtained (average of two plates), what is the transformation efficiency?

$$\frac{200\text{cfu}}{0.001\text{ng}} = 2 \times 10^5 \text{ cfu/ng} = 2 \times 10^8 \text{ cfu/}\mu\text{g DNA}$$

IV. Ligations Using the pGEM®-T and pGEM®-T Easy Vectors and the 2X Rapid Ligation Buffer

A. Protocol

1. Briefly centrifuge the pGEM®-T or pGEM®-T Easy Vector and Control Insert DNA tubes to collect contents at the bottom of the tube.
2. Set up ligation reactions as described below. Note: Use 0.5ml tubes known to have low DNA-binding capacity (e.g., VWR Cat.# 20170-310).
3. Vortex the 2X Rapid Ligation Buffer vigorously before each use.

	Standard Reaction	Positive Control	Background Control
2X Rapid Ligation Buffer, T4 DNA Ligase	5µl	5µl	5µl
pGEM®-T or pGEM®-T Easy Vector (50ng)	1µl	1µl	1µl
PCR product	Xµl*	–	–
Control Insert DNA	–	2µl	–
T4 DNA Ligase (3 Weiss units/µl)	<u>1µl</u>	<u>1µl</u>	<u>1µl</u>
deionized water to a final volume of	10µl	10µl	10µl

*Molar ratio of PCR product:vector may require optimization (see Section III.C).

4. Mix the reactions by pipetting. Incubate the reactions 1 hour at room temperature.

Alternatively, if the maximum number of transformants is required, incubate the reactions overnight at 4°C.

Notes:

1. Use only Promega T4 DNA Ligase supplied with this system in performing pGEM®-T and pGEM®-T Easy Vector ligations. Other commercial preparations of T4 DNA ligase may contain exonuclease activities that may remove the terminal thymidines from the vector.
2. 2X Rapid Ligation Buffer contains ATP, which degrades during temperature fluctuations. Avoid multiple freeze-thaw cycles and exposure to frequent temperature changes by making single-use aliquots of the buffer.
3. It is important to vortex the 2X Rapid Ligation Buffer before each use.
4. Longer incubation times will increase the number of transformants. Generally, incubation overnight at 4°C will produce the maximum number of transformants.



This protocol uses the **new 2X Rapid Ligation Buffer**. Use the appropriate volume when setting up the ligation reaction.

V. Transformations Using the pGEM[®]-T and pGEM[®]-T Easy Vector Ligation Reactions

Use high efficiency competent cells (1×10^8 cfu/ μ g DNA) for transformations. The ligation of fragments with a single-base overhang can be inefficient, so it is essential to use cells with a transformation efficiency of 1×10^8 cfu/ μ g DNA (or higher) in order to obtain a reasonable number of colonies (see Section III.E).

We recommend using JM109 High Efficiency Competent Cells (Cat.# L2001); these are provided with the pGEM[®]-T and pGEM[®]-T Easy Vector Systems II. Other host strains may be used, but they should be compatible with blue/white color screening and standard ampicillin selection. JM109 cells should be maintained on M9 minimal medium plates supplemented with thiamine hydrochloride prior to the preparation of competent cells. This selects for the presence of the F' episome, which carries both the *proAB* genes, which complement proline auxotrophy in a host with a (*proAB*) deletion, and *lacI^qZ Δ M15*, which is required in the blue/white color screening process. If you are using competent cells other than JM109 High Efficiency Competent Cells purchased from Promega, it is important that the appropriate transformation protocol be followed. Selection for transformants should be on LB/ampicillin/IPTG/X-Gal plates (see Section X.A). For best results, do not use plates more than 30 days old.

The genotype of JM109 is *recA1*, *endA1*, *gyrA96*, *thi*, *hsdR17* (*r_K-m_K+*), *relA1*, *supE44*, Δ (*lac-proAB*), [F', *traD36*, *proAB*, *lacI^qZ Δ M15*] (8).

A. Protocol

Reagents to Be Supplied by the User

(Solution compositions are provided in Section XI.A.)

- LB plates with ampicillin/IPTG/X-Gal
 - SOC medium
1. Prepare 2 LB/ampicillin/IPTG/X-Gal plates for each ligation reaction, plus two plates for determining transformation efficiency (see Section III.E). Equilibrate the plates to room temperature prior to plating (Step 10).
 2. Centrifuge the tubes containing the ligation reactions to collect contents at the bottom of the tube. Add 2 μ l of each ligation reaction to (a) sterile 1.5ml microcentrifuge tube(s) on ice (see Note 1). Set up another tube on ice with 0.1ng uncut plasmid for determination of the transformation efficiency of the competent cells (see Section III.E).
 3. Remove tube(s) of frozen JM109 High Efficiency Competent Cells from -70°C storage and place in an ice bath until just thawed (about 5 minutes). Mix the cells by **gently** flicking the tube.
 4. **Carefully** transfer 50 μ l of cells into each tube prepared in Step 2 (100 μ l cells for determination of transformation efficiency).
 5. **Gently** flick the tubes to mix and place them on ice for 20 minutes.
 6. Heat-shock the cells for 45-50 seconds in a water bath at exactly 42°C (DO NOT SHAKE).
 7. Immediately return the tubes to ice for 2 minutes.



Avoid excessive pipetting as the cells are very fragile.

8. Add 950 μ l room temperature SOC medium to the tubes containing cells transformed with ligation reactions and 900 μ l to the tube containing cells transformed with uncut plasmid (LB broth may be substituted, but colony number may be lower).
9. Incubate for 1.5 hours at 37°C with shaking (~150rpm).
10. Plate 100 μ l of each transformation culture onto duplicate antibiotic plates. For the transformation control, a 1:10 dilution with SOC medium is recommended for plating. If a higher number of colonies is desired, the cells may be pelleted by centrifugation at 1,000 x *g* for 10 minutes, resuspended in 200 μ l of SOC medium, and 100 μ l plated on each of 2 plates.
11. Incubate the plates overnight (16–24 hours) at 37°C. In our experience, approximately 100 colonies per plate are routinely seen when using competent cells that are 1 x 10⁸ cfu/ μ g DNA, if 100 μ l is plated. Longer incubations or storage of plates at 4°C (after 37°C overnight incubation) may be used to facilitate blue/white screening. White colonies generally contain inserts; however, inserts may also be present in blue colonies. Please see Section III.D for more information.

Notes:

1. In our experience, the use of larger (17 x 100mm) polypropylene tubes (e.g., Falcon Cat.# 2059) has been observed to increase transformation efficiency. Tubes from some manufacturers bind DNA, thereby decreasing the colony number, and should be avoided.
2. Colonies containing β -galactosidase activity may grow poorly relative to cells lacking this activity. After overnight growth, the blue colonies may be smaller than the white colonies which are approximately one millimeter in diameter.

VI. Isolation of Recombinant Plasmid DNA

A standard plasmid miniprep procedure, which takes 30-60 minutes to perform, is described in Promega's Protocols and Applications Guide (9). The miniprep process can be both laborious and time-consuming, particularly when large numbers of minipreps are required. A convenient and reliable method is the Wizard® *Plus* Minipreps DNA Purification System(9).

The Wizard® line of DNA purification products offers five alternatives for plasmid DNA preparation which are distinguished by the scale of the isolation desired (see Table 3). Wizard® *Plus* Miniprep DNA isolations can be completed in 15 minutes and the DNA is ready for other molecular biology applications without prior ethanol precipitations. For best results, a vacuum manifold, such as Promega's Vac-Man® (Cat.# A7231) or Vac-Man® Jr. (Cat.# A7660) Laboratory Vacuum Manifold, should be used to process the minipreps. If a vacuum source is not available, the minipreps may be processed individually using a disposable 3ml Luer-Lok® syringe and a microcentrifuge. A vacuum source and manifold is required for the Wizard® *Plus* Midipreps, Maxipreps, Megapreps and Series 9600™ DNA Purification Systems(h). Refer to Section X.B for ordering information.

Table 3. Wizard® *Plus* Plasmid DNA Purification Systems Selection Guide

Wizard® <i>Plus</i> DNA Purification System	Number of Isolations	Culture Volume	Typical DNA Yields*
Minipreps	50-250	1-3ml	3-10µg
Midipreps	25	10-100ml	10-200µg
Maxipreps	10	100-500ml	300-1,000µg
Megapreps	5	500-1,000ml	700-3,000µg
Series 9600™	8-192	1-5ml	5-20µg

*Typical DNA yields are based on results obtained with high copy number plasmids.

VII. Generation of Single-Stranded DNA from the pGEM®-T and pGEM®-T Easy Vectors

For induction of ssDNA production, bacterial cells containing either the pGEM®-T or pGEM®-T Easy Vector are infected with an appropriate helper phage. The plasmid then enters the f1 replication mode, and the resulting ssDNA is exported as an encapsulated virus-like particle. The ssDNA is purified from the supernatant by simple precipitation and extraction procedures, which are described in detail in Technical Bulletin #TB187, *pGEM® Vector Cloning and Single-Stranded DNA Generation*. For further information, please contact your local Promega Branch Office or Distributor. In the U.S., contact Technical Services at 1-800-356-9526.

VIII. Troubleshooting

Symptom	Possible Cause	Comments
No colonies	A problem has occurred with the transformation reaction or the cells have lost competence	Background undigested vector and religated non-T-tailed vector should yield 10–30 blue colonies, independent of the presence of insert DNA. Check the background control (Section III.E). Use high efficiency competent cells (1×10^8 cfu/ μ g DNA). Test the efficiency of the cells by transforming them with an uncut plasmid that allows for antibiotic selection, such as the pGEM [®] -5Zf(+) Vector. If the recommendations in Section V.A are followed, cells that are 1×10^8 cfu/ μ g DNA typically yield 100 colonies. Therefore, you would not see any colonies from cells that are 1×10^6 cfu/ μ g DNA (Section III.E).
Less than 10% white colonies with Control Insert DNA	Improper dilution of the 2X Rapid Ligation Buffer	The T4 DNA ligase buffer is provided at a concentration of 2X. Use 5μl in a 10μl reaction.
	Ligation reaction has failed	Ligase buffer may have low activity. The 2X Rapid Ligation Buffer contains ATP, which degrades during temperature fluctuations. Avoid multiple freeze-thaw cycles by making single-use aliquots of the buffer. Use a fresh vial of buffer. Ligase activity may be low. To test the activity of the ligase and buffer, set up a ligation with approximately 200ng of DNA markers (e.g., Lambda DNA/ <i>Hind</i> III Markers, Cat.# G1711). Compare ligated and nonligated DNA on a gel and check that the fragments have been religated into high molecular weight material.
	T-overhangs have been removed allowing blunt-end ligation of vector and giving rise to more blue colonies than white colonies	Avoid introduction of nucleases that may degrade the T-overhangs. Use only the T4 DNA Ligase provided with the system, which has been tested for minimal exonuclease activity.

For questions not addressed here, please contact your local Promega branch office or distributor (contact information available at: www.promega.com).

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VIII. Troubleshooting (continued)

Symptom	Possible Cause	Comments
High colony number, but low percentage of white with Control Insert DNA	The competent cells have a high transformation efficiency ($>1 \times 10^9$ cfu/ μ g DNA), and there is a ligation problem	Approximately 1,000 colonies can be obtained with cells that are 1×10^9 cfu/ μ g DNA with 70-90% of those being white. If the ligation is suboptimal or fails, the total number of colonies will be high, but the amount of white colonies will be low or zero. See comments under "Ligation reaction has failed" (above). Also, optimize vector:insert ratio.
Less than 60% white colonies with Control Insert DNA	Improper dilution of the 2X Rapid Ligation Buffer	The T4 DNA ligase buffer is provided at a concentration of 2X. Use 5μl in a 10μl reaction.
	T-overhangs have been removed allowing blunt-end ligation of vector and giving rise to more blue colonies than white colonies	Avoid introduction of nucleases that may degrade the T-overhangs. Use only the T4 DNA Ligase provided with the system, which has been tested for minimal exonuclease activity.
	Ligation temperature is too high	Higher temperatures ($>28^\circ\text{C}$) give rise to increased background and fewer recombinants.
Low number or no white colonies with PCR product	Improper dilution of the 2X Rapid Ligation Buffer	The T4 DNA ligase buffer is provided at a concentration of 2X. Use 5μl in a 10μl reaction.
	Ligation incubation is not long enough	Optimal results are seen with an overnight ligation.
	Failed ligation due to an inhibitory component in the PCR product	Mix some of the PCR product with the control ligation to see if it is inhibiting the reaction. If an inhibitor is suspected, repurify the PCR fragment.
	PCR product is not ligating because there are no 3'-A overhangs	As summarized in Table 1, not all thermostable DNA polymerases create a 3'-A overhang (3,4). Blunt-ended fragments may be subsequently A-tailed by treatment with an appropriate polymerase and dATP (10,11).

VIII. Troubleshooting (continued)

Symptom	Possible Cause	Comments
Low number or no white colonies with PCR product (continued)	PCR product cannot be ligated due to pyrimidine dimers formed from UV overexposure	This is a common problem with gel purified DNA. There is no way to fix this; the DNA must be remade. Exposure to shortwave UV should be limited as much as possible. Use of a glass plate between the gel and UV source will decrease UV overexposure. If possible, only visualize the PCR product using a longwave UV source.
	The PCR fragment is inserted, but it is not disrupting the <i>lacZ</i> gene	If there are a higher number of blue colonies resulting from the PCR fragment ligation than with the background control, some of these blue colonies may contain insert. Screen blue and pale blue colonies (see Section III.D).
	Insert:vector ratio is not optimal	Check the integrity and quantity of your PCR fragment by gel analysis. Optimize the insert:vector ratio (see Section III.C).
	Ligation temperature is too high	Higher temperatures (>28°C) give rise to increased background and fewer recombinants.
	There may be primer-dimers present in PCR fragment preparation	Primer-dimers will ligate into the pGEM®-T or pGEM®-T Easy Vector, but may not be seen after restriction digestion and gel analysis because of their small size. The vector will appear to contain no insert. The PCR fragment should be gel purified.
	Multiple PCR products are generated and cloned into pGEM®-T or pGEM®-T Easy Vector	Gel purify the PCR fragment of interest.
	DNA has rearranged	Check a number of clones to see whether the rearrangement is random. If so, the clone of interest should be present and can be identified by screening several clones. If the same rearrangement is found in all of the clones, use a repair deficient bacterial strain to protect the insert (e.g., SURE® cells), which may reduce recombination events.

For questions not addressed here, please contact your local Promega branch office or distributor (contact information available at: www.promega.com).

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VIII. Troubleshooting (continued)

Symptom	Possible Cause	Comments
PCR product ligation reaction produces white colonies only (no blue colonies are present)	Ampicillin is inactive, allowing ampicillin-sensitive cells to grow	Check that ampicillin plates are made properly and used within 2 weeks. Test ampicillin activity by streaking plates, with and without ampicillin, with an ampicillin-sensitive clone.
	The bacterial strain (e.g., JM109) has lost its F' episome	Check the background control. If these colonies are not blue, the cells may have lost the F' episome (assuming <i>lacI^qZΔM15</i> is located on the F' in the transformed strain and appropriate plates were used). Be sure that the cells are prepared properly for use with this system (see Section V).
	Plates are incompatible with blue/white screening	Check the background control. If these colonies are not blue, check that the plates have ampicillin/IPTG/X-Gal, and that they are fresh. If there is any question about the quality of the plates, repeat plating with fresh plates.
Not enough clones contain the PCR product of interest	Insufficient A-tailing of the PCR fragment	After purification of the PCR fragment, set up an A-tailing reaction (10,11). Clean up the sample and proceed with the protocol.
	Insert:vector ratio is not optimal	Check the integrity and quantity of your PCR fragment by gel analysis. Optimize the insert:vector ratio (see Section III.C).

IX. References

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X. Appendix A: Vector Sequences and Restriction Sites

A. Sequence of the pGEM[®]-T Vector

The sequence supplied below is that of the circular pGEM[®]-5Zf(+) Vector from which the pGEM[®]-T Vector is derived. The pGEM[®]-T Vector has been linearized with *EcoR* V at base 51 of this sequence (indicated by an asterisk) and a T added to both 3'-ends. The added T is not included in this sequence. The sequence shown corresponds to RNA synthesized by T7 RNA Polymerase and is complementary to RNA synthesized by SP6 RNA Polymerase. The strand shown is complementary to the ssDNA produced by this vector. Vector sequences are also available on the Internet at www.promega.com.

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1  GGGCGAATTG  GGCCCGACGT  CGCATGCTCC  CGGCCGCCAT  GGCCGCGGGA
51  T*ATCACTAGT GCGGCCGCCT  GCAGGTCGAC  CATATGGGAG  AGCTCCCAAC
101 GCGTTGGATG  CATAGCTTGA  GTATTCTATA  GTGTCACCTA  AATAGCTTGG
151 CGTAATCATG  GTCATAGCTG  TTTCTGTGT  GAAATTGTTA  TCCGCTCACA
201 ATTCCACACA  ACATACGAGC  CGGAAGCATA  AAGTGTAAG  CCTGGGGTGC
251 CTAATGAGTG  AGCTAACTCA  CATTAATTGC  GTTGCGCTCA  CTGCCCGCTT
301 TCCAGTCGGG  AAACCTGTCG  TGCCAGCTGC  ATTAATGAAT  CGGCCAACGC
351 GCGGGGAGAG  GCGGTTTGCG  TATTGGGCGC  TCTTCCGCTT  CCTCGCTCAC
401 TGACTCGCTG  CGCTCGGTCG  TTCGGCTGCG  GCGAGCGGTA  TCAGCTCACT
451 CAAAGGCGGT  AATACGGTTA  TCCACAGAAT  CAGGGGATAA  CGCAGGAAAG
501 AACATGTGAG  CAAAAGGCCA  GCAAAAGGCC  AGGAACCGTA  AAAAGGCCGC

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551 GTTGCTGGCG TTTTTCATA GGCTCCGCCC CCCTGACGAG CATCACAAAA
601 ATCGACGCTC AAGTCAGAGG TGGCGAAACC CGACAGGACT ATAAAGATAC
651 CAGGCGTTTC CCCCTGGAAG CTCCCTCGTG CGCTCTCCTG TTCCGACCCT
701 GCCGCTTACC GGATACCTGT CCGCCTTTCT CCCTTCGGGA AGCGTGGCGC
751 TTTCTCATAG CTCACGCTGT AGGTATCTCA GTTCGGTGTA GGTGTTTCGC
801 TCCAAGCTGG GCTGTGTGCA CGAACCCCCC GTTCAGCCCC ACCGCTGCGC
851 CTTATCCGGT AACTATCGTC TTGAGTCCAA CCCGGTAAGA CACGACTTAT
901 CGCCACTGGC AGCAGCCACT GGTAACAGGA TTAGCAGAGC GAGGTATGTA
951 GGCGGTGCTA CAGAGTTCTT GAAGTGGTGG CCTAACTACG GCTACACTAG
1001 AAGGACAGTA TTTGGTATCT GCGCTCTGCT GAAGCCAGTT ACCTTCGGAA
1051 AAAGAGTTGG TAGCTCTTGA TCCGGCAAAC AAACCACCGC TGGTAGCGGT
1101 GGTTTTTTTG TTTGCAAGCA GCAGATTACG CGCAGAAAAA AAGGATCTCA
1151 AGAAGATCCT TTGATCTTTT CTACGGGGTC TGACGCTCAG TGGAACGAAA
1201 ACTCACGTTA AGGGATTTTG GTCATGAGAT TATCAAAAAG GATCTTCACC
1251 TAGATCCTTT TAAATTAATA ATGAAGTTTT AAATCAATCT AAAGTATATA
1301 TGAGTAAACT TGGTCTGACA GTTACCAATG CTTAATCAGT GAGGCACCTA
1351 TCTCAGCGAT CTGTCTATTT CGTTCATCCA TAGTTGCCTG ACTCCCCGTC
1401 GTGTAGATAA CTACGATACG GGAGGGCTTA CCATCTGGCC CCAGTGCTGC
1451 AATGATACCG CGAGACCCAC GCTCACCGGC TCCAGATTTA TCAGCAATAA
1501 ACCAGCCAGC CGGAAGGGCC GAGCGCAGAA GTGGTCCTGC AACTTTTATCC
1551 GCCTCCATCC AGTCTATTAA TTGTTGCCGG GAAGCTAGAG TAAGTAGTTC
1601 GCCAGTTAAT AGTTTGCGCA ACGTTGTTGG CATTGCTACA GGCATCGTGG
1651 TGTCACGCTC GTCGTTTGGT ATGGCTTCAT TCAGCTCCGG TTCCAACGA
1701 TCAAGGCGAG TTACATGATC CCCCATGTTG TGCAAAAAG CGGTTAGCTC
1751 CTTCCGGTCT CCGATCGTTG TCAGAAGTAA GTTGGCCGCA GTGTTATCAC
1801 TCATGGTTAT GGCAGCACTG CATAATTCTC TTAAGTGCAT GCCATCCGTA
1851 AGATGCTTTT CTGTGACTGG TGAGTACTCA ACCAAGTCAT TCTGAGAATA
1901 GTGTATGCGG CGACCGAGTT GCTCTTGCCC GGCGTCAATA CGGGATAATA
1951 CCGCGCCACA TAGCAGAACT TTAAAAGTGC TCATCATTGG AAAACGTTCT
2001 TCGGGGCGAA AACTCTCAAG GATCTTACCG CTGTTGAGAT CCAGTTCGAT
2051 GTAACCCACT CGTGCACCCA ACTGATCTTC AGCATCTTTT ACTTTCACCA
2101 GCGTTTCTGG GTGAGCAAAA ACAGGAAGGC AAAATGCCGC AAAAAAGGGA
2151 ATAAGGGCGA CACGGAAATG TTGAATACTC AACTCTTCC TTTTCAATA
2201 TTATTGAAGC ATTTATCAGG GTTATTGTCT CATGAGCGGA TACATATTTG
2251 AATGTATTTA GAAAAATAAA CAAATAGGGG TTCCGCGCAC ATTTCCCCGA
2301 AAAGTGCCAC CTGTATGCGG TGTGAAATAC CGCACAGATG CGTAAGGAGA
2351 AAATACCGCA TCAGGCGAAA TTGTAAACGT TAATATTTTG TTAAAATTTCG
2401 CGTTAAATAT TTGTAAATC AGCTCATTTT TTAACCAATA GGCCGAAATC
2451 GGCAAAATCC CTTATAAATC AAAAGAATAG ACCGAGATAG GGTGAGTGT

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2501 TGTTCCAGTT TGGAACAAGA GTCCACTATT AAAGAACGTG GACTCCAACG
 2551 TCAAAGGGCG AAAAACCGTC TATCAGGGCG ATGGCCCACT ACGTGAACCA
 2601 TCACCCAAAT CAAGTTTTTTT GCGGTCGAGG TGCCGTAAAG CTCTAAATCG
 2651 GAACCCTAAA GGGAGCCCC GATTTAGAGC TTGACGGGGA AAGCCGGCGA
 2701 ACGTGGCGAG AAAGGAAGGG AAGAAAGCGA AAGGAGCGGG CGCTAGGGCG
 2751 CTGGCAAGTG TAGCGGTCAC GCTGCGCGTA ACCACCACAC CCGCCGCGCT
 2801 TAATGCGCCG CTACAGGGCG CGTCCATTCG CCATTCAGGC TCGCAACTG
 2851 TTGGGAAGGG CGATCGGTGC GGGCCTCTTC GCTATTACGC CAGCTGGCGA
 2901 AAGGGGGATG TGCTGCAAGG CGATTAAGTT GGGTAACGCC AGGGTTTTTC
 2951 CAGTCACGAC GTTGTAAAAC GACGGCCAGT GAATTGTAAT ACGACTCACT
 3001 ATA

B. pGEM[®]-T Vector Restriction Sites

The following restriction enzyme tables are based on those of the circular pGEM[®]-5Zf(+) Vector from which the pGEM[®]-T Vector is derived. The pGEM[®]-T Vector has been linearized at base 51 with *EcoR* V and a T added to both 3'-ends. This site will not be recovered upon ligation of the vector and insert. The tables were constructed using DNASTAR[®] sequence analysis software. Please note that we have not verified this information by restriction digestion with each enzyme listed. The location given specifies the 3'-end of the cut DNA (the base to the left of the cut site). Please contact your local Promega Branch Office or Distributor if you identify a discrepancy. In the U.S., contact Technical Services at 1-800-356-9526.

Table 4. Restriction Enzymes That Cut the pGEM[®]-T Vector Between 1 and 5 Times.

Enzyme	# of Sites	Location	Enzyme	# of Sites	Location
Aat II	1	20	BstZ I	2	31, 62
Acc I	1	76	Cfr10 I	2	1475, 2693
Acy I	2	17, 1932	Dde I	4	777, 1186, 1352, 1892
Afl III	2	99, 502	Dra I	3	1261, 1280, 1972
Alw26 I	2	1456, 2232	Dra III	1	2592
Alw44 I	2	816, 2062	Drd I	2	610, 2547
AlwNI	1	918	Dsa I	2	37, 43
Apa I	1	14	Eag I	2	31, 62
AspH I	4	94, 820, 1981, 2066	Ear I	3	386, 2190, 2881
Ava II	2	1533, 1755	EcIHK I	1	1395
Ban I	3	246, 1343, 2629	Eco52 I	2	31, 62
Ban II	3	14, 94, 2667	EcoI CR I	1	92
Bbu I	1	26	EcoR V	1	51 (see above)
Bgl I	3	39, 1515, 2836	Fok I	5	119, 1361, 1542, 1829, 2919
Bsa I	1	1456	Fsp I	2	1617, 2843
BsaA I	1	2592	Hae II	4	380, 750, 2743, 2751
BsaH I	2	17, 1932	Hga I	4	613, 1191, 1921, 2809
BsaJ I	5	37, 43, 241, 662, 2939	Hinc II	1	77
Bsp120 I	1	10	Hind II	1	77
BspH I	2	1222, 2230	Hsp92 I	2	17, 1932
BspM I	1	62	Ksp I	1	46
BssS I	2	675, 2059	Mae I	5	56, 997, 1250, 1585, 2743
BstO I	5	242, 530, 651, 664, 2940			
BstX I	1	103			

Enzyme	# of Sites	Location	Enzyme	#of Sites	Location
Mlu I	1	99	Sac II	1	46
Nae I	1	2695	Sal I	1	75
Nci I	4	30, 882, 1578, 1929	Sca I	1	1875
Nco I	1	37	Sfi I	1	39
Nde I	1	82	Sin I	2	1533, 1755
NgoM I	1	2693	Spe I	1	55
Not I	1	62	Sph I	1	26
Nsi I	1	112	Sse8387 I	1	73
Nsp I	2	26, 506	Ssp I	3	2199, 2384, 2408
Ppu10 I	1	108	Sty I	1	37
Pst I	1	73	Taq I	4	76, 602, 2046, 2625
Pvu I	2	1765, 2864	Tfi I	2	337, 477
Pvu II	2	326, 2893	Vsp I	3	273, 332, 1567
Rsa I	1	1875	Xmn I	1	1994
Sac I	1	94			

Note: The enzymes listed in boldface type are available from Promega.

Table 5. Restriction Enzymes That Do Not Cut the pGEM®-T Vector.

Acc III	<i>Bbs I</i>	<i>Bst1107 I</i>	EcoR I	<i>PaeR7 I</i>	Sma I
Acc65 I	Bcl I	Bst98 I	<i>Ehe I</i>	<i>PflM I</i>	SnaB I
AccB7 I	Bgl II	BstE II	<i>Fse I</i>	<i>PinA I</i>	<i>Spl I</i>
<i>Afl II</i>	<i>Blp I</i>	Bsu36 I	Hind III	<i>Pme I</i>	<i>Srf I</i>
Age I	<i>Bpu1102 I</i>	Cla I	Hpa I	<i>Pml I</i>	Stu I
<i>Asc I</i>	<i>BsaB I</i>	Csp I	I-Ppo I	<i>PpuM I</i>	<i>Swa I</i>
Ava I	BsaM I	Csp45 I	<i>Kas I</i>	<i>PshA I</i>	Tth111 I
<i>Avr II</i>	<i>BsiW I</i>	<i>Dra II</i>	Kpn I	<i>Psp5 II</i>	Xba I
Bal I	<i>Bsm I</i>	Eco47 III	Nar I	<i>PspA I</i>	<i>Xcm I</i>
BamH I	BsrBr I	<i>EcoN I</i>	Nhe I	<i>Rsr II</i>	Xho I
<i>Bbe I</i>	<i>BsrG I</i>	<i>Eco72 I</i>	Nru I	Sgf I⁽ⁱ⁾	Xma I
<i>BbrP I</i>	BssH II	<i>Eco81 I</i>	<i>Pac I</i>	<i>SgrA I</i>	

Note: The enzymes listed in boldface type are available from Promega.

Table 6. Restriction Enzymes That Cut the pGEM®-T Vector 6 or More Times.

<i>Aci I</i>	Bst71 I	Hae III	<i>Mae II</i>	MspA1 I	<i>SfaN I</i>
Alu I	<i>BstU I</i>	Hha I	<i>Mae III</i>	<i>Nla III</i>	Tru9 I
<i>Bbv I</i>	Cfo I	Hinf I	Mbo I	<i>Nla IV</i>	Xho II
BsaO I	Dpn I	<i>HinP I</i>	Mbo II	<i>Ple I</i>	
Bsp1286 I	<i>Dpn II</i>	Hpa II	<i>Mnl I</i>	Sau3A I	
<i>Bsr I</i>	<i>Eae I</i>	<i>Hph I</i>	<i>Mse I</i>	Sau96 I	
BsrS I	<i>Fnu4H I</i>	Hsp92 II	Msp I	<i>ScrF I</i>	

Note: The enzymes listed in boldface type are available from Promega.

C. Sequence of the pGEM[®]-T Easy Vector

The pGEM[®]-T Easy Vector has been linearized with *EcoR* V at base 60 of this sequence (indicated by an asterisk) and a T added to both 3'-ends. The added T is not included in this sequence. The sequence shown corresponds to RNA synthesized by T7 RNA Polymerase and is complementary to RNA synthesized by SP6 RNA Polymerase. The strand shown is complementary to the ssDNA produced by this vector. Vector sequences are also available on the Internet at www.promega.com.

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1  GGGCGAATTG  GGCCCGACGT  CGCATGCTCC  CGGCCGCCAT  GGCGGCCGCG
51  GGAATTCGAT* ATCACTAGTG  AATTCGCGGC  CGCCTGCAGG  TCGACCATAT
101 GGGAGAGCTC  CCAACGCGTT  GGATGCATAG  CTTGAGTATT  CTATAGTGTC
151 ACCTAAATAG  CTTGGCGTAA  TCATGGTCAT  AGCTGTTTCC  TGTGTGAAAT
201 TGTTATCCGC  TCACAATTCC  ACACAACATA  CGAGCCGAA  GCATAAAGTG
251 TAAAGCCTGG  GGTGCCTAAT  GAGTGAGCTA  ACTCACATTA  ATTGCGTTGC
301 GCTCACTGCC  CGCTTTCAG  TCGGGAAACC  TGTCGTGCCA  GCTGCATTAA
351 TGAATCGGCC  AACGCGCGGG  GAGAGGCGGT  TTGCGTATTG  GGCGCTCTTC
401 CGCTTCCTCG  CTCACTGACT  CGCTGCGCTC  GGTGTTCCGG  CTGCGGCGAG
451 CGGTATCAGC  TCACTCAAAG  GCGGTAATAC  GGTATCCAC  AGAATCAGGG
501 GATAACGCAG  GAAAGAACAT  GTGAGCAAAA  GGCCAGCAA  AGGCCAGGAA
551 CCGTAAAAAG  GCCGCGTTGC  TGGCGTTTTT  CCATAGGCTC  CGCCCCCTG
601 ACGAGCATCA  CAAAATCGA  CGCTCAAGTC  AGAGGTGGCG  AAACCCGACA
651 GGACTATAAA  GATACCAGGC  GTTTCCCCCT  GGAAGCTCCC  TCGTGCGCTC
701 TCCTGTTCCG  ACCCTGCCGC  TTACCGGATA  CCTGTCCGCC  TTTCTCCCTT
751 CGGGAAGCGT  GGCGCTTCT  CATAGCTCAC  GCTGTAGGTA  TCTCAGTTCG
801 GTGTAGGTCG  TTCGCTCAA  GCTGGGCTGT  GTGCACGAAC  CCCCCTTCA
851 GCCCGACCGC  TGCGCCTTAT  CCGGTAACTA  TCGTCTTGAG  TCCAACCCGG
901 TAAGACACGA  CTTATCGCCA  CTGGCAGCAG  CCACTGGTAA  CAGGATTAGC
951 AGAGCGAGGT  ATGTAGGCGG  TGCTACAGAG  TTCTTGAAGT  GGTGGCCTAA
1001 CTACGGCTAC  ACTAGAAGGA  CAGTATTTGG  TATCTGCGCT  CTGCTGAAGC
1051 CAGTTACCTT  CGGAAAAAGA  GTTGGTAGCT  CTTGATCCGG  CAAACAAACC
1101 ACCGCTGGTA  GCGGTGGTTT  TTTTGTGTTG  AAGCAGCAGA  TTACGCGCAG
1151 AAAAAAAGGA  TCTCAAGAAG  ATCCTTTGAT  CTTTTCTACG  GGGTCTGACG
1201 CTCAGTGGAA  CGAAACTCA  CGTTAAGGGA  TTTTGGTCAT  GAGATTATCA
1251 AAAAGGATCT  TCACCTAGAT  CCTTTTAAAT  TAAAAATGAA  GTTTTAAATC
1301 AATCTAAAGT  ATATATGAGT  AACTTGGTC  TGACAGTTAC  CAATGCTTAA
1351 TCAGTGAGGC  ACCTATCTCA  GCGATCTGTC  TATTTGTTTC  ATCCATAGTT
1401 GCCTGACTCC  CCGTCGTGTA  GATAACTACG  ATACGGGAGG  GCTTACCATC
1451 TGGCCCCAGT  GCTGCAATGA  TACCGCGAGA  CCCACGCTCA  CCGGCTCCAG
1501 ATTTATCAGC  AATAAACAG  CCAGCCGGAA  GGGCCGAGCG  CAGAAGTGGT
1551 CCTGCAACTT  TATCCGCTC  CATCCAGTCT  ATTAATTGTT  GCCGGAAGC
1601 TAGAGTAAGT  AGTTCGCCAG  TTAATAGTTT  GCGCAACGTT  GTTGGCATTG

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1651 CTACAGGCAT CGTGGTGTCA CGCTCGTCGT TTGGTATGGC TTCATTCAGC
1701 TCCGGTTCCC AACGATCAAG GCGAGTTACA TGATCCCCCA TGTTGTGCAA
1751 AAAAGCGGTT AGCTCCTTCG GTCCTCCGAT CGTTGTGAGA AGTAAGTTGG
1801 CCGCAGTGTT ATCACTCATG GTTATGGCAG CACTGCATAA TTCTCTTACT
1851 GTCATGCCAT CCGTAAGATG CTTTTCTGTG ACTGGTGAGT ACTCAACCAA
1901 GTCATTCTGA GAATAGTGTA TCGGGCGACC GAGTTGCTCT TGCCCGGCGT
1951 CAATACGGGA TAATACCGCG CCACATAGCA GAACTTTAAA AGTGCTCATC
2001 ATTGAAAAC GTTCTTCGGG GCGAAAAC TC AAGGATCT TACCGCTGTT
2051 GAGATCCAGT TCGATGTAAC CCACTCGTGC ACCCAACTGA TCTTCAGCAT
2101 CTTTTACTTT CACCAGCGTT TCTGGGTGAG CAAAAACAGG AAGGCAAAAT
2151 GCCGAAAAA AGGGAATAAG GGCGACACGG AAATGTTGAA TACTCATACT
2201 CTTCTTTTTT CAATATTATT GAAGCATTTA TCAGGGTTAT TGTCTCATGA
2251 GCGGATACAT ATTTGAATGT ATTTAGAAAA ATAAACAAAT AGGGGTTCCG
2301 CGCACATTC CCCGAAAAGT GCCACCTGTA TGCGGTGTGA AATACCGCAC
2351 AGATGCGTAA GGAGAAAATA CCGCATCAGG CGAAATTGTA AACGTTAATA
2401 TTTTGTTAAA ATTCGCGTTA AATATTTGTT AAATCAGCTC ATTTTTTAAC
2451 CAATAGGCCG AAATCGGCAA AATCCCTTAT AAATCAAAAG AATAGACCGA
2501 GATAGGGTTG AGTGTGTTC CAGTTTGAA CAAGAGTCCA CTATTAAAGA
2551 ACGTGGACTC CAACGTCAA GGGCGAAAA CCGTCTATCA GGGCGATGGC
2601 CCACTACGTG AACCATCACC CAAATCAAGT TTTTTCGGT CGAGGTGCCG
2651 TAAAGCTCTA AATCGGAACC CTAAAGGGAG CCCCAGATT AGAGCTTGAC
2701 GGGGAAAGCC GGCGAACGTG GCGAGAAAGG AAGGGAAGAA AGCGAAAGGA
2751 GCGGGCGCTA GGGCGCTGGC AAGTGTAGCG GTCACGCTGC GCGTAACCAC
2801 CACACCCGCC GCGCTTAATG CGCCGCTACA GGGCGCGTCC ATTCGCCATT
2851 CAGGCTGCGC AACTGTTGGG AAGGGCGATC GGTGCGGGCC TCTTCGCTAT
2901 TACGCCAGCT GGCGAAAGG GGATGTGCTG CAAGGCGATT AAGTTGGGTA
2951 ACGCCAGGT TTTCCAGTC ACGACGTTGT AAAACGACGG CCAGTGAATT
3001 GTAATACGAC TCACTATA

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D. pGEM®-T Easy Vector Restriction Sites

The pGEM®-T Easy Vector has been linearized at base 60 with *EcoR V* and a T added to both 3'-ends. This site will not be recovered upon ligation of the vector and insert. The tables were constructed using DNASTAR® sequence analysis software. Please note that we have not verified this information by restriction digestion with each enzyme listed. The location given specifies the 3'-end of the cut DNA (the base to the left of the cut site). Please contact your local Promega Branch Office or Distributor if you identify a discrepancy. In the U.S., contact Technical Services at 1-800-356-9526.

Table 7. Restriction Enzymes That Cut the pGEM®-T Easy Vector Between 1 and 5 Times

Enzyme	# of Sites	Location	Enzyme	# of Sites	Location
Aat II	1	20	Fok I	5	134, 1376, 1557, 1844, 2934
Acc I	1	91	<i>Fsp I</i>	2	1632, 2858
Acy I	2	17, 1947	Hae II	4	395, 765, 2758, 2766
<i>Afl III</i>	2	114, 517	<i>Hga I</i>	4	628, 1206, 1936, 2824
Alw26 I	2	1471, 2247	Hinc II	1	92
Alw44 I	2	831, 2077	<i>Hind II</i>	1	92
<i>AlwNI</i>	1	933	Hsp92 I	2	17, 1947
Apa I	1	14	<i>Ksp I</i>	1	49
<i>AspH I</i>	4	109, 835, 1996, 2081	<i>Mae I</i>	5	65, 1012, 1265, 1600, 2758
Ava II	2	1548, 1770	Mlu I	1	114
Ban I	3	261, 1358, 2644	Nae I	1	2710
Ban II	3	14, 109, 2682	Nci I	4	30, 897, 1593, 1944
Bbu I	1	26	Nco I	1	37
Bgl I	4	39, 42, 1530, 2851	Nde I	1	97
<i>Bsa I</i>	1	1471	NgoM I	1	2708
<i>BsaA I</i>	1	2607	Not I	2	43, 77
<i>BsaH I</i>	2	17, 1947	Nsi I	1	127
<i>BsaJ I</i>	5	37, 46, 256, 677, 2954	<i>Nsp I</i>	2	26, 521
<i>Bsp120 I</i>	1	10	<i>Ppu10 I</i>	1	123
<i>BspH I</i>	2	1237, 2245	Pst I	1	88
<i>BspM I</i>	1	77	Pvu I	2	1780, 2879
<i>BssS I</i>	2	690, 2074	Pvu II	2	341, 2908
BstO I	5	257, 545, 666, 679, 2955	Rsa I	1	1890
BstX I	1	118	Sac I	1	109
BstZ I	3	31, 43, 77	Sac II	1	49
<i>Cfr10 I</i>	2	1490, 2708	Sal I	1	90
Dde I	4	792, 1201, 1367, 1907	Sca I	1	1890
Dra I	3	1276, 1295, 1987	Sin I	2	1548, 1770
<i>Dra III</i>	1	2607	Spe I	1	64
<i>Drd I</i>	2	625, 2562	Sph I	1	26
<i>Dsa I</i>	2	37, 46	<i>Sse8387 I</i>	1	88
<i>Eag I</i>	3	31, 43, 77	Ssp I	3	2214, 2399, 2423
<i>Ear I</i>	3	401, 2205, 2896	Sty I	1	37
EcIHK I	1	1410	Taq I	5	56, 91, 617, 2061, 2640
Eco52 I	3	31, 43, 77	<i>Tfi I</i>	2	352, 492
EcoICR I	1	107	Vsp I	3	288, 347, 1582
EcoR I	2	52, 70	Xmn I	1	2009
EcoR V	1	60 (see above)			

Table 8. Restriction Enzymes That Do Not Cut the pGEM[®]-T Easy Vector.

<i>Acc III</i>	<i>Bbs I</i>	<i>Bst1107 I</i>	<i>Ehe I</i>	<i>PfM I</i>	<i>Sma I</i>
<i>Acc65 I</i>	<i>Bcl I</i>	<i>Bst98 I</i>	<i>Fse I</i>	<i>PinA I</i>	<i>SnaB I</i>
<i>AccB7 I</i>	<i>Bgl II</i>	<i>BstE II</i>	<i>Hind III</i>	<i>Pme I</i>	<i>Spl I</i>
<i>Afl II</i>	<i>Blp I</i>	<i>Bsu36 I</i>	<i>Hpa I</i>	<i>Pml I</i>	<i>Srf I</i>
<i>Age I</i>	<i>Bpu1102 I</i>	<i>Cla I</i>	<i>I-Ppo I</i>	<i>PpuM I</i>	<i>Stu I</i>
<i>Asc I</i>	<i>BsaB I</i>	<i>Csp I</i>	<i>Kas I</i>	<i>PshA I</i>	<i>Swa I</i>
<i>Ava I</i>	<i>BsaM I</i>	<i>Csp45 I</i>	<i>Kpn I</i>	<i>Psp5 II</i>	<i>Tth111 I</i>
<i>Avr II</i>	<i>BsiW I</i>	<i>Dra II</i>	<i>Nar I</i>	<i>PspA I</i>	<i>Xba I</i>
<i>Bal I</i>	<i>Bsm I</i>	<i>Eco47 III</i>	<i>Nhe I</i>	<i>Rsr II</i>	<i>Xcm I</i>
<i>BamH I</i>	<i>BsrBr I</i>	<i>EcoN I</i>	<i>Nru I</i>	<i>Sfi I</i>	<i>Xho I</i>
<i>Bbe I</i>	<i>BsrG I</i>	<i>Eco72 I</i>	<i>Pac I</i>	<i>Sgf I</i> ⁽ⁱ⁾	<i>Xma I</i>
<i>BbrP I</i>	<i>BssH II</i>	<i>Eco81 I</i>	<i>PaeR7 I</i>	<i>SgrA I</i>	

Note: The enzymes listed in boldface type are available from Promega.

Table 9. Restriction Enzymes That Cut the pGEM[®]-T Easy Vector 6 or More Times.

<i>Aci I</i>	<i>Bst71 I</i>	<i>Hae III</i>	<i>Mae II</i>	<i>MspA1 I</i>	<i>SfaN I</i>
<i>Alu I</i>	<i>BstU I</i>	<i>Hha I</i>	<i>Mae III</i>	<i>Nla III</i>	<i>Tru9 I</i>
<i>Bbv I</i>	<i>Cfo I</i>	<i>Hinf I</i>	<i>Mbo I</i>	<i>Nla IV</i>	<i>Xho II</i>
<i>BsaO I</i>	<i>Dpn I</i>	<i>HinP I</i>	<i>Mbo II</i>	<i>Ple I</i>	
<i>Bsp1286 I</i>	<i>Dpn II</i>	<i>Hpa II</i>	<i>Mnl I</i>	<i>Sau3A I</i>	
<i>Bsr I</i>	<i>Eae I</i>	<i>Hph I</i>	<i>Mse I</i>	<i>Sau96 I</i>	
<i>BsrS I</i>	<i>Fnu4H I</i>	<i>Hsp92 II</i>	<i>Msp I</i>	<i>ScrF I</i>	

Note: The enzymes listed in boldface type are available from Promega.

XI. Appendix B: Reference Information

A. Composition of Buffers and Solutions

IPTG stock solution (0.1M)

1.2g IPTG (Cat.# V3951)

Add water to 50ml final volume.
Filter-sterilize and store at 4°C.

X-Gal (2ml)

100mg 5-bromo-4-chloro-3-indolyl- β -D-galactoside
(Cat.#V3941)

Dissolve in 2ml N,N'-dimethyl-formamide. Cover with aluminum foil and store at -20°C.

LB medium (per liter)

10g Bacto®-Tryptone
5g Bacto®-Yeast Extract
5g NaCl

Adjust pH to 7.0 with NaOH.

LB plates with ampicillin

Add 15g agar to 1 liter of LB medium. Autoclave. Allow the medium to cool to 50°C before adding ampicillin to a final concentration of 100 μ g/ml. Pour 30-35ml of medium into 85mm petri dishes. Let the agar harden. Store at 4°C for up to one month or at room temperature for up to one week.

LB plates with ampicillin/IPTG/X-Gal

Make the LB plates with ampicillin as above; then supplement with 0.5mM IPTG and 80 μ g/ml X-Gal and pour the plates. Alternatively, 100 μ l of 100mM IPTG and 20 μ l of 50mg/ml X-Gal may be spread over the surface of an LB-ampicillin plate and allowed to absorb for 30 minutes at 37°C prior to use.

SOC medium (100ml)

2.0g Bacto®-tryptone
0.5g Bacto®-yeast extract
1ml 1M NaCl
0.25ml 1M KCl
1ml 2M Mg²⁺ stock, filter-sterilized (as prepared below)
1ml 2M glucose, filter-sterilized

Add Bacto®-tryptone, Bacto®-yeast extract, NaCl and KCl to 97ml distilled water. Stir to dissolve. Autoclave and cool to room temperature. Add 2M Mg²⁺ stock and 2M glucose, each to a final concentration of 20mM. Bring to 100ml with sterile, distilled water. Filter the complete medium through a 0.2 μ m filter unit. The final pH should be 7.0.

2M Mg²⁺ stock

20.33g MgCl₂ • 6H₂O
24.65g MgSO₄ • 7H₂O

Add distilled water to 100ml. Filter sterilize.

2X Rapid Ligation Buffer, T4 DNA Ligase (provided)

60mM Tris-HCl (pH 7.8)
20mM MgCl₂
20mM DTT
2mM ATP
10% polyethylene glycol (MW8000, ACS Grade)

Store in single-use aliquots at -20°C. Avoid multiple freeze/thaw cycles.

TYP broth (per liter)

16g Bacto®-Tryptone
16g Bacto®-Yeast Extract
5g NaCl
2.5g K₂HPO₄

B. Related Products

Thermostable Enzymes

Product	Size	Cat.#
PCR Core System I	200 reactions	M7660
PCR Core System II	200 reactions	M7665
<i>Taq</i> DNA Polymerase	100u	M1661
	500u	M1665
<i>Pfu</i> DNA Polymerase	100u	M7741
	500u	M7745
<i>Tli</i> DNA Polymerase	50u	M7101

Wizard® Plus SV Minipreps DNA Purification Systems(i)

Product	Cat.#
Wizard® Plus SV Minipreps DNA Purification System*	A1330
Wizard® Plus SV Minipreps DNA Purification System*	A1340
Each system contains sufficient reagents for 50 isolations from 1-10ml of culture. Cat.# A1340 includes 20 Miniprep Vacuum Adapters.	

Product	Cat.#
Wizard® Plus SV Minipreps DNA Purification System*	A1460
Wizard® Plus SV Minipreps DNA Purification System*	A1470
Each system contains sufficient reagents for 250 isolations from 1-10ml of culture. Cat.# A1470 includes 20 Miniprep Vacuum Adapters.	

Wizard® DNA Purification Systems

Product	Size	Cat.#
Wizard® Plus Minipreps DNA Purification System	50 preps	A7100
Wizard® Plus Minipreps DNA Purification System	100 preps	A7500
Wizard® Plus Minipreps DNA Purification System	250 preps	A7510
Wizard® Plus Midipreps DNA Purification System	25 preps	A7640
Wizard® Plus Maxipreps DNA Purification System	10 preps	A7270
Wizard® Plus Megapreps DNA Purification System	5 preps	A7300
Wizard® Plus Series 9600™ DNA Purification System	2 x 96 preps	A7000
Each system contains sufficient reagents for isolating DNA as described in Table 3.		

Product	Cat.#
Wizard® PCR Preps DNA Purification System	A7170
Each system contains sufficient reagents to purify 50 DNA samples, either with or without prior gel extraction.	

Product	Capacity	Cat.#
Vac-Man® Laboratory Vacuum Manifold	20 samples	A7231
Vac-Man® Jr. Laboratory Vacuum Manifold	2 samples	A7660

Competent Cells

Product	Efficiency	Size	Cat.#
JM109 Competent Cells, High Efficiency	>10 ⁸ cfu/μg	1ml (5 x 200μl)	L2001

Each order of competent cells is provided with control DNA.

*Components of the Wizard® Plus SV Minipreps DNA Purification System should not be exchanged for or replaced with components from any Wizard® Plus DNA Purification System. Components for the Wizard® Plus Systems and the Wizard® Plus SV Systems are not interchangeable.

Reagents and dNTPs

Product	Size	Cat.#
X-Gal	100mg	V3941
IPTG, Dioxane-Free	1g	V3955
	5g	V3951
	50g	V3953
dATP, 100mM	40µmoles	U1201
dCTP, 100mM	40µmoles	U1221
dGTP, 100mM	40µmoles	U1211
dTTP, 100mM	40µmoles	U1231
dATP, dCTP, dGTP, dTTP, each at 100mM	40µmoles of each	U1240
dATP, dCTP, dGTP, dTTP, each at 100mM	10µmoles of each	U1330
PCR Nucleotide Mix, 40mM	200µl	C1141
	1,000µl	C1145
Lambda DNA/ <i>Hind</i> III Markers	100µg	G1711
100bp DNA Ladder	250µl	G2101

(a) Licensed under one or both of U.S. Pat. No. 5,487,993 and European Pat. No. 0 550 693.

(b) U.S. Pat. No. 4,766,072 has been issued to Promega Corporation for transcription vectors having two different bacteriophage RNA polymerase promoter sequences separated by a series of unique restriction sites into which foreign DNA can be inserted.

(c) The PCR process is covered by patents issued and applicable in certain countries. Promega does not encourage or support the unauthorized or unlicensed use of the PCR process.

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(f) The method of recombinant expression of *Coleoptera* luciferase is covered by U.S. Pat. No. 5,583,024 assigned to The Regents of the University of California.

(g) Patent Pending.

(h) U.S. Pat. Nos. 5,658,548 and other patents.

(i) U.S. Pat. No. 5,391,487 has been issued to Promega Corporation for Restriction Endonuclease Sgf I.

(j) Patent Pending.

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**Ligations
Using the 2X
Rapid
Ligation
Buffer**
(Section IV.A)

1. Briefly centrifuge the pGEM[®]-T or pGEM[®]-T Easy Vector and Control Insert DNA tubes to collect contents at the bottom of the tube.
2. Set up ligation reactions as described below. **Note:** Use 0.5ml tubes known to have low DNA-binding capacity (e.g., VWR Cat.# 20170-310).
3. Vortex the 2X Rapid Ligation Buffer vigorously before each use.

	Standard Reaction	Positive Control	Background Control
2X Rapid Ligation Buffer, T4 DNA Ligase	5µl	5µl	5µl
pGEM [®] -T or pGEM [®] -T Easy Vector (50ng)	1µl	1µl	1µl
PCR product	Xµl	–	–
Control Insert DNA	–	2µl	–
T4 DNA Ligase (3 Weiss units/µl)	<u>1µl</u>	<u>1µl</u>	<u>1µl</u>
deionized water to a final volume of	10µl	10µl	10µl

4. Mix the reactions by pipetting. Incubate the reactions 1 hour at room temperature.

Alternatively, if the maximum number of transformants is required, incubate the reactions overnight at 4°C.



This protocol uses the **new 2X Rapid Ligation Buffer**. Use the appropriate volume when setting up the ligation reaction.