ZAP Express Predigested Vector Kit and
ZAP Express Predigested Gigapack Cloning Kits
EcoR I/CIAP-Treated

Instruction Manual
Catalog #239211 (ZAP Express Predigested Vector Kit) and
Catalog #239614 (ZAP Express Predigested Gigapack III Gold Cloning Kit)
Revision C.0

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239211-12
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ZAP Express Predigested Vector Kit
and ZAP Express Predigested Gigapack Cloning Kits
EcoRI/CIAP-Treated

CONTENTS

Limited Product Warranty ................................................................................................................ 2
Materials Provided .......................................................................................................................... 1
Storage Conditions .............................................................................................................................. 2
Notices to Purchaser ........................................................................................................................... 2
Introduction ......................................................................................................................................... 3

Overview of the ZAP Express Vector System ...................................................................... 3
The pBK-CMV Vector ......................................................................................................................... 4

Bacterial Host Strains ......................................................................................................................... 5

Host Strain Genotypes ......................................................................................................................... 5
XL1-Blue MRF’ Bacterial Strain Description ............................................................................... 5
Recommended Media ............................................................................................................................. 6
Establishing an Agar Plate Bacterial Stock ............................................................................... 6
Preparing a –80°C Bacterial Glycerol Stock ............................................................................... 7
Color Selection by IPTG and X-gal .................................................................................................. 7
Growth of Cells for Plating Phage ..................................................................................................... 7

Helper Phage ........................................................................................................................................... 8

Storing the Helper Phage .................................................................................................................... 8
Titering the Helper Phage ...................................................................................................................... 8
Amplifying the Helper Phage ............................................................................................................... 9

Ligating the Insert ................................................................................................................................. 10

Packaging and Titering .......................................................................................................................... 11

Packaging Extracts ............................................................................................................................... 11
Packaging Protocol ............................................................................................................................... 11
Titering the Packaging Reaction ......................................................................................................... 12
Testing the Efficiency of the Gigapack III Gold Packaging Extract with the Wild-Type Lambda Control DNA (Optional) .................................................................................................................. 14

Plating for Blue–White Color Selection ......................................................................................... 15

Amplifying the ZAP Express Library ............................................................................................... 16
Performing Plaque Lifts ..................................................................................................................... 17
Hybridizing and Screening .................................................................................................................. 19
Antibody Screening Protocol in *Escherichia coli* ................................................................. 19

Eukaryotic Screening with the ZAP Express Library ............................................................... 19
  Selective Assay .................................................................................................................. 19
  Panning Assay .................................................................................................................. 19
  Functional Assay .............................................................................................................. 20
  Protocol .............................................................................................................................. 21

Eukaryotic Expression ........................................................................................................... 21

In Vivo Excision of the pBK-CMV Phagemid Vector from the ZAP Express Vector .......... 22

In Vivo Excision Protocols Using ExAssist Helper Phage with XLOLR Strain ................. 23
  Single-Clone Excision Protocol ..................................................................................... 23
  Mass Excision Protocol ................................................................................................. 25

Appendix: Recovery of Single-Stranded DNA from Cells Containing the pBK-CMV Phagemid Vector ................................................................................................................. 27
  Single-Stranded Rescue Protocol .................................................................................... 28

Troubleshooting .................................................................................................................. 29
  Packaging ......................................................................................................................... 29
  Excision ............................................................................................................................ 29

Preparation of Media and Reagents .................................................................................. 30

References ......................................................................................................................... 32

Endnotes ............................................................................................................................. 32

MSDS Information ............................................................................................................. 32
ZAP Express Predigested Vector Kit
and ZAP Express Predigested Gigapack Cloning Kits
EcoR I/CIAP-Treated

MATERIALS PROVIDED

<table>
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<th>Catalog #239211</th>
<th>Catalog #239614</th>
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<td>ZAP Express vector digested with EcoR I, CIAP treated&lt;sup&gt;a&lt;/sup&gt;</td>
<td>10 μg</td>
<td>10 μg</td>
</tr>
<tr>
<td>pRheo test insert</td>
<td>1.25 μg</td>
<td>1.25 μg</td>
</tr>
<tr>
<td>XL1-Blue MRF&lt;sup&gt;+&lt;/sup&gt; strain&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.5-ml bacterial glycerol stock</td>
<td>0.5-ml bacterial glycerol stock</td>
</tr>
<tr>
<td>XLOLR strain&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.5-ml bacterial glycerol stock</td>
<td>0.5-ml bacterial glycerol stock</td>
</tr>
<tr>
<td>ExAssist interference-resistant helper phage&lt;sup&gt;c,d&lt;/sup&gt;</td>
<td>1 ml</td>
<td>1 ml</td>
</tr>
<tr>
<td>R408 Interference-Resistant Helper Phage&lt;sup&gt;c,e&lt;/sup&gt;</td>
<td>1 ml</td>
<td>1 ml</td>
</tr>
<tr>
<td>Gigapack III Gold packaging extract&lt;sup&gt;f&lt;/sup&gt;</td>
<td>—</td>
<td>11 × 25 μl</td>
</tr>
<tr>
<td>λcl857 Sam7 wild-type lambda control DNA&lt;sup&gt;g&lt;/sup&gt;</td>
<td>—</td>
<td>1.05 μg</td>
</tr>
<tr>
<td>VCS257 host strain&lt;sup&gt;h&lt;/sup&gt;</td>
<td>—</td>
<td>1-ml bacterial glycerol stock</td>
</tr>
</tbody>
</table>

<sup>a</sup> On arrival, store the ZAP Express vector at –20°C. After thawing, aliquot and store at –20°C. Do not pass through more than two freeze–thaw cycles. For short-term storage, store at 4°C for 1 month.

<sup>b</sup> Use the XLOLR strain for plating excised phagemids and the XL1-Blue MRF<sup>+</sup> strain for all other manipulations. For host strain shipping and storage conditions, see Bacterial Host Strains.

<sup>c</sup> The titer of the ExAssist interference-resistant helper phage is ~1.0 × 10<sup>10</sup> pfu/ml. This supercoiled single-stranded DNA migrates at ~5 kb on an agarose gel. The ExAssist helper phage is recommended for excision of the pBK-CMV phagemid vector from the ZAP Express vector. It should not be used for single-stranded rescue in general, because this f1 helper phage possesses α-complementing β-galactosidase sequences which may interfere with sequencing or site-directed mutagenesis where oligonucleotide primers hybridize to β-galactosidase sequences (e.g., M13–20 primer).

<sup>d</sup> Retiter after 1 month. (Take care not to contaminate the ZAP Express vector with this high-titer filamentous helper phage.) Store at –80°C.

<sup>e</sup> The titer of the R408 Interference-Resistant Helper Phage is ~7.5 × 10<sup>10</sup> pfu/ml. This supercoiled single-stranded DNA migrates at ~4 kb on an agarose gel. The R408 Interference-Resistant Helper Phage is recommended for single-stranded rescue (see Appendix: Recovery of Single-Stranded DNA from Cells Containing the pBK-CMV Phagemid Vector).

<sup>f</sup> Gigapack III packaging extract is very sensitive to slight variations in temperature. Storing the packaging extracts at the bottom of a –80°C freezer directly from the dry ice shipping container is required in order to prevent a loss of packaging efficiency. Transferring tubes from one freezer to another may also result in a loss of efficiency. **Do not allow the packaging extracts to thaw!** Do not store the packaging extracts in liquid nitrogen as the tubes may explode.

<sup>g</sup> The λcl857 Sam7 wild-type lambda control DNA is shipped frozen and should be stored at –80°C immediately on receipt.

<sup>h</sup> The VCS257 host strain, included for plating the λcl857 Sam7 wild-type lambda control DNA, is shipped as a frozen bacterial glycerol stock (see Bacterial Host Strains for additional storage instructions) and should also be stored at –80°C immediately on receipt. This control host strain is a derivative of DP50 supF and should be used only when plating the packaged lambda control DNA. The lambda control DNA used with Gigapack III Gold packaging extract requires a supF mutation in the bacterial host to plate efficiently.

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**Storage Conditions**

The ZAP Express Vector and Test Insert: –20°C  
Helper Phage: –80°C  
Bacterial Glycerol Stocks: –80°C  

**Notices to Purchaser**

The use of the CMV Promoter is covered under U.S. Patent Nos. 5,168,062 and 5,385,839 owned by the University of Iowa Research Foundation and licensed FOR RESEARCH USE ONLY. For further information, please contact UIRF at 319-335-4546.

This product is for research purposes only and must be used in accordance with NIH guidelines for recombinant DNA.
INTRODUCTION

Overview of the ZAP Express Vector System

The ZAP Express vector (see Figure 1) allows both eukaryotic and prokaryotic expression, while also increasing both cloning capacity and the number of unique lambda cloning sites. The ZAP Express vector has 12 unique cloning sites which will accommodate DNA inserts from 0 to 12 kb in length. The 12 unique cloning sites are Apa I, BamH I, EcoR I, Hind III, Kpn I, Not I, Sac I, Sal I, Sma I, Spe I, Xba I, and Xho I. Inserts cloned into the ZAP Express vector can be excised out of the phage in the form of the kanamycin-resistant pBK-CMV phagemid vector (see Figure 2) by the same excision mechanism found in the Lambda ZAP vectors.

![Figure 1: Map of the ZAP Express vector.](image)

Clones in the ZAP Express vector can be screened with either DNA probes or antibody probes, and in vivo rapid excision of the pBK-CMV phagemid vector allows insert characterization in a plasmid system. The polylinker of pBK-CMV phagemid has 17 unique cloning sites flanked by T3 and T7 promoters and has 5 primer sites for DNA sequencing. The plasmid has the bacteriophage f1 origin of replication allowing rescue of single-stranded DNA, which can be used for DNA sequencing or site-directed mutagenesis. Unidirectional deletions can be made using exonuclease III and mung bean nuclease by taking advantage of the unique positioning of 5' and 3' restriction sites. Transcripts made from the T3 and T7 promoters generate riboprobes useful in Southern and Northern blotting, and the lacZ promoter may be used to drive expression of fusion proteins suitable for Western blot analysis or protein purification.

Eukaryotic expression of inserts is driven by the cytomegalovirus (CMV) immediate early (IE) promoter with the SV40 transcription terminator and polyadenylation signal.

Stable selection of clones in eukaryotic cells is made possible by the presence of the neomycin-and kanamycin-resistance gene, which is driven by the SV40 early promoter with TK transcription polyadenylation signals to render transfecants resistant to G418 (geneticin).
The **pBK-CMV Vector**

![Diagram of the pBK-CMV Vector](image)

**pBK-CMV Multiple Cloning Site Region**
*(sequence shown 952–1196)*

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<thead>
<tr>
<th>Restriction Site</th>
<th>Fragment</th>
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<tr>
<td>Kpn I</td>
<td>T7 Promoter</td>
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<td>BstX I</td>
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<td></td>
</tr>
<tr>
<td>Sma I</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not I</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apa I</td>
<td></td>
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<td>Xba I</td>
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<td>Sca I</td>
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<tr>
<td>Xho I</td>
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<tr>
<td>Hind III</td>
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<td>EcoR I</td>
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<tr>
<td>BamH I</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spe I</td>
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</tr>
<tr>
<td>Acc I</td>
<td></td>
<td></td>
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<td>Sma I</td>
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**Feature**

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<td>SV40 polyA signal</td>
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<td>$\beta$-galactosidase $\alpha$-fragment coding sequence (lacZ')</td>
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<tr>
<td>multiple cloning site</td>
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<tr>
<td>lac promoter</td>
<td>1184–1305</td>
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<td>CMV promoter</td>
<td>1306–1895</td>
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<tr>
<td>pUC origin of replication</td>
<td>1954–2621</td>
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<tr>
<td>HSV-thymidine kinase (TK) polyA signal</td>
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<tr>
<td>neomycin/kanamycin resistance ORF</td>
<td>3209–4000</td>
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<tr>
<td>SV40 promoter</td>
<td>4035–4373</td>
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<tr>
<td>bla promoter</td>
<td>4392–4518</td>
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**FIGURE 2** The pBK-CMV phagemid vector. The complete sequence and list of restriction sites are available at [www.genomics.agilent.com](http://www.genomics.agilent.com).
# Bacterial Host Strains

## Host Strain Genotypes

<table>
<thead>
<tr>
<th>Host strain</th>
<th>Genotype</th>
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<tbody>
<tr>
<td>XL1-Blue MRF’ strain</td>
<td>(\Delta(mcrA)183\ \Delta(mcrCB-hsdSMR-mrr)173\ endA1\ supE44\ thi-1\ recA1\ gyrA96\ relA1\ lac\ [F’ proAB lac(\Delta M15) Tn10 (Tetr)])</td>
</tr>
<tr>
<td>XLOLR strain(^a)</td>
<td>(\Delta(mcrA)183\ \Delta(mcrCB-hsdSMR-mrr)173\ endA1\ thi-1\ recA1\ gyrA96\ relA1\ lac\ [F’ proAB lac(\Delta M15) Tn10 (Tetr)]) Su(^-) (nonsuppressing)(\lambda)’ (lambda resistant)</td>
</tr>
</tbody>
</table>

\(^a\) Use the XLOLR strain for excision only.

## XL1-Blue MRF’ Bacterial Strain Description

The RecA\(^+\) *E. coli* host strain XL1-Blue MRF’ is supplied with the ZAP Express cDNA synthesis kit.\(^3\) Because the pBK-CMV phagemid vector does not require a \(supF\) genotype, the amplified library grows very efficiently on the XL1-Blue MRF’ strain. In addition, use of the correct host strain is important when working with the pBK-CMV phagemid vector as the F’ episome present in the XL1-Blue MRF’ strain serves three purposes.

First, the \(\Delta M15\) lac\(Z\) gene present on the F’ episome is required for the \(\beta\)-galactosidase-based nonrecombinant selection strategy. When cDNA is present in the polylinker, expression from the lac\(Z\) gene is disrupted and white plaques are produced. In contrast, without insert in the polylinker, the amino terminus of \(\beta\)-galactosidase is expressed and nonrecombinants can be scored visually by the presence of blue plaques. To produce an enzymatically active \(\beta\)-galactosidase protein, two domains are required: the \(\alpha\)-region expressed by the vector and the \(\Delta M15\) lac\(Z\) domain expressed by the F’ episome. These two domains fold to form a functional protein, the \(\alpha\)-region complementing the missing amino acids resulting from the \(\Delta M15\) mutation. Therefore, in order to utilize the nonrecombinant selection strategy, the correct host strain must be used to produce a functional \(\beta\)-galactosidase protein.

Second, the F’ episome expresses the genes forming the F’ pili found on the surface of the bacteria. Without pili formation, filamentous phage (i.e., M13 or f1) infection could not occur. Because the conversion of a recombinant ZAP Express clone to a pBK-CMV phagemid vector requires superinfection with a filamentous helper phage, the F’ episome is required for in vivo excision (see In Vivo Excision of the pBK-CMV Phagemid Vector from the ZAP Express Vector).

Third, the F’ episome contains the lac repressor (lac\(I\)\(^q\) gene), which blocks transcription from the lac\(Z\) promoter in the absence of the inducer IPTG. This repressor is important for controlling expression of fusion proteins which may be toxic to the *E. coli*. Because the presence of the lac\(I\)\(^q\) repressor in the *E. coli* host strain can potentially increase the representation or completeness of the library, XL1-Blue MRF’ is useful for screening the amplified library.
**Note**  The strains used for the Lambda gt11 vector (i.e., Y1088, Y1089, and Y1090) are not suitable for use with the ZAP Express vector because these strains contain the plasmid pMC9, a pBR322 derivative, which contains many of the same sequences as those found in the phagemid portion of the ZAP Express vector. The SURE strain and the SOLR strain are not compatible with the ZAP Express system, since these strains contain the kanamycin-resistance gene found in the pBK-CMV phagemid vector. Using these strains with the ZAP Express vector could result in recombination between the homologous sequences.

### Recommended Media

<table>
<thead>
<tr>
<th>Host strain</th>
<th>Agar plates and liquid medium for bacterial streak and glycerol stock</th>
<th>Liquid medium for bacterial cultures prior to phage attachment</th>
<th>Agar plates and top agar for plaque formation</th>
<th>Agar plates for excision protocol</th>
</tr>
</thead>
<tbody>
<tr>
<td>XLOLR strain</td>
<td>LB-tetacycline&lt;sup&gt;a&lt;/sup&gt;</td>
<td>LB broth with supplements&lt;sup&gt;c&lt;/sup&gt;</td>
<td>—</td>
<td>LB-kanamycin&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>VCS257 strain&lt;sup&gt;d&lt;/sup&gt;</td>
<td>LB&lt;sup&gt;b&lt;/sup&gt;</td>
<td>LB broth with supplements&lt;sup&gt;c&lt;/sup&gt;</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>XL1-Blue MRF&lt;sup&gt;+&lt;/sup&gt; strain</td>
<td>LB-tetacycline&lt;sup&gt;a&lt;/sup&gt;</td>
<td>LB broth with supplements&lt;sup&gt;c&lt;/sup&gt;</td>
<td>NZY&lt;sup&gt;c&lt;/sup&gt;</td>
<td>—</td>
</tr>
</tbody>
</table>

<sup>a</sup> See Preparation of Media and Reagents.

<sup>b</sup> LB broth with 0.2% (w/v) maltose and 10 mM MgSO<sub>4</sub>.

<sup>c</sup> Maltose and magnesium supplements are required for optimal lambda phage receptor expression on the surface of the XL1-Blue MRF<sup>+</sup> host cell. The media supplements are not required for helper phage infection, but are included in both protocols for simplified media preparation.

<sup>d</sup> For use with Gigapack III Gold packaging extract and wild-type control only. Supplied with Gigapack III Gold packaging extract.

### Establishing an Agar Plate Bacterial Stock

The bacterial host strains are shipped as bacterial glycerol stocks. On arrival, prepare the following plates from the bacterial glycerol stocks.

**Note**  The host strains may thaw during shipment. The vials should be stored immediately at −20° or −80°C, but most strains remain viable longer if stored at −80°C. It is best to avoid repeated thawing of the host strains in order to maintain extended viability.

1. Revive the stored cells by scraping off splinters of solid ice with a sterile wire loop.

2. Streak the splinters onto an LB agar plate containing the appropriate antibiotic (see Recommended Media), if one is necessary.

3. Incubate the plate overnight at 37°C.

4. Seal the plate with Parafilm® laboratory film and store the plate at 4°C for up to 1 week.

5. Restreak the cells onto a fresh plate every week.
Preparing a –80°C Bacterial Glycerol Stock

1. In a sterile 50-ml conical tube, inoculate 10 ml of LB broth with the appropriate antibiotic (see Recommended Media) with one colony from the plate. Grow the cells to late log phase.

2. Add 4.5 ml of a sterile glycerol-liquid medium solution (prepared by mixing 5 ml of glycerol + 5 ml of the appropriate medium) to the bacterial culture from step 1. Mix well.

3. Aliquot into sterile centrifuge tubes (1 ml/tube).

This preparation may be stored at –20°C for 1–2 years or at –80°C for more than 2 years.

Color Selection by IPTG and X-gal

The color selection by α-complementation with the ZAP Express vector requires a high amount of IPTG and X-gal for generation of the blue color (see Plating for Blue-White Color Selection). Transcription and translation of the fusion protein are normal, but the large polylinker present within the pBK-CMV phagemid vector, which is present in the ZAP Express vector, is partly responsible for the reduced activity of the β-galactosidase protein—not the promoter. As would be expected, the copy number of the ZAP Express vector is much less per cell than the copy number of pBK-CMV phagemid vector derivatives. However, it is important to note that the color assay is used only for determining the ratio of recombinants to nonrecombinants within a newly constructed library and is not used for any other manipulations.

Growth of Cells for Plating Phage

Bacterial cultures for plating phage should be started from a fresh plate using a single colony and should be grown overnight with vigorous shaking at 30°C in 50 ml of LB broth supplemented with 0.2% (w/v) maltose and 10 mM MgSO₄. (Do not use tetracycline in the presence of magnesium.) The lower temperature ensures that the cells will not overgrow. The cells should be spun at 1000 × g for 10 minutes then gently resuspended in 10 ml of 10 mM MgSO₄. Before use, dilute cells to an OD₆₀₀ of 0.5 with 10 mM MgSO₄. Bacterial cells prepared in this manner can be used for all phage manipulations described within the manual. Highest efficiencies are obtained from freshly prepared cells.
**HELPER PHAGE**

Two different helper phages are provided with the ZAP Express cDNA synthesis kit: (1) the ExAssist interference-resistant helper phage with XLOLR strain and (2) the R408 helper phage. The ExAssist interference-resistant helper phage with XLOLR strain is designed to allow efficient in vivo excision of the pBK-CMV phagemid vector from the ZAP Express vector while preventing problems associated with helper phage co-infection. The ExAssist helper phage contains an amber mutation that prevents replication of the phage genome in a nonsuppressing *E. coli* strain (e.g., XLOLR cells). Only the excised phagemid can replicate in the host, removing the possibility of co-infection from the ExAssist helper phage. Because ExAssist helper phage cannot replicate in the XLOLR strain, single-stranded rescue cannot be performed in this strain using ExAssist helper phage. XLOLR cells are also resistant to lambda infection, preventing lambda DNA contamination after excision.

**Storing the Helper Phage**

The ExAssist helper phage and the R408 helper phage are supplied in 7% dimethylsulfoxide (DMSO) and should be stored at −80°C. The helper phage may be stored for short periods of time at −20°C or 4°C. It is important to titer the helper phage prior to each use. Expect titers of approximately $10^{10}$ pfu/ml for the ExAssist helper phage or $10^{10}$ pfu/ml for the R408 helper phage. If the titer drops over time, prepare a fresh high-titer stock of the helper phage as outlined in Amplifying the Helper Phage.

**Titering the Helper Phage**

1. Transfer a colony of XL1-Blue MRF’ cells into 10 ml of LB broth with supplements in a 50-ml conical tube. Incubate the conical tube with shaking at 37°C until growth reaches an OD$_{600}$ of 1.0.

2. Dilute the phage (10$^{-4}$–10$^{-7}$) in SM buffer (See Preparation of Media and Reagents) and combine 1 μl of each dilution with 200 μl of XL1-Blue MRF’ cells (OD$_{600}$ = 1.0).

3. Incubate the helper phage and the XL1-Blue MRF’ cells for 15 minutes at 37°C to allow the phage to attach to the cells.

4. Add 3 ml of NZY top agar, melted and cooled to ~48°C, and plate immediately onto dry, prewarmed NZY agar plates. Allow the plates to set for 10 minutes.
5. Invert the plates and incubate overnight at 37°C.

**Note**  
*ExAssist and R408 plaques will have a cloudier appearance than lambda phage plaques.*

6. To determine the titer [in plaque-forming units per milliliter (pfu/ml)], use the following formula:

\[
\text{Number of plaques (pfu) } \times \text{ dilution factor} \times \frac{\text{Volume plated (\(\mu l\))}}{1000 \, \mu l / \text{ml}}
\]

where the volume plated (in microliters) refers to the volume of the helper phage solution added to the cells.

### Amplifying the Helper Phage

1. Transfer a colony of XL1-Blue MRF´ cells into 10 ml of LB broth with supplements in a 50-ml conical tube. Incubate the conical tube with shaking at 37°C until growth reaches an OD$_{600}$ of 0.3.

**Note**  
*An OD$_{600}$ of 0.3 corresponds to $2.5 \times 10^8$ cells/ml.*

2. Add the helper phage at a multiplicity of infection (MOI) of 20:1 (phage-to-cells ratio).

3. Incubate the conical tube at 37°C for 15 minutes to allow the phage to attach to the cells.

4. Incubate the conical tube with shaking at 37°C for 8 hours.

5. Heat the conical tube at 65°C for 15 minutes.

6. Spin down the cell debris and transfer the supernatant to a fresh conical tube.

7. The titer of the supernatant should be between $7.5 \times 10^{10}$ and $1.0 \times 10^{12}$ pfu/ml for ExAssist helper phage or between $1.0 \times 10^{11}$ and $1.0 \times 10^{12}$ pfu/ml for R408 helper phage.

**Note**  
*ExAssist and R408 plaques will have a cloudier appearance than lambda phage plaques.*

8. Add dimethylsulfoxide (DMSO) to a final concentration of 7% (v/v) and store at −80°C.

9. For further details about helper phage titering or amplification, please see *Titering the Helper Phage* or Reference 4.
LIGATING THE INSERT

Note In all ligations, the final glycerol content should be less than 5% (v/v). Do not exceed 5% (v/v) glycerol. Due to the high molecular weight of the lambda vector, the contents may be very viscous. It is important to microcentrifuge the contents of the lambda vector tube briefly at 11,000 × g, then gently mix the solution by stirring with a yellow pipet tip prior to pipetting.

1. Set up a test ligation as follows to ligate the test insert into the ZAP Express vector:

   1.0 μl of the ZAP Express vector (1 μg/μl)
   0.8 μl of pRheo test insert (0.2 μg)
   0.5 μl of 10× ligase buffer
   0.5 μl of 10 mM rATP (pH 7.5)
   1.7 μl of water

   Then add

   0.5 μl of T4 DNA ligase (4 U/μl)

2. To prepare the sample ligation, add the following components:

   1.0 μl of the ZAP Express vector (1 μg/μl)
   X μl of sample insert
   0.5 μl of 10× ligase buffer
   0.5 μl of 10 mM rATP (pH 7.5)
   X μl of water for a final volume of 5 μl

   Then add

   0.5 μl of T4 DNA ligase (4 U/μl)

3. Incubate the reaction tubes overnight at 12–14°C or for up to 2 days at 4°C.

4. When ligating the insert, use a volume up to 2.5 μl. Use an equal molar ratio (or less to prevent multiple inserts) of the insert. The ZAP Express vector can accommodate inserts ranging from 0 to 12 kb. The ZAP Express vector is ~38,900 bp in length. If ligating a 4,000-bp insert to the vector, use 0.1 μg of insert for every 1 μg of vector. If the insert used is free from contaminants and contains a high percentage of ligatable ends, expect about 1 × 10^6–1.5 × 10^7 recombinant plaques when using high-efficiency packaging extracts, such as Gigapack III Plus or Gigapack III Gold packaging extracts [Catalog #200204 (Gigapack III Plus-4), #200205 (Gigapack III Plus-7), and #200206 (Gigapack III Plus-11) #200201 (Gigapack III Gold-4), #200202 (Gigapack III Gold-7), and #200203 (Gigapack III Gold-11)].
PACKAGING AND TITERING

Packaging Extracts

Packaging extracts are used to package recombinant lambda phage with high efficiency, which increases the size of gene libraries.

Gigapack III Gold packaging extract increases the efficiency and representation of libraries constructed from highly methylated DNA. The packaging extracts are restriction minus (HsdR⁻ McrA⁻ McrBC⁻ McrF⁻ Mrr⁻) to optimize packaging efficiency and library representation. When used in conjunction with restriction-deficient plating cultures, Gigapack III Gold packaging extract should improve the quality of DNA libraries constructed from methylated DNA.⁵⁻⁸

Optimal packaging efficiencies are obtained with lambda DNAs that are concatameric. Ligations should be carried out at DNA concentrations of 0.2 μg/μl or greater, which favors concatamers and not circular DNA molecules that only contain one cos site. DNA to be packaged should be relatively free from contaminants. DNA may be used directly from ligation reactions in most cases; however, polyethylene glycol (PEG), which is contained in some ligase buffers, has been shown to inhibit packaging. The volume of DNA added to each extract should be <5 μl.

Undigested wild-type lambda DNA will be packaged with efficiencies exceeding 1 × 10⁹ plaques/μg of vector when using Gigapack III Gold packaging extract. Predigested arms, when ligated to a test insert, will yield ~5 × 10⁶ recombinant plaques/μg of vector.

Packaging Protocol

1. Remove the appropriate number of packaging extracts from the ~80°C freezer and place the extracts on dry ice.

2. Quickly thaw the packaging extract by holding the tube between your fingers until the contents of the tube just begins to thaw.

3. Add the experimental DNA immediately (1–4 μl containing 0.1–1.0 μg of ligated DNA) to the packaging extract.

4. Stir the tube with a pipet tip to mix well. Gentle pipetting is allowable provided that air bubbles are not introduced.

5. Spin the tube quickly (for 3–5 seconds), if desired, to ensure that all contents are at the bottom of the tube.
6. Incubate the tube at room temperature (22°C) for 2 hours. **Do not exceed 2 hours.**

   **Note**  *The highest efficiency occurs between 90 minutes and 2 hours. Efficiency may drop dramatically during extended packaging times.*

7. Add 500 μl of SM buffer to the tube.

8. Add 20 μl of chloroform and mix the contents of the tube gently.

9. Spin the tube briefly to sediment the debris.

10. The supernatant containing the phage is now ready to be titered and may be stored at 4°C.

**Titering the Packaging Reaction**

**Preparing the Host Bacteria**

**Note**  *The VCS257 strain is for use with the Gigapack III Gold packaging extract and the positive wild-type lambda DNA control only.*

1. Streak the XL1-Blue MRF' and VCS257 cells onto LB agar plates containing the appropriate antibiotic (See *Recommended Media*). Incubate the plates overnight at 37°C.

   **Note**  *Do not add antibiotic to the medium in the following step. The antibiotic will bind to the bacterial cell wall and will inhibit the ability of the phage to infect the cell.*

2. Prepare separate 50-ml cultures of XL1-Blue MRF' and VCS257 cells in LB broth with supplements.

3. Incubate with shaking at 37°C for 4–6 hours (do not grow past an OD₆₀₀ of 1.0). Alternatively, grow overnight at 30°C, shaking at 200 rpm.

   **Note**  *The lower temperature keeps the bacteria from overgrowing, thus reducing the number of nonviable cells. Phage can adhere to nonviable cells resulting in a decreased titer.*

4. Pellet the bacteria at 1000 × g for 10 minutes.

5. Gently resuspend each cell pellet in 25 ml sterile 10 mM MgSO₄.

   **Note**  *For later use, store the cells at 4°C overnight in 10 mM MgSO₄.*
Titering Protocol

1. Dilute the XL1-Blue MRF’ cells (from step 5 of Preparing the Host Bacteria in Titering the Packaging Reaction) to an OD₆₀₀ of 0.5 with sterile 10 mM MgSO₄.

   **Note**  The bacteria should be used immediately following dilution.

2. To determine the titer of the packaged ligation product, mix the following components:

   - 1 μl of the final packaged reaction
   - 200 μl of XL1-Blue MRF’ cells at an OD₆₀₀ of 0.5

   and

   - 1 μl of a 1:10 dilution of the final packaged reaction
   - 200 μl of XL1-Blue MRF’ cells at an OD₆₀₀ of 0.5

3. Incubate the phage and the bacteria at 37°C for 15 minutes to allow the phage to attach to the cells.

4. Add 3 ml of NZY top agar, melted and cooled to ~48°C, and plate immediately onto dry, prewarmed NZY agar plates. Allow the plates to set for 10 minutes. Invert the plates and incubate at 37°C.

5. Plaques should be visible after 6–8 hours. Count the plaques and determine the titer in plaque-forming units per milliliter (pfu/ml).

   **Note**  If insert size is crucial, one may excise a few clones at this step or a few clones prior to single-clone amplification (see Single-Clone Excision Protocol in In Vivo Excision Protocols Using EXAssist Interference-Resistant Helper Phage with XLOLR Strain).
Testing the Efficiency of the Gigapack III Gold Packaging Extract with the Wild-Type Lambda Control DNA (Optional)

Use the following procedure to test the efficiency of Gigapack III Gold packaging extract with the λcI857 Sam7 wild-type lambda control DNA.

1. Thaw the frozen wild-type lambda control DNA on ice and gently mix the control after thawing.

2. Using 1 μl of the wild-type lambda control DNA (~0.2 μg), proceed with steps 1–10 in the Packaging Protocol.

   **Note** Because of the high titer achieved with the wild-type lambda control DNA, stop the control packaging reaction with 1 ml of SM buffer. This should make the plaques easier to count.

3. Prepare two consecutive 10⁻² dilutions of the packaging reaction from step 10 in the Packaging Protocol in SM buffer. (The final dilution is 10⁻⁴).

4. Add 10 μl of the 10⁻⁴ dilution to 200 μl of the VCS257 host strain at an OD₆₀₀ of 0.5. (This strain is recommended for plating the wild-type lambda control DNA only.) Incubate at 37°C for 15 minutes. Add 3 ml of NZY top agar, melted and cooled to ~48°C, and quickly pour the dilution onto dry, prewarmed NZY agar plates.

5. Incubate the plates for at least 12 hours at 37°C.

6. Count the plaques. Approximately 400 plaques should be obtained on the 10⁻⁴ dilution plate with Gigapack III Gold packaging extract.

7. Calculate the efficiency using the following equation:

\[
\text{Efficiency} = \frac{\text{Number of plaques} \times \text{dilution factor} \times \text{total packaging volume}}{\text{Total number of micrograms packaged} \times \text{number of microliters plated}}
\]
PLATING FOR BLUE–WHITE COLOR SELECTION

A background test can be completed by plating several hundred plaques on a plate (see Color Selection by IPTG and X-gal). Add 15 μl of 0.5 M IPTG (in water) and 50 μl of 250 mg/ml X-gal [in dimethylformamide (DMF)] to 2–3 ml of NZY top agar (48°C). The higher concentrations of IPTG–X-gal used in the plating often result in the formation of a precipitate, which disappears after incubation. Add the IPTG and X-gal to the NZY top agar separately, with mixing in between additions, to minimize the formation of this precipitate. Plate immediately on NZY agar plates. Plaques are visible after incubation for 6–8 hours at 37°C. Background plaques are blue, while recombinant plaques are white (clear).

1. To plate the packaged ligation product, mix the following components:

   1 μl of the final 500 μl packaged reaction
   200 μl of XL1-Blue MRF’ cells at an OD_{600} of 0.5

   and

   1 μl of a 1:10 dilution of packaged reaction
   200 μl of XL1-Blue MRF’ cells at an OD_{600} of 0.5

   **Note** Use of any other cell line may result in a dramatically reduced titer. XL1-Blue MRF’ is a RecA– McrA– and McrCB– Mrr– strain and does not restrict methylated DNA.

2. Incubate the phage and the bacteria at 37°C for 15 minutes to allow the phage to attach to the cells. (Best results are obtained with gentle shaking.)

3. Add the following components:

   2–3 ml of NZY top agar (48°C)
   15 μl of 0.5 M IPTG (in water)
   50 μl of X-gal [250 mg/ml (in DMF)]

   **Note** The increased concentrations of IPTG/X-gal used in the plating can result in the formation of a precipitate, but it will disappear after incubation.

4. Plate immediately onto the NZY agar plates and allow the plates to set for 10 minutes. Invert the plates and incubate at 37°C.

5. Plaques should be visible after 6–8 hours, although color detection requires overnight incubation. Background plaques are blue and should be <1 × 10⁵ pfu/μg of arms, while recombinant plaques will be white (clear) and should be 10- to 100-fold above the background. If the results of the test titer of the packaged sample ligation and test ligation give expected results, package the remainder of the sample ligation and titer.
AMPLIFYING THE ZAP EXPRESS LIBRARY

It is usually desirable to amplify libraries prepared in lambda vectors to make a large, stable quantity of a high-titer stock of the library. However, more than one round of amplification is not recommended, since slower growing clones may be significantly underrepresented. The following protocol is recommended for amplifying the ZAP Express library.

Day 1

1. Grow a 50-ml overnight culture of XL1-Blue MRF’ cells in LB broth with supplements at 30°C with shaking.

Day 2

2. Gently spin down the XL1-Blue MRF’ cells (1000 x g). Resuspend the cell pellet in 25 ml of 10 mM MgSO4. Measure the OD600 of the cell suspension, then dilute the cells to an OD600 of 0.5 in 10 mM MgSO4.

3. Combine aliquots of the packaged mixture or library suspension containing ~5 x 10^4 pfu of bacteriophage with 600 μl of XL1-Blue MRF’ cells at an OD600 of 0.5 in 14-ml BD Falcon® polypropylene round-bottom tubes. To amplify 1 x 10^6 plaques, use a total of 20 aliquots (each aliquot contains 5 x 10^4 plaques/150-mm plate).

   Note   Do not add more than 300 μl of phage/600 μl of cells.

4. Incubate the tubes containing the phage and host cells for 15 minutes at 37°C to allow the phage to attach to the cells.

5. Mix 6.5 ml of NZY top agar, melted and cooled to ~48°C, with each aliquot of infected bacteria and spread evenly onto a freshly poured 150-mm NZY agar plate. Allow the plates to set for 10 minutes.

6. Invert the plates and incubate at 37°C for 6–8 hours. Do not allow the plaques to get larger than 1–2 mm. On completion, the plaques should be touching.

7. Overlay the plates with ~8–10 ml of SM buffer. Store the plates at 4°C overnight (with gentle rocking if possible). This allows the phage to diffuse into the SM buffer.
Day 3

8. Recover the bacteriophage suspension from each plate and pool it into a sterile polypropylene container. Rinse the plates with an additional 2 ml of SM buffer and pool. Add chloroform to a 5% (v/v) final concentration. Mix well and incubate for 15 minutes at room temperature.

9. Remove the cell debris by centrifugation for 10 minutes at 500 × g.

10. Recover the supernatant and transfer it to a sterile polypropylene container. If the supernatant appears cloudy or has a high amount of cell debris, repeat steps 8 and 9. If the supernatant is clear, add chloroform to a 0.3% (v/v) final concentration and store at 4°C. Store aliquots of the amplified library in 7% (v/v) DMSO at −80°C.

11. Check the titer of the amplified library using host cells and serial dilutions of the library. (Assume ~10⁹–10¹¹ pfu/ml.)

Note Briefly spin the lambda phage stock to ensure that the chloroform is separated completely before removing the aliquot for titering.

PERFORMING PLAQUE LIFTS

1. Titer the amplified mixture or library suspension to determine the concentration using XL1-Blue MRF´ cells.

2. Combine the equivalent of 5 × 10⁴ pfu/plate and 600 μl of freshly prepared XL1-Blue MRF´ cells at an OD₆₀₀ of 0.5.

3. Incubate the bacteria and phage mixture at 37°C for 15 minutes to allow the phage to attach to the cells.

4. Add 6.5 ml of NZY top agar (~48°C) to the bacteria and phage mixture.

5. Quickly pour the plating culture onto a dry, prewarmed 150-mm NZY agar plate, which is at least 2 days old. Carefully swirl the plate to distribute the cells evenly. Allow the plates to set for 10 minutes. (Use 20 plates to screen 1 × 10⁶ pfu.)

6. Invert the plates and incubate at 37°C for ~8 hours.

7. Chill the plates for 2 hours at 4°C to prevent the top agar from sticking to the nitrocellulose membrane.

Note Use forceps and wear gloves for the following steps.
8. Place a nitrocellulose membrane onto each NZY agar plate for 2 minutes to allow the transfer of the phage particles to the membrane. Use a needle to prick through the membrane and agar for orientation. (If desired, waterproof ink in a syringe needle may be used.)

**Notes** If making duplicate nitrocellulose membranes, allow the second membrane to transfer for ~4 minutes.

*Pyrex® dishes are convenient for the following steps. All solutions should be at room temperature.*

9. Blot briefly on a Whatman® 3MM paper.

10. Crosslink the DNA to the membranes using the autocrosslink setting on the Stratalinker UV crosslinker* (120,000 μJ of UV energy) for ~30 seconds. Alternatively, oven bake at 80°C for ~1.5–2 hours.

11. Store the stock agar plates of the transfers at 4°C to use after screening.

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* Catalog #400071 (1800 model) or #400075 (2400 model).*
HYBRIDIZING AND SCREENING

Following the preparation of the membranes for hybridization, perform prehybridization, probe preparation, hybridization, and washes for either oligonucleotide probes or double-stranded probes and then expose the membranes to film as outlined in standard methodology texts. Following these procedures, perform secondary and tertiary screenings also as outlined in the standard methodology texts. When using the ZAP Express vector, perform in vivo excision on the isolates to obtain the insert-containing pBK-CMV phagemid vector (see In Vivo Excision of the pBK-CMV from the ZAP Express Vector and In Vivo Excision Protocols Using ExAssist Helper Phage with XLOLR Strain).

ANTIBODY SCREENING PROTOCOL IN Escherichia coli

A complete instruction manual for immunoscreening is supplied with the picoBlue immunoscreening kit. This kit is available with goat anti-rabbit antibodies or goat anti-mouse antibodies [picoBlue immunoscreening kit, Catalog #200371 (goat anti-rabbit) and #200372 (goat anti-mouse)].

EUKARYOTIC SCREENING WITH THE ZAP EXPRESS LIBRARY

Screening libraries in eukaryotic cells has proved to be an effective way of identifying clones otherwise nonidentifiable in prokaryotic screening systems. The screening technique used will depend on the clone of interest and on the type of assay available. An appropriate cell line for screening must be obtained, and an assay or reagent capable of identifying the cell or cells expressing the desired target protein must be developed. Three different techniques are available: selection, panning, and functional analysis of clone pools.

Selective Assay

Devising a selective assay for eukaryotic library screening requires a cell line that can grow in nonselective media and where expression of a transfected gene permits growth in selective media. An example of this method is screening for a thymidine kinase (TK) gene in L-TK– cells. If TK– cells are grown in HAT media, only those cells transfected with a clone coding for a protein capable of replacing TK will grow.

Panning Assay

Clone identification by "panning" requires the transfection of a library into a cell line deficient in the desired surface protein. When the clone of interest is translated and expressed on the surface of eukaryotic cells, the translated protein product is made accessible to an antibody, ligand, or receptor coupled either directly or indirectly to a solid-phase matrix. Eukaryotic transfectant clones expressing the appropriate insert will bind to the affinity matrix, while cells not adhering are washed away. Either transient or stable transfection protocols can be used.
Functional Assay

Functional assay screening can also be performed on either transient or stably transfected cells. Transient expression will likely require subdividing the amplified library into smaller pools of clones to prevent the dilution of a positive cell signal with an excess of negative clones. Each clone pool is amplified separately and transfected into the eukaryotic cells. The transfected cells are then tested for the expression of the desired clone. Once a pool is identified as containing the clone of interest, it is subdivided into smaller pools for a second round of prokaryotic amplification, eukaryotic transfection, and screening. After several rounds of enriching for the desired clone, a single clone can be isolated. The initial pool size is determined according to the sensitivity of the available assay so that a single clone within the pool is still theoretically detectable in the transfected cells. For example, if a positive assay signal is 1000-fold above background, pools containing 500–1000 members should still give a signal above background. The sensitivity of the assay dictates the initial size of the pools, as well as the number of pools required to screen. If stable transformants are created using G418 selection, pools of stable clones can be assayed. This simplifies the identification of isolated positive eukaryotic clones, because the eukaryotic colonies can be picked or diluted in microtiter tissue culture plates.

After a clone has been identified within the eukaryotic cells, the clone can be retrieved by several methods. Plasmid DNA within the tissue culture cells can be collected using the Hirt and Birnboin and Doly procedures, then transferred into E. coli cells for amplification and plasmid DNA preparations. Simmons et al. were able to screen libraries in COS cells, where the presence of the SV40 T antigen increases the copy number of phagemids containing the SV40 origin of replication. This results in a higher episomal copy number, which may help in the retrieval of the plasmids. Inserts can also be isolated by polymerase chain reaction (PCR) amplification of the tissue culture cells using T3/T7 primer sets. The resulting PCR fragment can be digested using restriction sites flanking the insert, then recloned into pBK-CMV phagemid DNA for further analysis.

Note Screening libraries in eukaryotic cells can be extremely laborious. Many functional assays are not sensitive enough to detect a clone from pools of nonrelated clones. Therefore, it is worth considering the use of techniques, such as differential PCR, selective hybridization, and degenerate oligonucleotides, to develop DNA probes for initial screening using prokaryotic plaques. Positive clones can then be screened by eukaryotic transfection and expression.
Protocol

Libraries are constructed in the ZAP Express vector (see Ligating the Insert and Packaging Protocol), and the libraries are titered to determine size (see Titering the Packaging Reaction). Amplify the library as described in Amplifying the ZAP Express Library. Eukaryotic screening can be performed with cesium-banded, double-stranded phagemid DNA prepared from the excised library (see In Vivo Excision of the pBK-CMV Phagemid Vector from the ZAP Express Vector). The library can be introduced into the eukaryotic cells as separate pools or as an entire library, depending on the assay system.

**EUKARYOTIC EXPRESSION**

The CMV promoter is considered to be a strong promoter and to function in many different cell lines. However, expression in eukaryotic cells is sensitive to many factors. If little or no expression is observed in eukaryotic cells, several factors can be considered.

1. For library screening where the insert may not be full length, the lacZ ATG allows the expression and detection of fusion protein. However, the 5’-untranslated sequences and amino-terminal fusion can affect expression levels of some inserts. If the clone is identified by prokaryotic screening techniques and is known to be full length (i.e., containing its own ATG and Kozak sequence), the prokaryotic 5’ UT sequences may be removed by digesting the excised phagemid with Nhe I and Spe I, by religating, and by screening for clones which lost the 200-bp fragment. Other cloning strategies may be used if these sites exist within the insert. Removal of this region has been shown to increase expression levels in some inserts. This effect may be due to increased specific activity of the expressed protein by eliminating the expression of fusion proteins, increased RNA stability, or increased translation efficiency by removing the competing upstream lacZ ATG.

2. Methylation of some insert DNA can prevent expression in some cell lines.

3. The promoter may not be functional in some cell lines and should be tested before screening a library.
In Vivo Excision of the PBK-CMV Phagemid Vector from the ZAP Express Vector

The ZAP Express vector is designed to allow simple, efficient in vivo excision and recircularization of any cloned insert contained within the lambda vector to form a phagemid containing the cloned insert. This in vivo excision depends on the placement of the DNA sequences within the lambda phage genome and on the presence of a variety of proteins, including filamentous (e.g., M13) bacteriophage-derived proteins. The M13 phage proteins recognize a region of DNA normally serving as the f1 bacteriophage "origin of replication." This origin of replication can be divided into two overlying parts: (1) the site of initiation and (2) the site of termination for DNA synthesis. These two regions are subcloned separately into the ZAP Express vector. The lambda phage (target) is made accessible to the M13-derived proteins by simultaneously infecting a strain of *E. coli* with both the lambda vector and the M13 helper phage.

Inside *E. coli*, the "helper" proteins (i.e., proteins from M13 phage) recognize the initiator DNA that is within the lambda vector. One of these proteins then nicks one of the two DNA strands. At the site of this nick, new DNA synthesis begins and duplicates whatever DNA exists in the lambda vector "downstream" (3') of the nicking site. DNA synthesis of a new single strand of DNA continues through the cloned insert until a termination signal, positioned 3' of the initiator signal, is encountered within the constructed lambda vector. The ssDNA molecule is circularized by the gene II product from the M13 phage, forming a circular DNA molecule containing the DNA between the initiator and terminator. In the case of the ZAP Express vector, this includes all sequences of the PBK-CMV phagemid vector and the insert, if one is present. This conversion is the "subcloning" step, since all sequences associated with normal lambda vectors are positioned outside of the initiator and terminator signals and are not contained within the circularized DNA. In addition, the circularizing of the DNA automatically recreates a functional f1 origin as found in f1 bacteriophage or phagemids.

Signals for "packaging" the newly created phagemid are linked to the f1 origin sequence. The signals permit the circularized ssDNA to be "packaged" into phagemid particles and secreted from the *E. coli*. Following secretion of the phagemid particle, the *E. coli* cells used for in vivo excision of the cloned DNA are killed, and the lambda phage is lysed by heat treatment at 70°C. The phagemid is not affected by the heat treatment. *Escherichia coli* is infected with the phagemid and can be plated on selective media to form colonies. DNA from excised colonies can be used for analysis of insert DNA, including DNA sequencing, subcloning, and mapping. Colonies from the excised PBK-CMV phagemid vector can also be used for subsequent production of ssDNA suitable for dideoxy-sequencing and site-specific mutagenesis.
In Vivo Excision Protocols Using ExAssist Helper Phage with XLOLR Strain

The ExAssist helper phage with XLOLR strain is designed to efficiently excise the pBK-CMV phagemid vector from the ZAP Express vector, while eliminating problems associated with helper phage co-infection. The ExAssist helper phage contains an amber mutation that prevents replication of the helper phage genome in a nonsuppressing *E. coli* strain such as XLOLR cells. This allows only the excised phagemid to replicate in the host, removing the possibility of co-infection from the ExAssist helper phage. Since the ExAssist helper phage cannot replicate in the XLOLR strain, single-stranded rescue cannot be performed in this strain using this helper phage.

Mass excision can be used to generate subtraction libraries and subtracted DNA probes. Converting the library to the phagemid form also allows screening of the phagemid library in eukaryotic cells by transformation of eukaryotic cells with supercoiled plasmid DNA.

**Single-Clone Excision Protocol**

**Day 1**

1. Core the plaque of interest from the agar plate and transfer the plaque to a sterile microcentrifuge tube containing 500 μl of SM buffer and 20 μl of chloroform. Vortex the microcentrifuge tube to release the phage particles into the SM buffer. Incubate the microcentrifuge tube for 1–2 hours at room temperature or overnight at 4°C. (This phage stock is stable for up to 6 months at 4°C.)

2. Grow separate 50-ml overnight cultures of XL1-Blue MRF’ and XLOLR cells in LB broth with supplements at 30°C.

**Day 2**

3. Gently spin down the XL1-Blue MRF’ and XLOLR cells (1000 × g). Resuspend each of the cell pellets in 25 ml of 10 mM MgSO₄. Measure the OD₆₀₀ of the cell suspensions, then adjust the concentration of the cells to an OD₆₀₀ of 1.0 (8 × 10⁸ cells/ml) in 10 mM MgSO₄.

4. Combine the following components in a 14-ml BD Falcon polypropylene tube:

   - 200 μl of XL1-Blue MRF’ cells at an OD₆₀₀ of 1.0
   - 250 μl of phage stock (containing >1 × 10⁴ phage particles)
   - 1 μl of the ExAssist helper phage (>1 × 10⁶ pfu/μl)

**Note**  Briefly spin the lambda phage stock to ensure that the chloroform is separated completely before removing the aliquot used in the excision reaction.
5. Incubate the polypropylene tube at 37°C for 15 minutes to allow the phage to attach to the cells.

6. Add 3 ml of LB broth with supplements and incubate the polypropylene tube for 2.5–3 hours at 37°C with shaking. Because clonal representation is not relevant, single-clone excision reactions can be safely performed overnight.

**Note** The turbidity of the media is not indicative of the success of the excision.

7. Heat the polypropylene tube at 65–70°C for 20 minutes to lyse the lambda phage particles and the cells. Spin the tube at 1000 × g for 15 minutes to pellet the cell debris.

8. Decant the supernatant into a sterile 14-ml BD Falcon polypropylene tube. This stock contains the excised pBK-CMV phagemid packaged as filamentous phage particles and may be stored at 4°C for 1–2 months.

9. To plate the excised phagemids, add 200 μl of freshly grown XLOLR cells from step 3 (OD₆₀₀ = 1.0) to two 1.5-ml microcentrifuge tubes. Add 100 μl of the phage supernatant (from step 8 above) to one microcentrifuge tube and 10 μl of the phage supernatant to the other microcentrifuge tube.

10. Incubate the microcentrifuge tubes at 37°C for 15 minutes.

11. Add 300 μl of NZY broth and incubate the tubes at 37°C for 45 minutes to allow sufficient expression of the kanamycin-resistance gene product prior to plating on selective medium.

12. Plate 200 μl of the cell mixture from each microcentrifuge tube on LB-kanamycin agar plates (50 μg/ml) and incubate the plates overnight at 37°C.

Due to the high-efficiency of the excision process, it may be necessary to titrate the supernatant to achieve single-colony isolation.

Colonies appearing on the plate contain the pBK-CMV double-stranded phagemid vector with the cloned DNA insert. Helper phage will not grow, since helper phage is unable to replicate in Su⁻ (nonsuppressing) XLOLR strain and does not contain kanamycin-resistance genes. XLOLR cells are also resistant to lambda phage infection, thus preventing lambda phage contamination after excision.

To maintain the pBK-CMV phagemid vector, streak the colony on a new LB–kanamycin agar plate. For long-term storage, prepare a bacterial glycerol stock and store at –80°C.

R408 helper phage is recommended for the single-stranded rescue procedure. (See Appendix: Recovery of Single-Stranded DNA from Cells Containing the pBK-CMV Phagemid Vector for the protocol.)
**Mass Excision Protocol**

**Day 1**

1. Grow separate 50-ml overnight cultures of XL1-Blue MRF\(^{-}\) and XLOLR cells in LB broth with supplements at 30\(^{\circ}\)C.

**Day 2**

2. Gently spin down the XL1-Blue MRF\(^{-}\) and XLOLR cells (1000 \(\times\) g). Resuspend each of the cell pellets in 25 ml of 10 mM MgSO\(_4\). Measure the OD\(_{600}\) of the cell suspensions, then adjust the concentration of the cells to an OD\(_{600}\) of 1.0 (8 \(\times\) 10\(^8\) cells/ml) in 10 mM MgSO\(_4\).

3. In a 50-ml conical tube, combine a portion of the amplified lambda bacteriophage library with XL1-Blue MRF\(^{-}\) cells at a MOI of 1:10 lambda phage-to-cell ratio. Excise 10- to 100-fold more lambda phage than the size of the primary library to ensure statistical representation of the excised clones. Add ExAssist helper phage at a 10:1 helper phage-to-cells ratio to ensure that every cell is co-infected with lambda phage and helper phage.

For example, use

- 10\(^7\) pfu of the lambda phage (i.e., 10- to 100-fold above the primary library size)
- 10\(^8\) XL1-Blue MRF\(^{-}\) cells (1:10 lambda phage-to-cell ratio, noting that an OD\(_{600}\) of 1.0 corresponds to 8 \(\times\) 10\(^8\) cells/ml)
- 10\(^9\) pfu of ExAssist helper phage (10:1 helper phage-to-cells ratio)

**Note** Briefly spin the lambda phage stock to ensure that the chloroform is separated completely before removing the aliquot used in the excision reaction.

4. Incubate the conical tube at 37\(^{\circ}\)C for 15 minutes to allow the phage to attach to the cells.

5. Add 20 ml of LB broth with supplements and incubate the conical tube for 2.5–3 hours at 37\(^{\circ}\)C with shaking.

**Notes** Incubation times for mass excision in excess of 3 hours may alter the clonal representation.

The turbidity of the media is not indicative of the success of the excision.

6. Heat the conical tube at 65–70\(^{\circ}\)C for 20 minutes to lyse the lambda phage particles and the cells.

7. Spin the conical tube at 1000 \(\times\) g for 10 minutes to pellet the cell debris and then decant the supernatant into a sterile conical tube.
8. To titer the excised phagemids, combine 1 μl of this supernatant with 200 μl of XLOLR cells from step 2 in a 1.5-ml microcentrifuge tube.

9. Incubate the microcentrifuge tube at 37°C for 15 minutes.

10. Add 40 μl of 5× NZY broth (for a final concentration of 1×) and incubate the tube at 37°C for 45 minutes to allow sufficient expression of the kanamycin-resistance gene product prior to plating on selective medium.

11. Plate 100 μl of the cell mixture onto LB–kanamycin agar plates (50 μg/ml) and incubate the plates overnight at 37°C.

   **Note**  *It may be necessary to further dilute the cell mixture to achieve single-colony isolation.*

Colonies may now be selected for plasmid preps, or the cell mixture may be plated directly onto filters for colony screening.
APPENDIX: RECOVERY OF SINGLE-STRANDED DNA FROM CELLS CONTAINING THE pBK-CMV PHAGEMID VECTOR

The pBK-CMV vector is a phagemid that can be secreted as single-stranded DNA (ssDNA) in the presence of M13 helper phage. These phagemids contain the intergenic (IG) region of a filamentous f1 phage. This region encodes all of the cis-acting functions of the phage required for packaging and replication. In E. coli with the F+ phenotype (containing an F’ episome), pBK-CMV phagemid vectors will be secreted as single-stranded f1 "packaged" phage when the bacteria have been infected by a helper phage. Because these filamentous helper phages (M13, f1) will not infect E. coli without an F’ episome coding for pili, it is essential to use the XL1-Blue MRF’ strain or a similar strain containing the F’ episome.21,22

We offer helper phage that preferentially package the pBK-CMV phagemid vector. Typically, 30–50 pBK-CMV molecules are packaged per helper phage DNA molecule. The pBK-CMV phagemid vector is offered with the IG region in the minus orientation.

Yields of ssDNA can depend on the specific insert sequence, but for most inserts >1 μg of ssDNA can be obtained from a 1.5-ml miniprep if grown in XL1-Blue MRF’. A faint single-stranded helper phage band may appear on a gel at ~4 kb for R408 helper phage. This DNA mixture can be sequenced with primers that are specific for the pBK-CMV phagemid vectors and do not hybridize to the helper phage genome.

R408 helper phage can be used to produce a large amount of single-stranded pBK-CMV phagemid vector. Use the ExAssist interference-resistant helper phage with the XLOLR strain for the excision of the pBK-CMV phagemid vector from the ZAP Express vector. Use the R408 helper phage for single-stranded rescue.
Single-Stranded Rescue Protocol

1. Inoculate a single colony into 5 ml of 2× YT broth§ containing 50 μg/ml kanamycin and R408 helper phage at 10⁷–10⁸ pfu/ml (MOI ~10).

2. Grow the culture at 37°C with vigorous aeration for 16–24 hours, or until growth has reached saturation.

3. Centrifuge 1.5 ml of the cell culture for 5 minutes in a microcentrifuge.

4. Remove 1 ml of the supernatant to a fresh tube, then add 150 μl of a solution containing 20% PEG8000 and 2.5 M NaCl. Allow phage particles to precipitate on ice for 15 minutes.

   **Note**  For increased yield, perform the PEG precipitation overnight at 4°C.

5. Centrifuge for 5 minutes in a microcentrifuge. (A pellet should be obvious.)

6. Remove supernatant. Centrifuge the PEG pellets a few seconds more to collect residual liquid, then remove and discard the residual liquid.

7. Resuspend the pellet in 400 μl of 0.3 M NaOAc (pH 6.0) and 1 mM EDTA by vortexing vigorously.

8. Extract with 1 volume phenol–chloroform and centrifuge for 1–2 minutes to separate phases.

9. Transfer the aqueous phase to a fresh tube and add 1 ml of ethanol. Centrifuge for 5 minutes.

10. Remove ethanol and dry the DNA pellet.

11. Dissolve the pellet in 25 μl of TE buffer.§

12. Analyze 1–2 μl on an agarose gel.

§ See Preparation of Media and Reagents.
## Troubleshooting

### Packaging

<table>
<thead>
<tr>
<th>Observations</th>
<th>Suggestions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Packaging efficiency is too low</td>
<td>Gigapack III packaging extract is very sensitive to slight variations in temperature; therefore, store the packaging extracts at the bottom of a −80°C freezer and avoid transferring tubes from one freezer to another</td>
</tr>
<tr>
<td></td>
<td>Do not allow the packaging extracts to thaw</td>
</tr>
<tr>
<td></td>
<td>Avoid use of ligase buffers containing PEG, which can inhibit packaging</td>
</tr>
<tr>
<td></td>
<td>Ensure the DNA concentration is sufficient. Ligate at DNA concentrations of 0.2 μg/μl or greater and package between 1 and 4 μl of the ligation reaction</td>
</tr>
<tr>
<td></td>
<td>Never package &gt;4 μl of the ligation reaction, which causes dilution of the proteins contained within the packaging extract</td>
</tr>
<tr>
<td>Neither a bacterial lawn nor plaques is observed on the plate when titering</td>
<td>The lambda phage stock aliquot used when determining titer and amplifying the library cannot contain chloroform, as chloroform lyases the <em>E. coli</em> cells. Briefly spin the lambda phage stock to ensure that the chloroform is separated completely before removing the aliquot</td>
</tr>
<tr>
<td>or amplifying the library</td>
<td></td>
</tr>
</tbody>
</table>

### Excision

<table>
<thead>
<tr>
<th>Observations</th>
<th>Suggestions</th>
</tr>
</thead>
<tbody>
<tr>
<td>The number of colonies is too low</td>
<td>Verify that the titer on the tubes is current and correct and use only calibrated pipettors. The molar ratios of lambda phage to cells to helper phage is critical</td>
</tr>
<tr>
<td></td>
<td>If an excision is unsuccessful, prepare a high-titer stock of the phage and repeat the excision procedure, as excision efficiencies are directly related to the ZAP Express phage titer</td>
</tr>
<tr>
<td></td>
<td>Ensure that the platings are performed using agar plates containing kanamycin</td>
</tr>
<tr>
<td></td>
<td>The lambda phage stock aliquot used for in vivo excision cannot contain chloroform, as chloroform lyases the <em>E. coli</em> cells. Briefly spin the lambda phage stock to ensure that the chloroform is separated completely before removing the aliquot</td>
</tr>
</tbody>
</table>
## PREPARATION OF MEDIA AND REAGENTS

<table>
<thead>
<tr>
<th>LB Agar (per Liter)</th>
<th>LB Broth (per Liter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 g of NaCl</td>
<td>10 g of NaCl</td>
</tr>
<tr>
<td>10 g of tryptone</td>
<td>10 g of tryptone</td>
</tr>
<tr>
<td>5 g of yeast extract</td>
<td>5 g of yeast extract</td>
</tr>
<tr>
<td>20 g of agar</td>
<td></td>
</tr>
<tr>
<td>Add deionized H₂O to a final volume of 1 liter</td>
<td>Add deionized H₂O to a final volume of 1 liter</td>
</tr>
<tr>
<td>Adjust pH to 7.0 with 5 N NaOH</td>
<td>Adjust to pH 7.0 with 5 N NaOH</td>
</tr>
<tr>
<td>Autoclave</td>
<td>Autoclave</td>
</tr>
<tr>
<td>Pour into petri dishes (≈25 ml/100-mm plate)</td>
<td>Autoclave</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LB Broth with Supplements</th>
<th>LB–Kanamycin Agar (per Liter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prepare 1 liter of LB broth</td>
<td>1 liter of LB agar</td>
</tr>
<tr>
<td>Autoclave</td>
<td>Autoclave</td>
</tr>
<tr>
<td>Add the following filter-sterilized supplements prior to use</td>
<td>Cool to 55°C</td>
</tr>
<tr>
<td>10 ml of 1 M MgSO₄</td>
<td>Add 7.5 ml of 10 mg/ml kanamycin (filter-sterilized)</td>
</tr>
<tr>
<td>3 ml of a 2 M maltose solution or</td>
<td>Pour into petri dishes (≈25 ml/100-mm plate)</td>
</tr>
<tr>
<td>10 ml of 20% (w/v) maltose</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LB–Tetracycline Agar (per Liter)</th>
<th>LB–Tetracycline Broth (per Liter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 liter of LB agar</td>
<td>Prepare 1 liter of LB broth</td>
</tr>
<tr>
<td>Autoclave</td>
<td>Autoclave</td>
</tr>
<tr>
<td>Cool to 55°C</td>
<td>Cool to 55°C</td>
</tr>
<tr>
<td>Add 1.5 ml of 10 mg/ml tetracycline (filter-sterilized)</td>
<td>Add 1.25 ml of 10 mg/ml-filter-sterilized tetracycline</td>
</tr>
<tr>
<td>Pour into petri dishes (≈25 ml/100-mm plate)</td>
<td>Store broth in a dark, cool place as tetracycline is light-sensitive</td>
</tr>
<tr>
<td>Store plates in a dark, cool place or cover plates with foil if left out at room temperature for extended time periods as tetracycline is light-sensitive</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>NZY Agar (per Liter)</th>
<th>NZY Broth (per Liter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 g of NaCl</td>
<td>5 g of NaCl</td>
</tr>
<tr>
<td>2 g of MgSO₄ • 7H₂O</td>
<td>2 g of MgSO₄ • 7H₂O</td>
</tr>
<tr>
<td>5 g of yeast extract</td>
<td>5 g of yeast extract</td>
</tr>
<tr>
<td>10 g of NZ amine (casein hydrolysate)</td>
<td>10 g of NZ amine (casein hydrolysate)</td>
</tr>
<tr>
<td>15 g of agar</td>
<td>Add deionized H₂O to a final volume of 1 liter</td>
</tr>
<tr>
<td>Add deionized H₂O to a final volume of 1 liter</td>
<td>Adjust the pH to 7.5 with NaOH</td>
</tr>
<tr>
<td>Adjust the pH to 7.5 with NaOH</td>
<td>Adjust the pH to 7.5 with NaOH</td>
</tr>
<tr>
<td>Pour into petri dishes (≈80 ml/150-mm plate)</td>
<td>Autoclave</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>NZY Broth (per Liter)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>5 g of NaCl</td>
<td></td>
</tr>
<tr>
<td>2 g of MgSO₄ • 7H₂O</td>
<td></td>
</tr>
<tr>
<td>5 g of yeast extract</td>
<td></td>
</tr>
<tr>
<td>10 g of NZ amine (casein hydrolysate)</td>
<td></td>
</tr>
<tr>
<td>Add deionized H₂O to a final volume of 1 liter</td>
<td></td>
</tr>
<tr>
<td>Adjust the pH to 7.5 with NaOH</td>
<td></td>
</tr>
<tr>
<td>Autoclave</td>
<td></td>
</tr>
<tr>
<td>NZY Top Agar (per Liter)</td>
<td>SM Buffer (per Liter)</td>
</tr>
<tr>
<td>--------------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td>1 liter of NZY broth</td>
<td>5.8 g of NaCl</td>
</tr>
<tr>
<td>Add 0.7% (w/v) agarose</td>
<td>2.0 g of MgSO₄ · 7H₂O</td>
</tr>
<tr>
<td>Autoclave</td>
<td>50.0 ml of 1 M Tris-HCl (pH 7.5)</td>
</tr>
<tr>
<td></td>
<td>5.0 ml of 2% (w/v) gelatin</td>
</tr>
<tr>
<td></td>
<td>Add deionized H₂O to a final volume of 1 liter</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>20× SSC Buffer (per Liter)</th>
<th>2× YT Broth (per Liter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>175.3 g of NaCl</td>
<td>10 g of NaCl</td>
</tr>
<tr>
<td>88.2 g of sodium citrate</td>
<td>10 g of yeast extract</td>
</tr>
<tr>
<td>800.0 ml of deionized H₂O</td>
<td>16 g of tryptone</td>
</tr>
<tr>
<td>10.0 N NaOH</td>
<td>Add deionized H₂O to a final volume of 1 liter</td>
</tr>
<tr>
<td>Adjust to pH 7.0 with a few drops of 10.0 N NaOH</td>
<td>Adjust to pH 7.5 with NaOH</td>
</tr>
<tr>
<td>Add deionized H₂O to a final volume of 1 liter</td>
<td>Autoclave</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TE Buffer</th>
<th>10× Ligase Buffer</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 mM Tris-HCl (pH 7.5)</td>
<td>500 mM Tris-HCl (pH 7.5)</td>
</tr>
<tr>
<td>1 mM EDTA</td>
<td>70 mM MgCl₂</td>
</tr>
<tr>
<td></td>
<td>10 mM dithiothreitol (DTT)</td>
</tr>
</tbody>
</table>

**Note**  
`rATP` is added separately in the ligation reaction.
REFERENCES


ENDNOTES

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MSDS INFORMATION

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