BaculoDirect™ GST Gateway® Expression Kit

For cloning and high-level expression of recombinant GST-fusion proteins using Gateway®-adapted Baculovirus DNA

Catalog nos. A10640, A10641
Rev. Date: 14 July 2010
Manual part no. A10607

MAN0000700
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# Kit Contents and Storage

## Types of Kits

This manual is supplied with the following kits.

<table>
<thead>
<tr>
<th>Product</th>
<th>Cat. no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaculoDirect™ GST Gateway® Transfection Kit</td>
<td>A10640</td>
</tr>
<tr>
<td>BaculoDirect™ GST Gateway® Expression Kit</td>
<td>A10641</td>
</tr>
</tbody>
</table>

## Kit Components

Each kit contains the components listed below. See the next page for a detailed description of other reagents supplied with each kit.

<table>
<thead>
<tr>
<th>Component</th>
<th>Cat. no. A10640</th>
<th>Cat. no. A10641</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaculoDirect™ N-GST Linear DNA</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Cellfectin® II Reagent</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>pENTR™-CAT Control Plasmid</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Ganciclovir</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Gateway® LR Clonase™ II Enzyme Mix for BaculoDirect™</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Sf21 Frozen Cells</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Grace’s Insect Cell Culture Medium, Unsupplemented</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>BaculoDirect™ GST Gateway® Expression Kit Manual</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Insect Cell Lines Manual</td>
<td></td>
<td>✓</td>
</tr>
</tbody>
</table>

## Shipping/Storage

The BaculoDirect™ GST Gateway® Transfection and Expression Kits are shipped as described below. Upon receipt, store the components as detailed. All components are guaranteed for six months if stored properly.

<table>
<thead>
<tr>
<th>Item</th>
<th>Shipping</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaculoDirect™ GST Gateway® Transfection Kit</td>
<td>Gel ice</td>
<td>4°C, except pENTR™-CAT: –20°C</td>
</tr>
<tr>
<td>Gateway® LR Clonase™ II Enzyme Mix for BaculoDirect™</td>
<td>Dry ice</td>
<td>–20°C</td>
</tr>
<tr>
<td>Ganciclovir</td>
<td>Dry ice</td>
<td>–20°C</td>
</tr>
<tr>
<td>Sf21 Frozen Cells</td>
<td>Dry ice</td>
<td>Liquid nitrogen</td>
</tr>
<tr>
<td>Grace’s Insect Cell Culture Medium, Unsupplemented</td>
<td>Room Temperature</td>
<td>4°C, protected from light</td>
</tr>
</tbody>
</table>

Continued on next page
## Transfection Kit Components

The BaculoDirect™ GST Gateway® Transfection Kits include the following components, sufficient to perform 5 reactions. Store components as detailed below.

<table>
<thead>
<tr>
<th>Item</th>
<th>Composition</th>
<th>Amount</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaculoDirect™ N-GST Linear DNA, linearized with Bsu36 I</td>
<td>300 ng per tube in 10 μl of TE buffer, pH 8.0*</td>
<td>6 tubes</td>
<td>4°C</td>
</tr>
<tr>
<td>Cellfectin® II Reagent</td>
<td>1 mg/ml in membrane-filtered water</td>
<td>125 μl</td>
<td>4°C</td>
</tr>
<tr>
<td>pENTR™-CAT Control Plasmid</td>
<td>40 μl of 0.5 ng/μl vector in TE buffer, pH 8.0</td>
<td>20 μg</td>
<td>–20°C</td>
</tr>
<tr>
<td>Ganciclovir</td>
<td>100 mM in deionized water</td>
<td>50 μl</td>
<td>–20°C, protected from light</td>
</tr>
</tbody>
</table>

*TE buffer, pH 8.0: 10 mM Tris-HCl, 1 mM EDTA, pH 8.0

## Expression Kit Components

The BaculoDirect™ GST Expression Kits include the following components. Store components as detailed below.

<table>
<thead>
<tr>
<th>Item</th>
<th>Composition</th>
<th>Amount</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaculoDirect™ N-GST Linear DNA, linearized with Bsu36 I</td>
<td>300 ng per tube in 10 μl of TE buffer, pH 8.0</td>
<td>6 tubes</td>
<td>4°C</td>
</tr>
<tr>
<td>Cellfectin® II Reagent</td>
<td>1 mg/ml in membrane-filtered water</td>
<td>125 μl</td>
<td>4°C</td>
</tr>
<tr>
<td>pENTR™-CAT Control Plasmid</td>
<td>40 μl of 0.5 ng/μl vector in TE buffer, pH 8.0</td>
<td>20 μg</td>
<td>–20°C</td>
</tr>
<tr>
<td>Ganciclovir</td>
<td>100 mM in deionized water</td>
<td>50 μl</td>
<td>–20°C, protected from light</td>
</tr>
<tr>
<td>Gateway® LR Clonase™ II Enzyme Mix for BaculoDirect™ Kits</td>
<td>–</td>
<td>40 μl</td>
<td>–20°C for up to 6 months; (–80°C for long-term storage)</td>
</tr>
<tr>
<td>Sf21 Frozen Cells</td>
<td>1 × 10⁷ cells/ml in: 60% complete TNM-FH 30% FBS 10% DMSO</td>
<td>1 ml</td>
<td>Liquid nitrogen</td>
</tr>
<tr>
<td>Grace’s Insect Cell Culture Medium, Unsupplemented</td>
<td>Sterile-filtered medium contains L-glutamine</td>
<td>500 ml</td>
<td>4°C, protected from light</td>
</tr>
</tbody>
</table>
Accessory Products

**Additional Products**

Many of the reagents supplied with the BaculoDirect™ GST Gateway® Transfection and Expression Kits as well as other products suitable for use with the kits are available separately from Invitrogen. Ordering information is provided below.

<table>
<thead>
<tr>
<th>Product</th>
<th>Amount</th>
<th>Cat. no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gateway® LR Clonase II Enzyme Mix for BaculoDirect®</td>
<td>10 reactions</td>
<td>11791-023</td>
</tr>
<tr>
<td>Cellfectin® II Reagent</td>
<td>1 ml</td>
<td>10362-100</td>
</tr>
<tr>
<td>Grace’s Insect Cell Culture Medium, Unsupplemented</td>
<td>500 ml</td>
<td>11595-030</td>
</tr>
<tr>
<td>Grace’s Insect Cell Culture Medium, Supplemented</td>
<td>500 ml</td>
<td>11605-094</td>
</tr>
<tr>
<td>Sf-900 II SFM</td>
<td>500 ml</td>
<td>10902-096</td>
</tr>
<tr>
<td>Sf-900™ III SFM</td>
<td>500 ml</td>
<td>12658-019</td>
</tr>
<tr>
<td>Sf-900 Medium (1.3X)</td>
<td>100 ml</td>
<td>10967-032</td>
</tr>
<tr>
<td>Express Five® SFM</td>
<td>1000 ml</td>
<td>10486-025</td>
</tr>
<tr>
<td>Penicillin-Streptomycin</td>
<td>20 ml</td>
<td>15140-148</td>
</tr>
<tr>
<td>Fungizone® Antimycotic</td>
<td>20 ml</td>
<td>15290-018</td>
</tr>
<tr>
<td>Fetal Bovine Serum Qualified, Heat-Inactivated</td>
<td>100 ml</td>
<td>16140-063</td>
</tr>
<tr>
<td>Easy-DNA™ Kit</td>
<td>15–200 reactions</td>
<td>K1800-01</td>
</tr>
<tr>
<td>PureLink™ Genomic DNA Mini Kit</td>
<td>10 preps</td>
<td>K1820-00</td>
</tr>
<tr>
<td></td>
<td>50 preps</td>
<td>K1820-01</td>
</tr>
<tr>
<td></td>
<td>250 preps</td>
<td>K1820-02</td>
</tr>
<tr>
<td>PureLink™ HiPure Plasmid Miniprep Kit</td>
<td>25 preps</td>
<td>K2100-02</td>
</tr>
<tr>
<td>BaculoTiter™ Assay Kit</td>
<td>30 titers</td>
<td>K1270</td>
</tr>
<tr>
<td>4% Agarose Gel</td>
<td>40 ml</td>
<td>18300-012</td>
</tr>
<tr>
<td>β-Gal Staining Kit</td>
<td>1 kit</td>
<td>K1465-01</td>
</tr>
<tr>
<td>Bluo-gal</td>
<td>1 g</td>
<td>15519-028</td>
</tr>
<tr>
<td>CAT Antiserum</td>
<td>50 μl</td>
<td>R902-25</td>
</tr>
<tr>
<td>Sterile, cell culture grade, distilled water</td>
<td>500 ml</td>
<td>15230-162</td>
</tr>
<tr>
<td>Proteinase K</td>
<td>100 mg</td>
<td>25530-015</td>
</tr>
<tr>
<td>UltraPure™ Glycogen</td>
<td>100 μl</td>
<td>10814-010</td>
</tr>
<tr>
<td>NuPAGE® LDS Sample Preparation Buffer (4X)</td>
<td>10 ml</td>
<td>NP0007</td>
</tr>
<tr>
<td></td>
<td>250 ml</td>
<td>NP0008</td>
</tr>
<tr>
<td>Novex® Tris-Glycine SDS Sample Buffer (2X)</td>
<td>20 ml</td>
<td>LC2676</td>
</tr>
</tbody>
</table>

*Continued on next page*
Accessory Products, continued

Insect Cells

Invitrogen offers a variety of insect cell lines for protein expression studies. We recommend using Sf9 or Sf21 cells to generate high-titer viral stocks with the BaculoDirect™ GST Gateway® Expression Kits. Once you have generated high-titer viral stocks, you may use Sf9, Sf21, High Five™, or Mimic™ Sf9 cells for protein expression studies. For more information, refer to www.invitrogen.com or contact Technical Support (page 45).

<table>
<thead>
<tr>
<th>Item</th>
<th>Amount</th>
<th>Cat. no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sf9 Frozen Cells</td>
<td>1 ml tube, 1 × 10⁷ cells/ml</td>
<td>B825-01</td>
</tr>
<tr>
<td>Sf21 Frozen Cells</td>
<td>1 ml tube, 1 × 10⁷ cells/ml</td>
<td>B821-01</td>
</tr>
<tr>
<td>High Five™ Cells</td>
<td>1 ml tube, 3 × 10⁶ cells/ml</td>
<td>B855-02</td>
</tr>
<tr>
<td>Mimic™ Sf9 Insect Cells</td>
<td>1 ml tube, 1 × 10⁷ cells/ml</td>
<td>12552-014</td>
</tr>
</tbody>
</table>

Gateway® Entry Vectors

A variety of Gateway® entry vectors are available from Invitrogen. Depending on your application, you may choose entry vectors with specific features such as a ribosome binding site (RBS). For more information on the Gateway® cloning technology, as well as the features and vector maps of available Gateway® entry vectors, refer to www.invitrogen.com or contact Technical Support (page 45).

<table>
<thead>
<tr>
<th>Item</th>
<th>Amount</th>
<th>Cat. no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>pENTR™/TEV/D-TOPO® Cloning Kit</td>
<td>20 reactions</td>
<td>K2535-20</td>
</tr>
<tr>
<td>pENTR™/D-TOPO® Cloning Kit</td>
<td>20 reactions</td>
<td>K2400-20</td>
</tr>
</tbody>
</table>

Note: A selection guide for choosing the most appropriate Gateway® entry vector for your application can be found on our website at www.invitrogen.com/Gateway.

Continued on next page
You may use western blot analysis to detect and affinity chromatography on glutathione agarose to purify your recombinant fusion protein that is expressed in frame with the N-terminal peptide containing the GST (glutathione S-transferase) tag. You may also use Dynabeads® complexed with anti-GST antibodies to isolate your recombinant protein. For more information, refer to www.invitrogen.com or contact Technical Support (page 45).

<table>
<thead>
<tr>
<th>Item</th>
<th>Amount</th>
<th>Cat. no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-glutathione S-transferase, rabbit IgG fraction, 3 mg/ml</td>
<td>0.5 ml</td>
<td>A-5800</td>
</tr>
<tr>
<td>Anti-glutathione S-transferase, rabbit IgG fraction, Alexa Fluor® 488 conjugate, 2 mg/ml</td>
<td>0.5 ml</td>
<td>A-11131</td>
</tr>
<tr>
<td>Glutathione Transferase Fusion Protein Purification Kit</td>
<td>5 purifications</td>
<td>G-21801</td>
</tr>
<tr>
<td>Glutathione agarose, linked through sulfur (sedimented bead suspension)</td>
<td>10 ml</td>
<td>G-2879</td>
</tr>
<tr>
<td>Purification Columns (10 ml polypropylene columns)</td>
<td>50</td>
<td>R640-50</td>
</tr>
<tr>
<td>Dynabeads® M-280 Sheep anti-Rabbit IgG</td>
<td>2 ml</td>
<td>112-03D</td>
</tr>
<tr>
<td>Dynabeads® Protein A</td>
<td>2 ml</td>
<td>100-01D</td>
</tr>
<tr>
<td>Dynabeads® Protein G</td>
<td>2 ml</td>
<td>100-03D</td>
</tr>
<tr>
<td>WesternBreeze® Chromogenic Kit, Anti-Mouse</td>
<td>1 kit</td>
<td>WB7103</td>
</tr>
<tr>
<td>WesternBreeze® Chromogenic Kit, Anti-Goat</td>
<td>1 kit</td>
<td>WB7107</td>
</tr>
<tr>
<td>WesternBreeze® Chromogenic Kit, Anti-Rabbit</td>
<td>1 kit</td>
<td>WB7105</td>
</tr>
<tr>
<td>WesternBreeze® Chemiluminescent Kit, Anti-Mouse</td>
<td>1 kit</td>
<td>WB7104</td>
</tr>
<tr>
<td>WesternBreeze® Chemiluminescent Kit, Anti-Goat</td>
<td>1 kit</td>
<td>WB7106</td>
</tr>
<tr>
<td>WesternBreeze® Chemiluminescent Kit, Anti-Rabbit</td>
<td>1 kit</td>
<td>WB7108</td>
</tr>
<tr>
<td>WesternBreeze® Blocker/Diluent (part A and B)</td>
<td>80 ml each</td>
<td>WB7050</td>
</tr>
<tr>
<td>WesternBreeze® Wash Solution (16X)</td>
<td>2 × 100ml</td>
<td>WB7003</td>
</tr>
</tbody>
</table>
Introduction

Overview

Introduction
The BaculoDirect™ GST Gateway® Transfection and Expression Kits use Gateway® Technology to facilitate direct transfer of the gene of interest into the baculovirus genome in vitro without the need for additional cloning or recombination in bacterial or insect cells. The resulting recombinant baculovirus DNA is transfected directly into insect cells to generate recombinant virus and to screen for expression. The ability to clone and express genes from baculovirus without plaque purification or selection in bacteria makes the BaculoDirect™ GST Gateway® Transfection and Expression Kits the fastest procedure for baculovirus expression.

Advantages of the BaculoDirect™ GST Gateway® Kits
Using the BaculoDirect™ GST Gateway® Transfection and Expression Kits to obtain purified recombinant virus suitable for production of high titer stocks offers the following advantages:

- Saves time by allowing rapid cloning of the gene of interest into the baculovirus genome without the need for traditional homologous recombination or site-specific transposition methods.
- Produces high-level expression of a GST-tagged recombinant protein for easy detection and purification.
- Linearized baculovirus DNA and ganciclovir selection inhibits replication of non-recombinant virus, and eliminates the need for plaque purification.
- The GST tag is helpful for solubilization of the overexpressed protein of interest as it prevents the fusion protein from being sequestered into inclusion bodies.

BaculoDirect™ N-GST Linear DNA
The major features of the BaculoDirect™ N-GST Linear DNA include:

- attR1 and attR2 sites for recombinational cloning of the gene of interest from a Gateway® entry clone
- Herpes simplex virus thymidine kinase gene (HSV1 tk) located between the two attR sites for negative selection using ganciclovir
- lacZ gene located between the two attR sites for determination of viral purity using β-galactosidase staining
- N-terminal GST fusion tag for detection and purification of recombinant fusion proteins

Note: If you are planning on expressing a secreted protein, be aware that the presence of the GST-tag may interfere with secretion.

Continued on next page
Overview, continued

The Gateway® Technology

Gateway® Technology is a universal cloning method based on the site-specific recombination properties of bacteriophage lambda (Landy, 1989). The Gateway® Technology provides a rapid and highly efficient way to move DNA sequences into multiple vector systems for functional analysis and protein expression. To produce recombinant baculovirus using the BaculoDirect™ GST Gateway® Expression Kit, simply:

1. Clone the gene of interest into a Gateway® entry vector of choice to create an entry clone.
2. Perform an LR recombination reaction to transfer the gene of interest from the entry clone to the BaculoDirect™ N-GST Linear DNA.
3. Transfect insect cells with recombinant baculovirus DNA and harvest recombinant baculovirus.

For more detailed information about Gateway® Technology, generating an entry clone, and performing the LR recombination reaction, refer to the Gateway® Technology with Clonase™ II manual (part no. 25-0749). This manual is available for downloading from www.invitrogen.com or by contacting Technical Support (page 45).

Important

The BaculoDirect™ GST Gateway® Expression System is designed to help you construct a recombinant baculovirus to deliver and express a gene of interest in insect cells. Use of this system is geared towards those users who are familiar with the principles of baculovirus expression systems and Gateway® Technology. We highly recommend that users possess a working knowledge of viral and insect cell culture techniques.

For more information about the baculovirus life cycle, viral structure, and laboratory techniques, refer to the following published reviews: King and Possee, 1992; O’Reilly et. al., 1992; and Richardson et. al., 1995.

Insect Cell Lines Manual

Before starting baculoviral expression experiments, we recommend that users refer to the Insect Cell Lines manual (part no. 25-0127) for additional information on insect cell culture. This manual contains information on:

- Thawing frozen cells
- Maintaining and passaging cells
- Freezing cells
- Scaling up cell culture

This manual is provided with the BaculoDirect™ GST Gateway® Expression Kits and is also available from www.invitrogen.com or by contacting Technical Support (page 45).
The Gateway® Technology

Introduction

The Gateway® Technology is based on the bacteriophage lambda site-specific recombination system which facilitates the integration of lambda into the *E. coli* chromosome and the switch between the lytic and lysogenic pathways (Ptashne, 1992). In the Gateway® Technology, the components of the lambda recombination system are modified to improve the specificity and efficiency of the system (Bushman *et al.*, 1985). This section provides a brief overview of the Gateway® Technology. For detailed information, refer to the Gateway® Technology with Clonase™ II manual (part no. 25-0749).

Characteristics of Recombination Reactions

Lambda integration into the *E. coli* chromosome occurs via intermolecular DNA recombination that is mediated by a mixture of lambda and *E. coli*-encoded proteins (i.e., LR Clonase™ II Enzyme Mix for BaculoDirect™). The hallmarks of lambda recombination are listed below. For more detailed information about lambda recombination, see published references and reviews (Landy, 1989; Ptashne, 1992).

- Recombination occurs between specific (att) sites on interacting DNA molecules.
- Recombination is conservative (i.e., there is no net gain or loss of nucleotides) and does not require DNA synthesis. The DNA segments flanking the recombination sites are switched, such that after recombination, the att sites are hybrid sequences comprised of sequences donated by each parental vector. For example, attP sites are comprised of sequences from attR and attL sites.
- Strand exchange occurs within a core region that is common to all att sites.

att Sites

Lambda recombination occurs between site-specific attachment (att) sites: attB on the *E. coli* chromosome and attP on the lambda chromosome. The att sites serve as the binding site for recombination proteins and have been well characterized (Weisberg & Landy, 1983). Upon lambda integration, recombination occurs between attB and attP sites to give rise to attL and attR sites. The actual crossover occurs between homologous 15 bp core regions on the two sites, but surrounding sequences are required as they contain the binding sites for the recombination proteins (Landy, 1989).

Continued on next page
The Gateway® Technology, continued

**Gateway® LR Recombination Reaction**

By using the BaculoDirect™ GST Gateway® Transfection Kit (available separately or as a part of the BaculoDirect™ GST Expression Kit), you will take advantage of the LR reaction to transfer your gene of interest into the BaculoDirect™ N-GST Linear DNA. The LR reaction facilitates recombination of an `attL` substrate (entry clone) with an `attR` substrate (BaculoDirect™ N-GST Linear DNA) to create an `attB`-containing expression virus (see diagram below). This reaction is catalyzed by LR Clonase™ II Enzyme Mix for BaculoDirect™.

![Diagram showing the LR reaction process](image)

- **entry clone**
  - `attL` gene
- **BaculoDirect™ N-GST Linear DNA**
  - `attR` TK gene lacZ

LR Clonase™ II for BaculoDirect™

- **expression virus**
  - `attB` gene
- **by-product**
  - `attP` TK gene lacZ
Ganciclovir is a nucleoside analog [9-(1,3-Dihydroxy-2-propoxymethyl) guanine] that is enzymatically phosphorylated by Herpes Simplex Virus type 1 thymidine kinase (HSV1 tk). Once phosphorylated, the active analog incorporates into DNA and inhibits DNA replication (Rubsam et al., 1999). Ganciclovir selection has been used in Sf9 cells to purify recombinant viruses that have lost the counter-selectable gene marker (HSV1 tk) due to homologous recombination (Godeau et al., 1992).

In the BaculoDirect™ GST Gateway® Transfection and Expression Kits, the HSV1 tk gene is under the control of an immediate early promoter (Pie-1(0)) which drives synthesis of the first viral transcript produced in infected cells (Kovacs et al., 1991).

Ganciclovir is a hazardous material that may cause harm if ingested, inhaled, or absorbed through the skin. Exercise caution and wear suitable protective clothing, gloves, and safety goggles while handling solutions containing ganciclovir. Before handling ganciclovir, review the Safety Data Sheet available from our website at www.invitrogen.com/sds or by contacting Technical Support (page 45).
Experimental Overview

The following diagram summarizes the general steps required to express your gene of interest (GOI) using the BaculoDirect™ GST Gateway® Expression Kit.

1. Generate a Gateway® entry clone containing your gene of interest

2. Perform an LR reaction between the entry clone and the BaculoDirect™ N-GST Linear DNA to generate recombinant baculovirus DNA.

3. Directly transfet insect cells with recombinant baculovirus DNA and select with ganciclovir. Collect P1 viral stock.

4. Screen for recombinant protein expression (optional)

5. Infect insect cells with P1 viral stock to amplify virus and select with ganciclovir. Collect P2 viral stock.

6. Screen for recombinant protein expression (required)

7. Infect insect cells with P2 viral stock to amplify and scale-up virus. Collect P3 viral stock.

The experimental steps necessary to express your protein of interest using the BaculoDirect™ GST Gateway® Transfection and Expression Kits are outlined below. For more details on each step, refer to the indicated pages.

<table>
<thead>
<tr>
<th>Step</th>
<th>Action</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Generate an entry clone containing your gene of interest.</td>
<td>8</td>
</tr>
<tr>
<td>2</td>
<td>Perform the LR recombination reaction between the BaculoDirect™ N-GST Linear DNA and an entry clone containing your gene of interest using BaculoDirect™ GST Gateway® Transfection Kit</td>
<td>9–10</td>
</tr>
<tr>
<td>3</td>
<td>Directly transfect insect cells with recombinant baculovirus DNA and collect P1 viral stock. Screen for recombinant protein expression, if desired.</td>
<td>11–16</td>
</tr>
<tr>
<td>4</td>
<td>Infect insect cells with P1 viral stock to generate a high-titer viral stock. Screen for recombinant protein expression.</td>
<td>17–19</td>
</tr>
<tr>
<td>5</td>
<td>Determine titer of viral stock by plaque assay.</td>
<td>20–23</td>
</tr>
<tr>
<td>6</td>
<td>Isolate recombinant viral DNA and analyze by PCR, if desired.</td>
<td>24–28</td>
</tr>
<tr>
<td>7</td>
<td>Infect insect cells and optimize conditions for recombinant protein expression.</td>
<td>29–32</td>
</tr>
<tr>
<td>8</td>
<td>Purify recombinant protein, if desired.</td>
<td>33</td>
</tr>
</tbody>
</table>
Methods

Before Starting

Introduction
Before you start your experiments, you will need to have an entry clone containing your gene of interest, cultures of Sf9 or Sf21 cells growing, and frozen master stocks available. Refer to the guidelines below for more information.

Gateway® Entry Vectors
To recombine your gene of interest into the BaculoDirect™ N-GST Linear DNA, you will need an entry clone containing your gene of interest. For your convenience, Invitrogen offers a variety of Gateway® entry vectors (see page vii). A selection guide for choosing the most appropriate Gateway® entry vector for your application can be found on our website at www.invitrogen.com/Gateway. For detailed information on constructing an entry clone, refer to the manual for the specific entry vector you are using.

Points to Consider for BaculoDirect™ N-GST Linear DNA
Keep the following points in mind when constructing your entry clone to be used in the recombination reaction with the BaculoDirect™ N-GST Linear DNA:

- Design your gene of interest to be in frame with the N-terminal GST tag after recombination. Refer to page 28 for a diagram of the recombinant baculovirus DNA.

  Tip: Keep the translation reading frame of your protein of interest in frame with the AAA AAA triplet in the attL1 site of the entry clone.

- Make sure your insert contains a stop codon.

Recommended Cells
We recommend using Sf9 or Sf21 cells to generate high-titer viral stocks with the BaculoDirect™ GST Gateway® Expression Kits. Note that Sf21 cells are provided with the BaculoDirect™ GST Gateway® Expression Kits. We do not recommend using High Five™ cells to generate viral stocks due to lower transfection efficiency. Once you have generated high-titer viral stocks, you may use Sf9, Sf21, High Five™, or Mimic™ Sf9 cells for protein expression studies. See page vii for ordering information.

Recommended Media
For the highest transfection efficiency, we recommend using Grace’s Insect Cell Culture Medium, Unsupplemented (provided with the BaculoDirect™ GST Gateway® Expression Kits) for the transfection experiment. For infection, expression studies, and general culturing of insect cells, you may use any complete growth medium (e.g., Sf-900 II SFM, Sf-900™ III SFM, complete TNM-FH, or other suitable medium). Refer to page 40 for a recipe for complete TNM-FH.

When working with recombinant or wild-type viral stocks, always maintain separate media bottles for cell culture and for virus work. Baculovirus particles can survive and be maintained in media at 4°C and will contaminate your cell cultures if added to tissue culture plates or flasks during passaging.
Performing the LR Recombination Reaction

**Introduction**

After you have generated an entry clone using most appropriate Gateway® entry vector for your application, perform the LR recombination reaction to transfer the gene of interest into the BaculoDirect™ N-GST Linear DNA. We recommend that you include the pENTR™-CAT positive control supplied with the kit in your experiments to help you evaluate your results.

**Important**

The LR recombination reaction protocol provided on the next page contains optimized amounts of each reagent. To obtain the best possible results, follow the protocol exactly as described.

**LR Clonase™ II Enzyme Mix for BaculoDirect™**

LR Clonase™ II Enzyme Mix for BaculoDirect™ is supplied with the BaculoDirect™ GST Gateway® Expression Kits to catalyze the LR recombination reaction. The LR Clonase™ II Enzyme Mix for BaculoDirect™ combines the proprietary enzyme formulation and 5X LR Clonase™ Reaction Buffer. Use the protocol provided on the next page to perform the LR recombination reaction using LR Clonase™ II Enzyme Mix for BaculoDirect™.

*Note:* For the LR recombination reaction, use LR Clonase™ II Enzyme Mix for BaculoDirect™ only, do not use LR Clonase™ II or LR Clonase™ from other kits.

**Materials Needed**

- Purified plasmid DNA of your entry clone (50–150 ng/μl in TE buffer, pH 8.0)
  *Note:* Use PureLink™ HiPure Plasmid Prep Kit, not a silica-based miniprep kit, for the purification of the entry clone.
- BaculoDirect™ N-GST Linear DNA (300 ng/tube; provided with the BaculoDirect™ GST Gateway® Expression and Transfection Kits)
- pENTR™-CAT control plasmid, optional (100 ng/μl; provided with the kits)
- LR Clonase™ II Enzyme Mix for BaculoDirect™ (provided with the kits; keep at −20°C until immediately before use)
- 1X TE Buffer, pH 8.0 (10 mM Tris-HCl, pH 8.0, 1 mM EDTA)
- 25°C water bath

*Continued on next page*
Performing the LR Recombination Reaction, continued

**LR Recombination Reaction Protocol**

Perform Steps 1–4 in a **sterile laminar flow hood** to reduce the chances of contamination.

1. To set up your sample and positive control reaction, add the following components **directly** to the BaculoDirect™ N-GST Linear DNA tubes containing 10 μl (300 ng) of DNA at room temperature and mix the contents. **Do not vortex or pipette up and down as this will shear the baculovirus DNA and reduce transfection efficiency.**

<table>
<thead>
<tr>
<th>Component</th>
<th>Sample</th>
<th>Positive Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaculoDirect™ N-GST Linear DNA</td>
<td>10 μl (in tube)</td>
<td>10 μl (in tube)</td>
</tr>
<tr>
<td>pENTR™-CAT control (100 ng/μl)</td>
<td>–</td>
<td>1 μl</td>
</tr>
<tr>
<td>Entry clone (100–300 ng/reaction)</td>
<td>1–2 μl</td>
<td>–</td>
</tr>
<tr>
<td>1X TE Buffer, pH 8.0</td>
<td>4–5 μl</td>
<td>5 μl</td>
</tr>
</tbody>
</table>

**Total volume** 16 μl 16 μl

**Note:** To include a negative control, set up a third sample reaction and substitute 4 μl of 1X TE Buffer, pH 8.0 for the enzyme mix (see step 4).

2. Remove the LR Clonase™ II Enzyme Mix for BaculoDirect™ from −20°C and thaw on ice (~ 2 minutes).

3. Vortex the LR Clonase™ II Enzyme Mix for BaculoDirect™ briefly twice (~ 2 seconds each time).

4. To each sample above, add 4 μl of LR Clonase™ II Enzyme Mix for BaculoDirect™ or 4 μl of 1X TE Buffer, pH 8.0 (if preparing a negative control) for a total reaction volume of 20 μl.

5. Mix well by tapping the tube several times. **Do not vortex or pipette up and down as this will shear the baculovirus DNA and reduce transfection efficiency.**

6. Incubate the reactions at 25°C for 1 hour.

**Note:** Extending the incubation time up to 18 hours typically increases the efficiency of the LR recombination reaction.

---

**Note**

After incubation, you may analyze the LR reaction by PCR. Dilute a 2 μl aliquot of the LR reaction 200-fold and use 2 μl of the dilution in a 25 μl PCR reaction.

For the PCR amplification, you may use the Polyhedrin forward primer (5’-AAATGATAAACCATCTCGC-3’) and a primer of your own design that binds within your gene of interest.

See page 28 for Polyhedrin forward primer binding site. PCR reaction conditions must be optimized.

---

**The Next Step**

Once the LR reaction is completed, you are ready to directly transfect the recombinant baculovirus DNA into insect cells. Proceed to the next section for transfection guidelines.
Transfecting Sf9 or Sf21 Cells

Introduction
This section provides detailed guidelines for transfecting your LR recombination reaction into Sf9 or Sf21 insect cells. Sf21 cells are provided with the BaculoDirect™ GST Gateway® Expression Kits.

Cellfectin® II Reagent
Cellfectin® II Reagent is supplied with the BaculoDirect™ GST Gateway® Transfection and Expression Kits for lipid-mediated transfection of your insect cells. Cellfectin® II Reagent is a proprietary liposome formulation of a cationic lipid in membrane-filtered water, and is ideally suited for the transfection of Sf9 and Sf21 insect cells.

Serum-Free Medium
We recommend using Grace’s Insect Cell Culture Medium, Unsupplemented, however, you may use serum-free medium during the transfection experiment. Note that components in serum-free medium may interfere with transfection resulting in a decrease in transfection efficiency.

Note: If you are already culturing Sf9 or Sf21 cells in Sf-900 II SFM or Sf-900™ III SFM, you can perform the transfection in Grace’s Insect Cell Culture Medium, Unsupplemented, then easily switch back to Sf-900 II SFM or Sf-900™ III SFM after transfection.

Use of complete growth medium that contains antibiotics and antimycotics in addition to ganciclovir for the last step of the transfection protocol (see next page) is optional. If so desired, you can use 100 U/ml of penicillin, 100 μg/ml of streptomycin, and 0.25 μg/ml of amphotericin B (i.e., Fungizone™ Antimycotic) in this last step (see page vi for ordering information).

Preparing and Storing Ganciclovir
We recommend setting aside the amount of complete growth medium needed for the experiment requiring ganciclovir selection and adding the appropriate amount of ganciclovir to a final concentration of 100 μM. Aliquot the remaining ganciclovir into multiple tubes to reduce the number of freeze/thaw cycles.

Additional ganciclovir may be purchased in powder form from InvivoGen (Cat. no. sud-gcv). Refer to page 40 for instructions on reconstituting and storing ganciclovir.

Ganciclovir provided with the BaculoDirect™ GST Gateway® Transfection and Expression Kits has a concentration of 100 mM and might form a precipitate upon thawing. Incubating the ganciclovir at 37°C for 10 minutes and vortexing will redisolve the ganciclovir and eliminate the precipitate. If you experience ganciclovir precipitation, we recommend diluting the 100 mM stock solution in half with sterile distilled water before refreezing for storage.

Continued on next page
Transfecting Sf9 or Sf21 Cells, continued

Materials Needed

- Sf21 cells (provided with the BaculoDirect™ GST Gateway® Expression Kits) or Sf9 cells
- LR reaction from LR Recombination Reaction Protocol, page 10
- Grace’s Insect Cell Culture Medium, Unsupplemented (provided with the BaculoDirect™ GST Gateway® Expression Kits; also available separately, see page vi)
- Cellfectin® II Reagent (provided with the BaculoDirect™ GST Gateway® Transfection and Expression Kits; also available separately, see page vi)
- Complete growth medium (e.g., Sf-900 II SFM, or other suitable medium) with antibiotics and 100 μM ganciclovir (see Note on previous page)
- Six-well tissue culture plates
- 27°C incubator
- Air-tight bags or containers
- Inverted microscope

Note: All the reagents and cell lines necessary for transfection are also available separately from Invitrogen. For ordering information, see page vi–vii.

Controls

We recommend that you include the following controls in your experiments:

- LR recombination reaction using pENTR™-CAT plasmid as a positive control
- Cellfectin® II Reagent only (mock transfection) as a negative control

Important

Use Grace's Insect Medium, Unsupplemented which does not contain any supplements, FBS, or antibiotics for the transfection procedure. The proteins in the FBS and supplements will interfere with the Cellfectin® II Reagent, causing the transfection efficiency to decrease.

Continued on next page
Transfecting Sf9 or Sf21 Cells, continued

**Transfection Procedure**

For Sf9 or Sf21 insect cells cultured in Grace’s Insect Medium, Supplemented containing 10% FBS, use the following protocol to transfect your cells in a 6-well format. All amounts and volumes are given on a per well basis.

1. Verify that the Sf9 or Sf21 cells are in the log phase (1.5–2.5 × 10⁶ cells/ml) with greater than 95% viability.

2. If the cell density is in range of 1.5–2.5 × 10⁶ cells/ml and the culture is without antibiotics, proceed to step 2a. If the cell density is not in this range or the cell culture contains antibiotics, follow steps 2b–2c:
   a. Add 2 ml of Grace’s Insect Medium, Unsupplemented (without antibiotics and serum) in each well. Seed 8 × 10⁵ Sf9 or Sf21 cells from Step 1 per well. Do not change medium or wash the cells. The medium carried over will enhance the transfection efficiency. Allow cells to attach for 15 minutes at room temperature in the hood. Proceed to step 3.
   b. Prepare 10ml plating medium by mixing 1.5 ml Grace’s Insect Medium, Supplemented containing 10% FBS (without antibiotics) and 8.5 ml Grace’s Insect Medium, Unsupplemented (without FBS and antibiotics).
   c. Plate 8 × 10⁵ Sf9 or Sf21 cells from Step 1 per well. Allow cells to attach for 15 minutes at room temperature in the hood. Remove the medium. Add 2.5 ml plating medium from step 2b per well. Proceed to step 3.

3. Prepare the following solutions in 1.5 ml microcentrifuge tubes for each transfection sample. Cellfectin® II Reagent in the Transfection Mixture A can be left at room temperature for up to 30 minutes.

   **Transfection Mixture A:**
   - Cellfectin® II Reagent 8 µl
   - Grace’s Insect Medium, Unsupplemented (without supplements, serum, or antibiotics) 100 µl

   **Transfection Mixture B:**
   - LR recombination reaction 10 µl
   - Grace’s Insect Medium, Unsupplemented (without supplements, FBS, or antibiotics) 100 µl

Procedure continued on next page
Transfecting Sf9 or Sf21 Cells, continued

Transfection Procedure, continued

Procedure continued from previous page

4. Combine Transfection Mixture A and Transfection Mixture B. Mix gently by tapping the tube and incubate at room temperature for 25–35 minutes.

5. After 25–35 minutes incubation, add the transfection mix from Step 4 (total volume ~210 μl) dropwise onto the cells from step 2. Repeat for all transfections.
   Note: With Cellfectin® II, you do not have to remove the medium from cells and wash cells prior to adding the DNA:lipid complex to cells.

6. Incubate the cells in a 27°C incubator for 3 to 5 hours.

7. Remove the transfection mixture and replace with 2 ml of complete growth medium (e.g., Grace’s Insect Medium, Supplemented and 10% FBS) with 100 μM ganciclovir to each well. Addition of antibiotics is optional (see page 11). Repeat for all transfections.
   Note: Distribute the drops evenly to avoid disturbing the monolayer.

8. Place the plates in a sealed plastic bag with moist paper towels to prevent evaporation. Incubate the cells at 27°C for 72 hours or until you start to see signs of viral infection.
Isolating P1 Viral Stock

Introduction

Budded virus should be released into the medium 72 hours after transfection. However, depending on transfection efficiency, cells may not show all of the signs of viral infection for up to a week. Beginning at 72 hours after transfection, visually inspect the cells daily for signs of infection (see below). Once the cells appear infected, harvest the virus from the cell culture medium using the procedure below. You may also perform an initial screen for expression of your recombinant fusion protein, if desired.

Materials Needed

- Transfected insect cells from Step 8, previous page
- Inverted microscope
- 15 ml tubes

Characteristics of Infected Cells

Virus infected insect cells typically display the following characteristics as observed from visual inspection using an inverted phase microscope at 250–400X magnification.

<table>
<thead>
<tr>
<th>Time Period</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early (first 24 hours)</td>
<td>Increased cell diameter</td>
</tr>
<tr>
<td></td>
<td>Increased size of cell nuclei</td>
</tr>
<tr>
<td>Late (24–72 hours)</td>
<td>Cessation of cell growth</td>
</tr>
<tr>
<td></td>
<td>Detachment</td>
</tr>
<tr>
<td>Very Late (&gt;72 hours)</td>
<td>Cell lysis</td>
</tr>
</tbody>
</table>

Isolating P1 Viral Stock

1. Once the transfected cells demonstrate signs of very late stage infection (e.g., 72 hours post-transfection) collect 2 ml of medium from each well and transfer to sterile 15 ml tubes. Centrifuge the tubes at 3000–5000 rpm for 5 minutes to remove cells and large debris.
2. Transfer the supernatant to fresh 15 ml tubes. This is the P1 viral stock. Store at 4°C, protected from light. See the next page for additional storage information.
3. If you wish to screen for expression of your recombinant fusion protein, proceed to Screening for Expression, next page.

Continued on next page
Isolating P1 Viral Stock, continued

**Screening for Expression**

You may perform a small-scale or preliminary expression experiment on the transfected cells to verify expression of your recombinant protein. Follow the general guidelines below to assay for expression.

- If you are expressing a secreted protein, remove a sample from the medium to analyze protein expression and secretion. You may also harvest cells to analyze intracellular levels of your recombinant protein (see below).
- To harvest cells, transfer transfected cells into microcentrifuge tubes and centrifuge. Wash cells 2X with PBS to remove traces of serum.
- Assay for expression by Western blot analysis. For information on preparing protein samples and detecting expression, refer to pages 31–32.

**Storing Viral Stocks**

Store viral stocks as follows:

- If medium is serum-free, add serum to 10%. Serum proteins act as substrates for proteases and therefore prevent degradation of viral coat proteins.
- Store viral stock at 4°C, protected from light.
- Store an aliquot of the viral stock at –80°C.
- Do not store routinely used viral stocks at temperatures below 4°C. Repeated freeze/thaw cycles can result in a 10- to 100-fold decrease in viral titer.

**The Next Step**

Once you have obtained your P1 viral stock, you may:

- Amplify the viral stock by infecting Sf9 or Sf21 cells (refer to Preparing a High-Titer Viral Stock and Screening for Protein Expression, page 17). We recommend this procedure to obtain the highest viral titers and optimal results in your expression studies.
- Perform a plaque assay to amplify your viral stock from a single viral clone or to determine the titer of your P1 viral stock (refer to Performing a Plaque Assay, page 20).
Preparing a High-Titer Viral Stock and Screening for Protein Expression

Introduction

The P1 viral stock is a low-titer stock (1 × 10^5 to 1 × 10^6 pfu/ml). You will infect cells with the P1 stock to generate a high-titer P2 stock of approximately 5 × 10^7 to 1 × 10^8 pfu/ml. This P2 viral stock can then be used to generate a large-scale, high-titer viral stock suitable for expression studies. Guidelines are provided in this section to amplify the recombinant baculovirus and to screen for recombinant protein expression.

Materials Needed

- Sf21 cells (provided with the BaculoDirect™ GST Gateway® Expression Kits or available separately, see page vii) or Sf9 cells (available separately, see page vii for ordering information)
- Complete growth medium with 100 μM ganciclovir
- P1 viral stock (from Step 2, page 15)
- β-Gal Staining Kit (recommended; see page vi for ordering information) or other suitable kit
- Six-well tissue culture plates
- 27°C incubator
- Inverted microscope
- Air-tight bags or containers

You will infect duplicate samples of Sf9 or Sf21 cells with P1 viral stock. One set of cells will be assayed for the presence of non-recombinant virus by β-galactosidase staining. The other set will be assayed for expression of your recombinant protein.

Preparing Cells

Use log phase Sf9 or Sf21 cells with greater than 95% viability.

1. Seed 8 × 10^5 Sf9 or Sf21 cells per well in 2 ml of complete growth medium with 100 μM ganciclovir in a six-well tissue culture plate. Remember to seed duplicate wells (see Note above). Gently tip the plate from side to side 4–6 times to evenly distribute the cells.
2. Incubate the cells at 27°C for one hour to allow the cells to fully attach to the bottom of the plate.
3. Verify that the cells have attached by inspecting them under an inverted microscope.

Continued on next page
Preparing a High-Titer Viral Stock and Screening for Protein Expression, continued

### Isolating P2 Viral Stock

1. Add 5 μl of the P1 viral stock to each well. Place the plates in a sealed plastic bag with moist paper towels to prevent evaporation. Incubate infected cells for 72 hours at 27ºC.

2. At 72 hours post-infection, collect 2 ml of medium from each well and transfer to sterile 15 ml tubes. Centrifuge the tubes at 3,000–5,000 rpm for 5 minutes to remove debris.

3. Transfer the supernatant to fresh 15 ml tubes. **This is the P2 viral stock.**
   Store at 4ºC, protected from light. Refer to page 16 for additional storage information.

4. With one set of infected cells, proceed to β-Galactosidase Staining. With the other set of infected cells, proceed to Screening for Expression. We recommend performing both procedures before scaling up your viral stock and performing expression experiments.

### β-Galactosidase Staining

Because the BaculoDirect™ Linear DNA contains the *lacZ* gene, you may assay for the presence of non-recombinant virus by staining the infected cells for β-galactosidase expression. Recombinant virus will not stain blue because the gene of interest replaces the *lacZ* gene after the LR recombination reaction (see diagram on page 4). If you see blue-stained cells, we recommend that you perform a plaque assay to isolate a recombinant viral clone (see page 20).

### Screening for Expression

You will need to verify expression of your recombinant protein before further amping your viral stock. Follow the general guidelines below to assay for expression:

- If you are expressing a secreted protein, remove a sample from the medium to analyze protein expression and secretion. You may also harvest cells to analyze intracellular levels of your recombinant protein (see below).
  Note: The presence of the GST-tag on your recombinant protein may interfere with its secretion.

- To harvest cells, transfer transfected cells into microcentrifuge tubes and centrifuge to collect cells. Wash cells 2X with PBS to remove traces of serum.

- Assay for expression by western blot analysis. For information on preparing protein samples and detecting expression, refer to pages 31–32.
Preparing a High-Titer Viral Stock and Screening for Protein Expression, continued

Scaling Up the Amplification Procedure

If you are satisfied with the purity of the viral stock and have confirmed expression of your recombinant protein, you may scale-up the amplification procedure to any volume of your choice. To produce a large-scale, high-titer P3 stock, we recommend doing the following:

• Perform a plaque assay to determine the titer of the P2 viral stock (see next page).
• Use the equation provided below to determine the amount of P2 viral stock to use to infect at a specific MOI.
• Scale up the amount of cells and volume of virus appropriately and follow the guidelines outlined in this section.

Note that ganciclovir selection is not required for generation of the P3 viral stock.

Multiplicty of Infection (MOI)

To amplify your viral stock, infect cells at a multiplicity of infection (MOI) ranging from 0.1 to 1.0. MOI is defined as the number of virus particles per cell. Use the following formula to calculate how much viral stock to add to obtain a specific MOI.

\[
\text{Inoculum required (ml)} = \frac{\text{MOI (pfu/cell)} \times \text{number of cells}}{\text{viral titer (pfu/ml)}}
\]

Note: If you have not determined the titer of your P2 viral stock, you may assume that the titer ranges from \(5 \times 10^7\) to \(1 \times 10^8\) pfu/ml.

Generating High-Titer Stocks From Frozen Master Stock

If you start with a frozen viral master stock, we recommend generating a new high-titer stock as viral titer generally decreases from storage at \(-80^\circ\text{C}\). To generate another high-titer stock from the master stock, re-infect insect cells and amplify the viral stock using the guidelines outlined in this section.

The Next Step

Now that you have a high-titer viral stock, you will need to determine the titer of your viral inoculum. Proceed to the next section to perform a plaque assay and calculate viral titer.
Performing a Plaque Assay

Introduction
We recommend you perform a plaque assay to determine the titer of your viral stock. You may also perform a plaque assay to purify a single viral clone, if desired. In this procedure, you will infect cells with dilutions of your viral stock and identify focal points of infection (plaques) on an agarose overlay. You may also titer your viral stock by the end-point dilution method described in O'Reilly et. al., 1992.

BaculoTiter™ Assay Kit
We recommend using the BaculoTiter™ Assay Kit, available separately from Invitrogen, to determine the titer of your baculoviral stock. The BaculoTiter™ Assay Kit rapidly determines the titer of an unknown baculovirus sample with minimal handling steps, providing both accuracy and convenience in an easy-to-use kit format in two days as opposed to ten days with the serial dilution assays. See page vi for ordering information.

Blue/White Screening
You will use a chromogenic substrate to distinguish colorless plaques containing recombinant virus from blue plaques containing non-recombinant virus. We recommend using Bluo-gal instead of X-gal for blue/white screening because Bluo-gal generally produces a darker blue color than X-gal. Add Bluo-gal directly to the overlay solution before pouring over the infected cells.

Materials Needed
- Sf21 cells (provided with BaculoDirect™ GST Gateway® Expression Kits or available separately, see page vii) or Sf9 cells (available separately, see page vii)
- Sf-900 II, Sf-900™ III SFM, or other appropriate complete growth medium (see Note below)
- Sf-900 Medium (1.3X) or other appropriate plaquing medium (see Note below)
- 4% Agarose Gel
- Sterile, cell-culture grade, distilled water.
- 100 ml sterile, glass bottle
- Serial dilutions of viral stock (see page 21)
- Bluo-gal (50 mg/ml, see page 41 for a recipe)
- Six-well tissue culture plates
- Sterile hood
- Water baths at 47°C and 70°C
- 27°C incubator
- Inverted microscope
- Air-tight bags or containers

Continued on next page
Performing a Plaque Assay, continued

If you are culturing your Sf9 or Sf21 cells in serum-supplemented media (i.e., complete TNM-FH), you should have the following reagents on hand (see page vi for ordering information):

• Grace’s Insect Cell Culture Medium, Supplemented
• Grace’s Insect Cell Culture Medium (2X)
• Fetal Bovine Serum (FBS), Qualified, Heat-Inactivated

Diluting Virus

You will be infecting cells with serial dilutions of your viral stock. Keep in mind the following points when preparing the 10-fold serial dilutions:

• Prepare dilutions in complete growth medium
• Vortex viral stocks or dilutions before making the next dilution to ensure virus is evenly resuspended
• Prepare 3 ml (for duplicate wells) or 4 ml (for triplicate wells) of each viral dilution
• Make sure to return your viral stock to 4ºC
• Prepare dilutions according to the viral stock you are using to perform the plaque assay (see recommended dilutions below)

<table>
<thead>
<tr>
<th>Viral Stock</th>
<th>Dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>$10^{-1}, 10^{-2}, 10^{-3}, 10^{-4}, 10^{-5}, 10^{-6}$</td>
</tr>
<tr>
<td>P2</td>
<td>$10^{-3}, 10^{-4}, 10^{-5}, 10^{-6}, 10^{-7}, 10^{-8}$</td>
</tr>
<tr>
<td>P3</td>
<td>$10^{-4}, 10^{-5}, 10^{-6}, 10^{-7}, 10^{-8}, 10^{-9}$</td>
</tr>
</tbody>
</table>

Infecting Cells with Virus

The quality of the cell monolayer is critical for a successful plaque assay. Be sure to include a cells-only control to assess cell viability, contamination, and monolayer quality.

1. Seed $8 \times 10^5$ Sf9 or Sf21 cells per well in 2 ml complete growth medium in a six-well plate. Use 2 to 3 wells for each viral dilution to be tested (see Diluting Virus, above). Gently tip the plate from side to side 4–6 times to evenly distribute the cells.

2. Incubate the cells at 27ºC for one hour to allow the cells to fully attach to the bottom of the plate.

3. Verify that the cells have attached by inspecting them under an inverted microscope.

4. Aspirate the medium from the wells. Carefully add 1 ml of each viral dilution dropwise to the appropriately labeled well. Be careful not to disturb the monolayer.

5. Incubate the cells at 27ºC for 1 hour. While cells are incubating, prepare the Plaquing Medium (next page).

Continued on next page
Performing a Plaque Assay, continued

Preparing the Plaquing Medium

Plaquing medium, a mixture of culture medium and agarose, is used to immobilize the infected cells for the plaque assay. Prepare plaquing medium immediately before use. If you are culturing your cells in Sf-900 II SFM or Sf-900™ III SFM, prepare Sf-900 Plaquing Medium. If you are culturing cells in TNM-FH, prepare Grace’s Plaquing Medium.

Note: This procedure provides instructions to prepare 40 or 50 ml of Sf-900 and Grace’s Plaquing Medium, respectively. You will need 2 ml of Plaquing Medium per well. To prepare more Plaquing Medium, scale up the volume of reagents used accordingly.

1. Melt the 4% Agarose Gel by placing the bottle in a 70°C water bath for 20 to 30 minutes or heating the agarose in a microwave oven. While the 4% agarose gel is melting, place the following in the 45°C water bath:
   • Empty, sterile 100 ml bottle
   • Sf-900 Medium (1.3X) or Grace’s Insect Cell Culture Medium (2X), as appropriate

2. Once the 4% agarose gel has liquefied, move the agarose gel, medium, and empty 100 ml bottle to a sterile hood.

3. Working quickly, prepare the plaquing medium as follow:
   **Sf-900 Plaquing Medium:** Combine 30 ml of Sf-900 Medium (1.3X) and 10 ml of the melted 4% Agarose Gel in the empty 100 ml bottle and mix gently.
   **Grace’s Plaquing Medium:** Add 20 ml of heat-inactivated FBS to the 100 ml bottle of Grace’s Insect Medium (2X) and mix. Combine 25 ml of the Grace’s Insect Medium (2X) containing serum with 12.5 ml of cell-culture grade, sterile, distilled water and 12.5 ml of the melted 4% Agarose Gel in the empty 100 ml bottle and mix gently.

4. Add Bluo-gal to a final concentration of 150 μg/ml. Mix immediately by pipetting up and down. Place the bottle in a 47°C water bath until use.

Agarose Overlay

Pouring the agarose overlay may require some practice if you are unfamiliar with this technique. You should already have your plaquing medium prepared (see above; remove from water bath). It is important to work quickly and efficiently.

1. After the 1 hour incubation period (Step 5, previous page), remove the cells from the incubator and completely aspirate the medium from each well containing cells and virus. If you have multiple plates, follow this protocol for one plate before proceeding to the next. Do not let the cells dry out.

2. Withdraw 2 ml of the plaquing medium and slowly stream the solution down the side of the well. Repeat for all wells. **Do not move the plate until the agarose overlay has set.**

3. Repeat Steps 1–2 until all plates have been completed.

4. Place the plates in a sealed plastic bag with moist paper towels to prevent evaporation.
   **Note:** Once condensation appears on the plastic bag or container, open the bag or container. Moisture can destroy the monolayer, preventing plaque formation.

5. Incubate the cells at 27°C for 4–6 days or until plaques are well formed. Proceed to **Calculating Viral Titer**, next page.

Continued on next page
Performing a Plaque Assay, continued

Neutral Red Overlay

To improve visualization of plaques, you may perform a neutral red overlay 4 days post-infection. Do not use this procedure if you plan to plaque purify your virus as neutral red is a known mutagen that can alter your recombinant virus.

1. Prepare a 1 mg/ml Neutral Red solution in complete growth medium and filter sterilize.

2. Combine the following reagents in a 50 ml tube and place in a 37°C water bath.
   
   Neutral Red (1 mg/ml) 1.5 ml
   Complete growth medium 16.5 ml

3. Microwave 4% Agarose Gel until melted, then place in a 47°C water bath for 5 minutes.

4. Move the 50 ml tube of neutral red solution and the 4% Agarose Gel to a sterile hood. Add 6 ml of 4% Agarose Gel to the neutral red solution.

5. Add 1 ml of the Neutral Red overlay to each well containing plaquing overlay. Once the agarose has hardened, return plates to a 27°C incubator until plaques are ready to count. Plaques will appear as clear spots on a red monolayer.

Calculating Viral Titer

Use the equation below to calculate your viral titer.

\[
\text{pfu/ml} = \frac{\text{number of plaques (pfu)}}{\text{dilution factor} \times \text{ml of inoculum}}
\]

Example

A well with a viral dilution of \(10^{-8}\) contains 18 white plaques. The viral titer is:

\[
\text{pfu/ml} = \frac{18 \text{ pfu}}{10^{-8} \times 1 \text{ ml}} = 1.8 \times 10^9 \text{ pfu/ml}
\]

The Next Step

Once you have a viral stock of suitable titer (\(\geq 1 \times 10^8\) pfu/ml), you may infect cells and perform expression studies (see page 29). To plaque purify the virus or to analyze the recombinant DNA, proceed to the next section.
Isolating Virus From a Single Plaque

Introduction

This section provides detailed guidelines for plaque purifying your virus. Isolated virus can be used to generate a viral stock from a single viral clone or for PCR analysis of the recombinant baculovirus DNA. If you do not wish to plaque purify your virus, proceed to Expressing Recombinant Protein, page 29.

Materials Needed

- Sf21 cells (provided with the BaculoDirect® GST Gateway® Expression Kits; also available separately, see page vii) or Sf9 cells (available separately, see page vii)
- Complete growth medium
- Plates containing plaques from Step 5, page 22
- Six-well tissue culture plates
- 27°C incubator
- Inverted microscope
- Sterile Pasteur pipette and bulb
- Air-tight bags or containers

Preparing Cells

Use log phase Sf9 or Sf21 cells with greater than 95% viability.

1. Seed 8 × 10^5 Sf9 or Sf21 cells per well in 2 ml of complete growth medium in a six-well tissue culture plate. Gently tip the plate from side to side 4–6 times to evenly distribute the cells.

2. Incubate the cells at 27°C for one hour to allow the cells to fully attach to the bottom of the plate.

3. Verify that the cells have attached by inspecting them under an inverted microscope.

Infecting Cells

1. Using a sterile Pasteur pipette and bulb, carefully penetrate and remove the agarose containing the desired plaque.

2. Transfer the agarose plug containing the plaque to a 1.5 ml microcentrifuge tube containing 500 μl of complete growth medium. Mix well by vortexing.

3. Add 100 μl of the agarose plug solution from Step 2 to each well.

4. Place the plates in a sealed plastic bag with moist paper towels to prevent evaporation. Incubate at 27°C for 72 hours.

5. If you wish to isolate viral DNA, proceed to Isolating Viral DNA for PCR Analysis next page. If you wish to amplify your viral stock, proceed to Isolating Virus for Amplification, next page.

Continued on next page
**Isolating Virus From a Single Plaque, continued**

| **Isolating Viral DNA for PCR Analysis** | 1. Collect 2 ml of medium from each well from Step 4, previous page, and transfer to sterile 15 ml tubes. Centrifuge the tubes at 3,000–5,000 rpm for 5 minutes to remove cells and large debris.  
2. Transfer the supernatant to fresh 15 ml tubes.  
3. Proceed to **Analyzing Recombinant Viral DNA**, next page. |
|---|---|

| **Isolating Virus for Amplification** | 1. Incubate the cells for 2 more days. At 5 days post-infection, collect 2 ml of medium from each well and transfer to sterile 15 ml tubes. Centrifuge the tubes at 3,000–5,000 rpm for 5 minutes to remove cells and large debris.  
2. Transfer the supernatant to fresh 15 ml tubes. Store 1 ml of the viral clone stock at −80°C as a frozen stock and 1 ml at 4°C as a reserve stock. Refer to page 16 for additional storage information.  
3. Proceed to **Preparing a High-Titer Viral Stock and Screening for Expression**, page 17. |
|---|---|
Analyzing Recombinant Viral DNA

Introduction
You may analyze your recombinant viral DNA by PCR to verify the presence and orientation of your gene of interest. You may also use the PCR procedure on the next page to confirm your recombinant baculovirus DNA after the LR reaction. We recommend including a negative control (no DNA template) in your experiments to help you evaluate your results.

Viral DNA Purification
Invitrogen offers a variety of products that enable high-yield, high-purity DNA extraction from a wide range of sample types. For fast and easy isolation of DNA from baculoviruses, we recommend using the PureLink™ Genomic DNA Mini Kit or the Easy-DNA™ Kit (see page vi for ordering information). Follow the protocol provided with the kit manual for isolating baculovirus DNA. All Invitrogen manuals are available for downloading from our website (www.invitrogen.com) or by contacting Technical Support (page 45). An alternative protocol is also provided on the next page to isolate your baculovirus DNA.

Materials Needed
- Viral supernatant from Isolating Viral DNA for PCR Analysis, previous page
- 20% PEG 8000 in 1 M NaCl at 4°C (see page 41 for a recipe)
- Lysis buffer (0.1% Triton X-100 in PBS or TBS)
- Proteinase K (5–10 mg/ml, see page vi for ordering information)
- Phenol:chloroform:isoamyl alcohol (25:24:1)
- 3 M sodium acetate
- Glycogen (2 mg/ml, see page vi for ordering information)
- 100% ethanol
- 70% ethanol
- 50°C water bath

Continued on next page
Isolating Viral DNA

Perform the following protocol to lyse cells and extract the viral DNA.

1. Transfer 750 μl of your viral supernatant from Step 2 of Isolating Viral DNA for PCR Analysis, page 25, to a fresh 1.5 ml microcentrifuge tube.

2. Add 750 μl of cold (4°C) 20% PEG 8000 in 1 M NaCl. Invert the tube twice to mix and incubate at room temperature for 30 minutes.

3. Centrifuge at maximum speed for 10 minutes at room temperature to spin down the virus particles. Remove all medium from the pellet.

   Note: An additional quick spin may be required to remove trace amounts of medium. The pellet may not be visible at this point.

4. Add 100 μl of lysis buffer (0.1% Triton X-100 in PBS or TBS) to the pellet. Carefully wash the sides of the tubes to ensure that all of the viral particles are resuspended.

5. Add 10 μl of Proteinase K (5–10 mg/ml) and mix gently by inverting the tube. Incubate at 50°C for 1 hour.

6. Add 110 μl of phenol : chloroform : isoamyl alcohol (25:24:1) and mix gently by inverting the tube. Centrifuge at maximum speed for 5 minutes at room temperature. Transfer the upper aqueous phase to a fresh microcentrifuge tube.

7. Add the following reagents to the aqueous phase:
   - 3 M sodium acetate: 10 μl
   - Glycogen (2 mg/ml): 5 μl
   - 100% ethanol: 250 μl

   Incubate tubes at –20°C for at least 20 minutes.

8. Centrifuge at maximum speed for 15 minutes at 4°C. Wash the pellet with 70% ethanol. Centrifuge again and remove all traces of ethanol.

9. Resuspend the pellet in 10 μl of sterile water. Proceed to PCR Procedure, below.

PCR Procedure

You will need to optimize PCR conditions for your specific primers and template. For the PCR amplification, you may use the Polyhedrin forward primer (5’-AAATGATAACCATCTCGC-3’) and a primer of your own design that binds within your gene of interest. See page 28 for Polyhedrin forward primer binding site.

Continued on next page
Analyzing Recombinant Viral DNA, continued

**Analyzing PCR Results**

Calculate the expected size of your PCR fragment based on the location of the primer binding sites (see below for a diagram). After running your PCR reactions on a 1% agarose gel, you should see a band of the expected size for recombinant viral DNA and no bands for the negative control.

---

**Recombination Region for BaculoDirect™ N-GST Linear DNA**

The recombination region of the recombinant baculovirus resulting from BaculoDirect™ N-GST Linear DNA × entry clone is shown below.

**Features of the Recombination Region:**

- Shaded regions correspond to DNA sequence transferred from the entry clone into the BaculoDirect™ N-GST Linear DNA by recombination. Non-shaded regions are derived from the BaculoDirect™ N-Term Linear DNA.
- The underlined nucleotides flanking the shaded region correspond to bases 5272 and 10716, respectively, of the BaculoDirect™ N-GST Linear DNA sequence.
Expressing Recombinant Protein

Introduction
Once you have generated a viral stock of suitable titer (e.g., \(1 \times 10^8\) pfu/ml), you are ready to use the viral stock to infect insect cells and assay for expression of your recombinant protein. Guidelines for infection and expression are provided below.

Positive Control
If you generated a high-titer stock from the positive control construct pENTR™-CAT, we recommend infecting cells with this viral stock to help determine the optimal MOI for your particular cell line and application. Once you have infected cells with the positive control virus, the gene encoding chloramphenicol acetyltransferase (CAT) will be constitutively expressed and can be easily assayed (see page 32).

Guidelines for Expression
General guidelines are provided below to infect insect cells with the recombinant baculovirus to express your protein of interest.

- **Cell line:** Depending on your application and gene of interest, you may use any insect cell line (e.g., SF9, SF21, High Five™, Mimic™ SF9) for expression. Cells may be grown in adherent or suspension culture in the culture vessel of choice.
  
  **Note:** If you are expressing a secreted protein, you may improve expression by using High Five™ cells.

- **Culture Conditions:** We generally culture cells in serum-free conditions using Sf-900 II SFM, Sf-900™ III SFM, or Express Five® SFM as appropriate (see page vi). Depending on your application and the protein of interest, note that it may be necessary to supplement the culture post-infection with 0.1% to 0.5% FBS or BSA to protect the recombinant protein from proteolysis. Protein-based protease inhibitors are generally less expensive and more effective than many synthetic protease inhibitors.

- **Infection Conditions:** We recommend infecting cultures while cells are in the mid-logarithmic phase of growth at a density of \(1.5 \times 10^6\) to \(2.5 \times 10^6\) cells/ml. Make sure that the culture is not rate-limited by nutritional (i.e., amino acid or carbohydrate utilization) or environmental factors (i.e., pH, dissolved O₂, or temperature) during infection.

- **MOI:** Optimal MOI will vary between cell lines and the relative infection kinetics of the virus isolate or clone used. A dose response should be established for each virus, medium, reactor, and cell line employed to determine the optimal infection parameters to use for protein expression. As a starting point, infect cells using an MOI of 5 and 10. Refer to page 19 for an equation to determine how much virus stock to add to obtain a specific MOI.

- **Time course:** We recommend performing a time course to determine the expression kinetics for your recombinant protein as many proteins may be degraded by cellular proteases released in cell culture.
  
  **Note:** Maximum expression of secreted proteins is generally observed between 30–72 hours and non-secreted proteins between 48–96 hours post-infection.

- **Secreted proteins:** If you cloned your gene of interest with a secretion signal sequence, be aware that the N-terminal GST-tag may interfere with the secretion of your recombinant protein. We recommend that you verify the expression of protein using samples from both extracellular medium and cells lysates.

*Continued on next page*
Expressing Recombinant Protein, continued

**Optimizing Expression**

You may perform the following to determine the optimal conditions to use to express your recombinant protein of interest:

- **Cell line:** Infect Sf9, Sf21, High Five™, or Mimic™ Sf9 cells at a constant MOI. Assay for recombinant protein expression at different times post-infection (e.g., 24, 48, 72, 96 hours post-infection). Choose the cell line that provides the optimal level of recombinant protein expression.

- **MOI:** Infect a population of cells at varying MOIs (e.g., 1, 2, 5, 10, 20) and assay for protein expression. Use the MOI that provides the optimal level of recombinant protein expression.

- **Time course:** Infect cells at a constant MOI and assay for recombinant protein expression at different times post-infection (e.g., 24, 48, 72, 96 hours post-infection). Choose the time point at which optimal recombinant protein expression is obtained.

**Materials Needed**

- Insect cells of choice
- Complete growth medium
- Viral stock of known titer, \( \geq 1 \times 10^8 \text{ pfu/ml} \)
- SDS-PAGE Loading Buffer (NuPAGE® LDS Sample Buffer or Novex® Tris-Glycine SDS Sample Buffer, see page vi)
- Six-well tissue culture plate
- 27ºC incubator
- Inverted microscope

**Preparing Cells**

1. Seed \( 1 \times 10^6 \)–\( 2 \times 10^6 \) insect cells per well in 2 ml complete growth medium in a six-well tissue culture plate.
2. Incubate the cells at 27ºC for one hour to allow the cells to fully attach to the bottom of the plate.
3. Verify that the cells have attached by inspecting them under an inverted microscope.

*Continued on next page*
Expressing Recombinant Protein, continued

Preparing Protein Samples

Use the following procedure to prepare samples of your recombinant protein. This procedure is designed to allow expression analysis in a six-well format from cells harvested 24 to 96 hours post-infection. Other protocols are suitable.

1. Add the viral stock to each well at the desired MOI. Include the appropriate controls (e.g., mock-infected (uninfected) cells, positive control baculovirus, previously characterized recombinant baculoviruses).

2. Incubate infected cells at 27°C.

3. At the appropriate time (e.g., 24, 48, 72, 96 hours post-infection), harvest the cells and media and place in a 15 ml tube. Gently spin to pellet the cells. Transfer the cell medium to a fresh tube.

4. For analysis of intracellular protein, wash the cells 2X with PBS and resuspend the cells in 2 ml of PBS. Remove a 15 μl sample and add 5 μl of 4X SDS-PAGE Buffer.

   For analysis of secreted protein, remove a 15 μl sample of the cell medium and add 5 μl of 4X SDS-PAGE Buffer.

   Note: The N-terminal GST-tag may interfere with the secretion of your recombinant protein. We recommend that you verify the expression of protein using samples from both the extracellular medium and the cells lysates.

5. Freeze samples at –20°C or boil samples for at least 3 minutes and separate proteins by SDS-PAGE.
Analyzing Recombinant Protein

Introduction

You may analyze the expression of your recombinant protein by polyacrylamide gel electrophoresis or by Western blot analysis. General information for analyzing your recombinant GST-fusion protein is provided below. For detailed instructions, consult the literature provided with each product.

Polyacrylamide Gel Electrophoresis

To facilitate separation and visualization of your recombinant fusion protein by polyacrylamide gel electrophoresis, a wide range of pre-cast NuPAGE® and Novex® Tris-Glycine polyacrylamide gels and electrophoresis apparatus are available from Invitrogen. Invitrogen also carries a large selection of molecular weight protein standards and staining kits. For more information, refer to our website at www.invitrogen.com or contact Technical Support (page 45).

Detecting Recombinant Proteins

To detect expression of your recombinant fusion protein by western blot analysis, you may use antibodies against the N-terminal GST tag available from Invitrogen or an antibody to your protein of interest. The ready-to-use WesternBreeze® Chromogenic Kits and WesternBreeze® Chemiluminescent Kits are available from Invitrogen to facilitate detection of antibodies by colorimetric or chemiluminescent methods. See page viii for ordering information. For more information, refer to our website at www.invitrogen.com or contact Technical Support (page 45).

Assay for CAT Protein

If you used the control plasmid pENTR™-CAT to produce baculovirus expressing the CAT protein, you may assay for CAT expression using your method of choice. CAT will be fused to the N-terminal peptide containing the GST tag, allowing you to use western blot analysis with an anti-GST antibody. CAT Antiserum is also available separately from Invitrogen (see page vi for ordering information). Other commercial kits are available for assaying CAT expression. The molecular weight of the CAT fusion protein is approximately 53 kDa.

Due to the presence of the attB sites, there will be additional amino acids between your gene of interest and the N-terminal GST tag (see page 28 for a diagram). Expression of your protein with the GST tag will increase the size of your recombinant protein by approximately 27.8 kDa.

Note
Purifying Recombinant Protein

Introduction

Once you have optimized expression levels, you may purify your recombinant GST-fusion protein either by affinity chromatography on glutathione agarose or by immunoprecipitation using anti-GST antibodies. General information for purifying your recombinant GST-fusion protein is provided below. For detailed instructions, consult the literature provided with each product, which are also available through our website at www.invitrogen.com or by contacting Technical Support (page 45).

Glutathione Agarose

Invitrogen offers glutathione agarose for the purification of your recombinant GST-fusion protein by affinity chromatography in a single step (see page vi for ordering information). The glutathione agarose consists of glutathione linked via the sulfur atom to crosslinked beaded agarose and has a binding capacity of approximately 5-6 mg of bovine liver GST per ml of gel. Adding excess free glutathione liberates the GST fragment from the matrix, which can then be regenerated by washing with a high salt buffer. If you are using another resin, follow the manufacturer’s instructions.

Glutathione agarose is available from Invitrogen either as a sedimented bead suspension (10 ml or 100 ml) or as a part of the Glutathione Transferase Fusion Protein Purification Kit containing anti-GST antibodies and purification columns. For ordering information, see page viii.

Anti-Glutathione S-Transferase Antibody

Invitrogen also offers the highly purified rabbit polyclonal anti-GST antibody that can be used to purify GST fusion proteins by immunoprecipitation. This highly specific antibody, which was generated against a 260-amino acid N-terminal fragment of the Schistosoma japonica enzyme expressed in Escherichia coli, is also useful for detecting GST fusion proteins on western blots (see page 32). The intensely green-fluorescent Alexa Fluor® 488 conjugate of anti-glutathione S-transferase is also available for direct detection of GST fusion proteins. For ordering information, see page viii.

Removal of the GST tag

Your purified GST-fusion protein expressed from the recombinant baculovirus (BaculoDirect™ N-GST Linear DNA × entry clone) does not contain a cleavage site for the removal of the N-terminal GST tag. However, you can engineer your Gateway® entry clone to encode a recognition site for a specific protease, such as TEV, thrombin, or factor Xa, between the attL1 recombination site and your gene of interest. The LR reaction which facilitates the recombination of your entry clone (attL substrate) and the BaculoDirect™ N-GST Linear DNA (attR substrate) to create the recombinant expression baculovirus will leave the protease recognition site intact, allowing you to remove the GST tag with the appropriate protease.

For site-specific proteases available from Invitrogen, refer to www.invitrogen.com or contact Technical Support (page 45).
## Troubleshooting

### Transfection

The table below lists some potential problems and possible solutions to help you troubleshoot your transfection and initial protein expression screening experiments.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
</table>
| Transfected cells are dead, but Cellfectin® II-only control is fine | Contamination or cytotoxicity from the LR recombination reaction             | • Include a no-Cellfectin® II, entry clone-only negative control. \  
  • Use PureLink™ HiPure Plasmid Prep Kit, not a silica-based miniprep kit, for the purification of the entry clone. |
| Transfected cells do not show signs of infection | Kinetics of infection are slower than expected                               | • Virus production can take up to a week after transfection. Observe cells until 8 or 9 days after infection. If no signs of infection appear, investigate other possible causes. |
| Low transfection efficiency                  |                                                                               | • Use Cellfectin® II Reagent that is less than 6 months old and do not freeze Cellfectin® II Reagent for storage. \  
  • Perform transfection in Grace's Insect Medium Unsupplemented that does not contain supplements, antibiotics, or FBS. |
| Cells are not viable                         |                                                                               | • Cells should be in log phase and 95–98% viable. \  
  • Refer to the Insect Cell Lines manual for tips on culturing Sf9 and Sf21 cells. |
| Cells are not confluent enough               |                                                                               | Plate cells at 50–70% confluence.                                                                                                          |
| Cells are of high passage                    |                                                                               | Use cells that are between 8 to 15 passages.                                                                                              |
| Cells are of too low passage                 |                                                                               | After reviving cells, grow them for at least 5 passages before transfection.                                                             |
| Cells are too dense                          |                                                                               | Plate $8 \times 10^5$ cells per well for six-well plates. Split the cells if they are too confluent 3 days after transfection. Add complete growth medium containing $100 \mu M$ ganciclovir and incubate for 1 to 2 more days at 27°C. |
| LR recombination reaction unsuccessful      |                                                                               | • Make sure you added the LR Clonase™ II Enzyme Mix for BaculoDirect™ Kits to the LR reaction. \  
  • Check the pENTR™-CAT positive control transfection to verify that the LR reaction was successful. \  
  • Check your LR reaction using PCR as described on page 10. \  
  • Incubate LR reactions for up to 18 hours to increase recombinational efficiency. |
Troubleshooting, continued

**Transfection, continued**

The table below lists some potential problems and possible solutions to help you troubleshoot your transfection and initial protein expression screening experiments.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>No or little recombinant fusion protein detected in initial expression screen</td>
<td>Entry clone constructed incorrectly</td>
<td>Refer to page 8 of this manual and the instructions in the manual for the specific entry vector you are using for guidelines on constructing your entry clone.</td>
</tr>
</tbody>
</table>
|                                                                        | Insert not in frame with the N-terminal GST tag          | • Refer to the diagram on page 28 to verify the correct reading frame of the resulting recombinant baculovirus DNA following the LR reaction.  
                                                                             |                                                                        | • Analyze recombinant viral DNA by PCR to confirm correct size and orientation.  
                                                                             |                                                                        | • Sequence PCR product to verify proper reading frame for expression of the GST tag. |

**High-titer viral stock production**

The table below lists some potential problems and possible solutions to help you troubleshoot your high-titer P2 and P3 viral amplification experiments.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>MOI is lower than 0.01</td>
<td>Transfect 70% confluent cells on a 6-well plate with a dilution series of P1 viral stock, and monitor the cells every 24 hours. Use the MOI that does not produce any morphological changes within 24 hours in 70–80% of the infected cells.</td>
<td></td>
</tr>
</tbody>
</table>
| Cells are of high passage or passed their logarithmic phase of growth  |                                                                 | • Cells used for viral amplification should be younger than 20–25 passages.  
                                                                             |                                                                        | • If using suspension culture for amplification, cell density should be between 8 × 10⁵ and 1 × 10⁶ cells/ml. |
| Enlarged cell diameter observed 24 hours post-infection               | MOI is higher than 1                                     | High MOI will decrease the viral stock quality. Transfect 70% confluent cells on a six-well plate with a dilution series of P1 viral stock, and monitor the cells every 24 hours. Use the MOI that does not produce any morphological changes within 24 hours in 70–80% of the infected cells. |
**Troubleshooting, continued**

**Plaque Assay**

The following table lists some potential problems and possible solutions to help you troubleshoot your plaque assay.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>No plaques</td>
<td>Kinetics of infection are slower than expected</td>
<td>Observe plates until 8 or 9 days after infection. If no plaques appear, investigate other possible causes.</td>
</tr>
<tr>
<td></td>
<td>No confluent monolayer on Day 2 or Day 3 post-infection</td>
<td>Seed $8 \times 10^5$ cells in a six-well plate with 70% confluence. Cells should double at least once before infection stops growth.</td>
</tr>
<tr>
<td></td>
<td>Excessive condensation during incubation at 27ºC</td>
<td>Remove paper towels or open the container containing plates as soon as condensation appears.</td>
</tr>
<tr>
<td></td>
<td>Viral titer too low</td>
<td>Use higher concentrations of viral titer. You may need to re-infect your cells and collect a higher titer of your viral stock.</td>
</tr>
<tr>
<td>Small plaques that are difficult to visualize</td>
<td>Too many cells seeded</td>
<td>Seed fewer cells. We recommend seeding $8 \times 10^5$ cells per well for a six-well plate.</td>
</tr>
<tr>
<td></td>
<td>Viral titer not dilute enough</td>
<td>Prepare additional dilutions of your viral stock for infection.</td>
</tr>
<tr>
<td>Cells are dead</td>
<td>Temperature of the plaquing medium is too high</td>
<td>Prepare plaquing medium, then place in a 47ºC water bath until use.</td>
</tr>
<tr>
<td>Cracks in the agarose overlay</td>
<td>Growth medium not completely removed</td>
<td>Completely aspirate the growth medium before adding the plaquing medium. Any remaining growth medium can interfere with the gelling process.</td>
</tr>
</tbody>
</table>

*Continued on next page*
## Troubleshooting, continued

### Protein Expression

The following table lists some potential problems and possible solutions to help you troubleshoot your expression studies. We recommend including both positive and negative controls in your experiments to verify that correct reagents and protocols were used and to narrow down potential causes of the problem.

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very little or no recombinant fusion protein detected, but cells are infected and dead</td>
<td>Entry clone constructed incorrectly</td>
<td>Refer to page 8 of this manual and the instructions in the manual for the specific entry vector you are using for guidelines on constructing your entry clone.</td>
</tr>
<tr>
<td></td>
<td>Insert not in frame with N-terminal GST tag</td>
<td>• Refer to the diagram on page 28 to verify the correct reading frame of the resulting recombinant baculovirus DNA following the LR reaction.</td>
</tr>
<tr>
<td></td>
<td>Recombination error during LR reaction, or presence of a premature stop codon in construct</td>
<td>• Analyze recombinant viral DNA by PCR to confirm correct size and orientation (page 10).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Sequence PCR product to verify proper reading frame for expression of the GST tag.</td>
</tr>
<tr>
<td>Incorrect MOI used</td>
<td></td>
<td>• Run initial expression studies with an MOI of 5 and 10.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Recalculate the amount of viral stock needed to infect cells using the equation on page 19.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• You may need to test a range of MOIs depending on the kinetics of expression of your recombinant protein.</td>
</tr>
<tr>
<td>Cells harvested too late</td>
<td>Do a time course experiment and harvest cells at different time points (e.g., 48, 60, 72, and 96 hrs).</td>
<td></td>
</tr>
<tr>
<td>Protein is lost during cell lysis</td>
<td>If you are trying to detect an intracellular protein, analyze the supernatant to determine if the protein is being lost due to cell lysis.</td>
<td></td>
</tr>
<tr>
<td>Protein is degraded or unstable</td>
<td>Add protease inhibitors to your cell lysates.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Check mRNA levels.</td>
<td></td>
</tr>
<tr>
<td>Protein is toxic to cells</td>
<td>Harvest cells at earlier time points (e.g., 18-24 hours post-infection).</td>
<td></td>
</tr>
</tbody>
</table>

*Continued on next page*
## Troubleshooting, continued

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very little or no recombinant fusion protein detected, but cells are infected and dead</td>
<td>Viral stock a mixture of recombinant and non-recombinant virus</td>
<td>Add 100 μM ganciclovir at the end of transfection for P1 viral production. Use the same amount of ganciclovir during P2 viral amplification to select against non-recombinant virus.</td>
</tr>
<tr>
<td>Cell density too low</td>
<td></td>
<td>For protein expression using suspension culture, Sf9 and Sf21 cell density should be between $2.5 \times 10^6$ and $3.0 \times 10^6$ cells/ml.</td>
</tr>
<tr>
<td>High passage viral stock is used for protein expression</td>
<td></td>
<td>Cells used for protein expression should have less than 4 passages.</td>
</tr>
<tr>
<td>Protein is expressed, but escaped detection</td>
<td></td>
<td>• If expressing a secretion protein, make sure to check cells for the presence of the protein, because secretion will never be 100% efficient and sometimes could be very low.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• If your protein of interest is expressed intracellularly, make sure the check the cell lysate pellet for its presence.</td>
</tr>
<tr>
<td>Very little or no recombinant fusion protein detected, but cells are healthy and not dying after 72 hours</td>
<td>Viral stock is revived from frozen aliquots</td>
<td>The titer of the frozen viral stock will decrease after reviving. If the titer is too low, amplification may be needed</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Do a plaque assay or end-point dilution to accurately determine viral titer stock.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Do an MOI test with different MOI (e.g., 1, 5, and 10).</td>
</tr>
<tr>
<td>MOI is too low</td>
<td></td>
<td>For protein expression using suspension culture, Sf9 and Sf21 cell density should be between $2.5 \times 10^6$ and $3.0 \times 10^6$ cells/ml.</td>
</tr>
<tr>
<td>Cell density too high or cells are too old</td>
<td></td>
<td>Do a time course experiment and harvest cells at different time points (e.g., 48, 60, 72, and 96 hrs).</td>
</tr>
<tr>
<td>Protein is expressed but also see degradation</td>
<td>Harvesting time is not optimal</td>
<td>Do a time course experiment and harvest cells at different time points (e.g., 48, 60, 72, and 96 hrs).</td>
</tr>
<tr>
<td></td>
<td>MOI is too low</td>
<td>• Do a plaque assay or end-point dilution to accurately determine viral titer stock.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Do an MOI test with different MOI (e.g., 1, 5, and 10).</td>
</tr>
</tbody>
</table>

*Continued on next page*
## Troubleshooting, continued

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein is expressed but also see degradation</td>
<td>Premature stop codon in sequence</td>
<td>Check the sequence of your entry clone to verify that it does not contain a premature stop codon or mutations.</td>
</tr>
<tr>
<td>Protein is expressed, but is insoluble</td>
<td>A binding partner or other parts of protein may be needed for proper folding</td>
<td>Identify the partner and coexpress.</td>
</tr>
<tr>
<td>Protein expressed well small scale, but lost expression when scaled up</td>
<td>MOI and harvesting time may need to fine be tuned after scaling up</td>
<td>Using the MOI and harvesting time established during small scale test expression as a guideline, change harvesting time ±6 hours while keeping the MOI constant.</td>
</tr>
<tr>
<td>Viral stock worked well initially, but after a couple of months, expression levels decreased considerably</td>
<td>Viral stock was originally amplified using high MOI</td>
<td>Re-amplify virus from lower passage stock using low MOI (0.01–0.1).</td>
</tr>
<tr>
<td></td>
<td>Did not centrifuge and discard cells when harvesting viral supernatant</td>
<td>Re-amplify virus from lower passage stock using low MOI (0.01–0.1). If this viral stock is P2, it can be used in amplification.</td>
</tr>
<tr>
<td></td>
<td>For certain genes, the virus can become very unstable</td>
<td>Freeze the aliquoted viral stock and perform one round of amplification after reviving the virus.</td>
</tr>
</tbody>
</table>
Appendix

Recipes

Complete TNM-FH Medium

Complete TNM-FH medium is Grace’s Insect Medium with supplements (lactalbumin hydrolysate, L-glutamine, TC-yeastolate) and 10% fetal bovine serum (FBS).

1. If you are using Grace’s Insect Medium, Supplemented, add 55 ml of FBS. Mix well.
2. To include antibiotics and antifungals, add the following at the recommended concentration:
   - Penicillin 100U/ml
   - Streptomycin 100 μg/ml
   - Amphotericin 0.25 μg/ml
3. Filter-sterilize the solution through a 0.2 μm filter into a sterile container. A pre-filter may be required.
4. Store at 4°C and warm to 27°C before use.

Ganciclovir Stock Solution

Ganciclovir (100 mM in deionized water)

1. Add 26 mg of ganciclovir powder to 800 μl of deionized water.
2. Add 1 M NaOH dropwise until the solution reaches pH 12 and the ganciclovir dissolves into solution.
3. Add HCl dropwise until the solution reaches pH 11.
4. Bring up the final volume to 1 ml with deionized water.
5. Filter-sterilize the solution through a 0.2 μm filter.
6. Aliquot the solution into multiple tubes, and thaw each aliquot only once. Store at −20°C, protected from light, for up to 6 months. Thawed aliquots are stable at 4°C for up to 1 month.

Ganciclovir Working Solution

We recommend setting aside the amount of complete growth medium needed for the experiment requiring ganciclovir selection and adding the appropriate amount of ganciclovir to a final concentration of 100 μM. Aliquot the remaining ganciclovir into multiple tubes to reduce the number of freeze/thaw cycles.

Additional ganciclovir may be purchased in powder form from InvivoGen (Cat. no. sud-gcv).

Important

Ganciclovir provided with the BaculoDirect™ GST Transfection and Expression Kits has a concentration of 100 mM and might form a precipitate upon thawing. Incubating the ganciclovir at 37°C for 10 minutes and vortexing will redissolve the ganciclovir and eliminate the precipitate. If you experience ganciclovir precipitation, we recommend diluting the 100 mM stock solution in half with sterile distilled water before refreezing for storage.
Recipes, continued

**PEG/NaCl Solution**

20% Polyethylene glycol (PEG) 8000
1 M NaCl

1. Add the following reagents to 80 ml of deionized water:
   - PEG 8000 20 g
   - NaCl 5.84 g

2. Bring the final volume to 100 ml with deionized water.
3. Autoclave 20 minutes on liquid cycle.
4. While the solution is still warm (~55°C), swirl carefully to mix thoroughly.

**Bluo-gal**

Follow the guidelines below to prepare a 50 mg/ml stock solution of Bluo-gal.

1. Dissolve the Bluo-gal in dimethylformamide or dimethyl sulfoxide (DMSO) to make a 50 mg/ml stock solution. Use a glass or polypropylene tube.
   **Important:** Exercise caution when working with dimethylformamide. Dispense solutions in a vented chemical hood only.

2. Do not filter the stock solution.
**Map of BaculoDirect™ N-GST Linear DNA**

**Description**

BaculoDirect™ N-GST Linear DNA was constructed by homologous recombination between wild type *Autographa californica* multiple nuclear polyhedrosis virus (AcMNPV) DNA and a transfer plasmid containing a Gateway® cassette (see map below). After recombination, the Gateway® cassette replaces the native polyhedrin gene resulting in β-galactosidase positive, polyhedra negative recombinant virus. The modified baculovirus genome is linearized at the *Bsu*36 I site located at the 5’ end of the lacZ gene to produce BaculoDirect™ N-Term Linear DNA.

**Map**

The map below shows the Gateway® cassette elements of the BaculoDirect™ N-GST Linear DNA. The first nucleotide of the BaculoDirect™ N-GST Linear DNA sequence corresponds to the first *EcoR* I site in Homologous Region 1 (hr1). For the complete sequence of *Autographa californica* nuclear polyhedrosis virus, refer to GenBank Accession #NC_001623 or Ayres, M.D. *et. al.*, 1994.

---

**BaculoDirect™ N-GST Linear DNA Gateway® Cassette:**

- Polyhedrin promoter (*P*$_{PH}$): bases 4430-4556
- Polyhedrin Forward priming site: bases 4444-4461
- Initiation ATG: bases 4577-4579
- GST tag: bases 4580-5249
- *attR*1 recombination site: bases 5265-5362
- Herpes simplex virus thymidine kinase gene (*HSV1 tk*): bases 5649-6779 (c)
- Immediate early promoter [*P*$_{IE-1(0)}$]: bases 6808-7359 (c)
- *p10* promoter (*P*_p10): bases 7407-7504
- *Bsu*36 I linearization site: base 7755
- lacZ ORF: bases 7516-10590
- *attR2* recombination site: bases 10606-10730

*(c) = complementary strand*
## Features of the BaculoDirect™ N-GST Linear DNA

Features of the BaculoDirect™ N-GST Linear DNA Gateway® cassettes are described below. All features have been functionally tested.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Benefit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyhedrin promoter</td>
<td>Allows efficient, high-level expression of your recombinant protein.</td>
</tr>
<tr>
<td>Polyhedrin Forward priming site</td>
<td>Allows PCR detection and sequencing of the insert.</td>
</tr>
<tr>
<td>GST tag</td>
<td>Allows purification of the recombinant fusion protein by affinity chromatography on sulfur-linked glutathione agarose. Allows detection of the recombinant fusion protein with anti-GST antibodies. Helps solubilize the recombinant fusion protein</td>
</tr>
<tr>
<td>$attR1$ and $attR2$ sites</td>
<td>Allows recombination cloning of the gene of interest from an entry clone.</td>
</tr>
<tr>
<td>Immediate-early promoter (PIE-1(0))</td>
<td>Allows expression of the herpes simplex virus thymidine kinase gene (Kovacs et al., 1991).</td>
</tr>
<tr>
<td>Herpes simplex virus thymidine kinase gene (HSV1 tk)</td>
<td>Allows negative selection of non-recombinant virus in the presence of ganciclovir (Godeau et al., 1992).</td>
</tr>
<tr>
<td>p10 promoter</td>
<td>Allows expression of the $lacZ$ gene.</td>
</tr>
<tr>
<td>$lacZ$ gene</td>
<td>Allows detection non-recombinant virus through blue/white screening.</td>
</tr>
</tbody>
</table>
Map of pENTR™-CAT

Description

pENTR™-CAT (3231 bp) is a control vector containing the chloramphenicol acetyltransferase (CAT) gene. The CAT gene was amplified using PCR primers containing attB recombination sites. The amplified PCR product was then used in a BP recombination reaction with pDONR™221 to generate the entry clone. For more information about the BP recombination reaction, refer to the Gateway® Technology with Clonase™ II manual (part no. 25-0749).

Following an LR recombination reaction between pENTR™-CAT control vector and BaculoDirect™ Linear DNA, CAT will be expressed as a fusion to the N-terminal GST tag. The molecular weight of the CAT fusion protein is approximately 53 kDa.

Map

The map below shows the elements of the pENTR™-CAT control vector. The vector sequence of pENTR™-CAT is available at www.invitrogen.com or by contacting Technical Support (page 45).

Comments for pENTR™-CAT
3231 nucleotides

attL1 recombination site: bases 569-668
CAT ORF: bases 698-1354
attL2 recombination site: bases 1356-1455
Kanamycin resistance gene: bases 1625-2434
pUC origin: bases 2555-3228
Technical Support

Web Resources
Visit the Invitrogen website at www.invitrogen.com for:
• Technical resources, including manuals, vector maps and sequences, application notes, SDSs, FAQs, formulations, citations, handbooks, etc.
• Complete technical support contact information
• Access to the Invitrogen Online Catalog
• Additional product information and special offers

Contact Us
For more information or technical assistance, call, write, fax, or email. Additional international offices are listed on our website (www.invitrogen.com).

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SDS
Safety Data Sheets (SDSs) are available at www.invitrogen.com/sds.

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*Continued on next page*
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Gateway® Clone Distribution Policy

For additional information about Invitrogen’s policy for the use and distribution of Gateway® clones, see the section entitled Gateway® Clone Distribution Policy, next page.
Gateway® Clone Distribution Policy

**Introduction**

The information supplied in this section is intended to provide clarity concerning Invitrogen’s policy for the use and distribution of cloned nucleic acid fragments, including open reading frames, created using Invitrogen’s commercially available Gateway® Technology.

**Gateway® Entry Clones**

Invitrogen understands that Gateway® entry clones, containing attL1 and attL2 sites, may be generated by academic and government researchers for the purpose of scientific research. Invitrogen agrees that such clones may be distributed for scientific research by non-profit organizations and by for-profit organizations without royalty payment to Invitrogen.

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