pOG44

Flp-recombinase expression vector designed for use with the Flp-In $^{^{\text{\tiny{TM}}}}$ System

Catalog no. V6005-20

Version C 111110 25-0352



Table of Contents

Table of Contents	iii
Important Information	v
Methods	1
Overview	1
Using pOG44	3
Appendix	5
pOG44 Vector	5
Technical Service	
Purchaser Notification	9
References	11

Important Information

Contents

20 µg pOG44, lyophilized in TE, pH 8.0

Shipping/Storage

Lyophilized plasmid is shipped at room temperature and should be stored at -20°C.

Product Specifications

The pOG44 vector is qualified by restriction enzyme digestion with specific restriction enzymes as listed below. Restriction digests must demonstrate the correct banding pattern when electrophoresed on an agarose gel (see below).

Vector	Restriction Enzymes	Expected Results (bp)
pOG44	Kpn I	5438, 347
	Xba I	5785

Accessory Products

Many of the reagents used in the $\text{Flp-In}^{\text{TM}}$ System are available separately from Invitrogen. See the table below for ordering information.

Product	Amount	Catalog no.
pFRT/lacZeo	20 μg, lyophilized	V6015-20
pFRT/lacZeo2	20 μg, lyophilized	V6022-20
pcDNA5/FRT	20 μg, lyophilized	V6010-20
T7 Promoter Primer	2 μg, lyophilized	N560-02
Zeocin™	1 g	R250-01
	5 g	R250-05
Hygromycin	1 g	R220-05

Flp-In[™] Expression Vectors

Additional Flp-In^{$^{\text{TM}}$} expression vectors are available from Invitrogen. For more information about the features of each vector or to download a manual for a vector, refer to our World Wide Web site (www.invitrogen.com) or call Technical Service (see page 7).

Product	Amount	Catalog no.
pcDNA5/FRT/V5-His TOPO® TA Expression Kit	1 kit	K6020-01
pSecTag/FRT/V5-His TOPO® TA Expression Kit	1 kit	K6025-01
pEF5/FRT/V5 Directional TOPO® Expression Kit	1 kit	K6035-01
pEF5/FRT/V5-DEST Gateway [™] Vector Pack	6 µg	V6020-20

Important Information, continued

Flp-In[™] Host Cell Lines

For your convenience, Invitrogen has available several mammalian Flp-In[™] host cell lines that stably express the *lacZ-Zeocin*[™] fusion gene from pFRT/*lacZ*eo or pFRT/*lacZ*eo2. Each cell line contains a single integrated FRT site as confirmed by Southern blot analysis. The cell lines should be maintained in medium containing Zeocin[™]. For more information about the Flp-In[™] Cell Lines, see our World Wide Web site (www.invitrogen.com) or call Technical Service (see page 7).

Cell Line	Amount	Catalog no.
Flp-In [™] -293	3 x 10 ⁶ cells, frozen	R750-07
Flp-In [™] -CV-1	3 x 10 ⁶ cells, frozen	R752-07
Flp-In [™] -CHO	3 x 10 ⁶ cells, frozen	R758-07
Flp-In [™] -BHK	3 x 10 ⁶ cells, frozen	R760-07
Flp-In [™] -3T3	3 x 10 ⁶ cells, frozen	R761-07
Flp-In [™] -Jurkat	3 x 10 ⁶ cells, frozen	R762-07

Methods

Overview

Introduction

pOG44 is a 5.8 kb Flp recombinase expression vector designed for use with the Flp-In[™] System (Catalog nos. K6010-01 and K6010-02) available from Invitrogen. When cotransfected with the pcDNA5/FRT plasmid into a Flp-In[™] mammalian host cell line, the Flp recombinase expressed from pOG44 mediates integration of the pcDNA5/FRT vector containing the gene of interest into the genome via Flp Recombination Target (FRT) sites. The vector contains the following elements:

- The human cytomegalovirus (CMV) immediate-early enhancer/promoter for high-level constitutive expression of the Flp recombinase in a wide range of mammalian cells (Andersson *et al.*, 1989; Boshart *et al.*, 1985; Nelson *et al.*, 1987)
- Synthetic intron to enhance expression of the *FLP* gene (Huang and Gorman, 1990; O'Gorman *et al.*, 1991)
- *FLP* gene encoding the Flp recombinase (Buchholz *et al.*, 1996) to mediate integration of the pcDNA5/FRT expression plasmid into the genome

For more information about the Flp-InTM System, the pcDNA5/FRT plasmid, and generation of the Flp-InTM host cell line, refer to the Flp-InTM System manual. The Flp-InTM System manual is supplied with the Flp-InTM Complete or Core Systems, but is also available for downloading from our Web site (www.invitrogen.com) or by contacting Technical Service (see page 7).

FLP Gene

The *FLP* gene was originally isolated from the *Saccharomyces cerevisiae* 2µ plasmid (Broach *et al.*, 1982; Broach and Hicks, 1980), and encodes a site-specific recombinase that is a member of the integrase family of recombinases (Argos *et al.*, 1986). The Flp recombinase mediates a site-specific recombination reaction between interacting DNA molecules via the pairing of interacting FRT sites. For more information about site-specific recombination, refer to the next page and published reviews (Craig, 1988; Sauer, 1994).

The native *FLP* gene encodes a protein of 423 amino acids with a calculated molecular weight of 49 kDa. The *FLP* gene expressed from pOG44 encodes a temperature-sensitive Flp recombinase which carries a point mutation (flp-F70L) that results in a change in amino acid 70 from phenylalanine to leucine (Buchholz *et al.*, 1996). For more information about the properties of the flp-F70L protein, see below and Buchholz *et al.*, 1996.

Activity of the Flp Recombinase

When tested in mammalian cells, the native Flp recombinase has been shown to possess optimum recombination activity near 30°C and relatively low activity at 37°C, a result consistent with its physiological role in yeast (Buchholz *et al.*, 1996).

The flp-F70L protein expressed from pOG44 exhibits increased thermolability at 37°C in mammalian cells when compared to the native Flp recombinase (Buchholz *et al.*, 1996). Studies have shown that the Flp recombinase expressed from pOG44 possesses only **10%** of the activity of the native Flp recombinase at 37°C (Buchholz *et al.*, 1996).

Overview, continued

FIp Recombinase-Mediated DNA Recombination

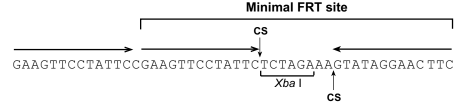
In the Flp-In[™] System, integration of the pcDNA5/FRT expression construct containing your gene of interest into the genome occurs via Flp recombinase-mediated intermolecular DNA recombination. The hallmarks of Flp-mediated recombination are listed below.

- Recombination occurs between specific FRT sites (see below) on the interacting DNA molecules
- Recombination is conservative and requires no DNA synthesis; the FRT sites are
 preserved following recombination and there is minimal opportunity for introduction
 of mutations at the recombination site
- Strand exchange requires only the small 34 bp minimal FRT site (see below)

For more information about the Flp recombinase and conservative site-specific recombination, refer to published reviews (Craig, 1988; Sauer, 1994).

FRT Site

The FRT site, originally isolated from *Saccharomyces cerevisiae*, serves as a binding site for Flp recombinase and has been well-characterized (Gronostajski and Sadowski, 1985; Jayaram, 1985; Sauer, 1994; Senecoff *et al.*, 1985). The minimal FRT site consists of a 34 bp sequence containing two 13 bp imperfect inverted repeats separated by an 8 bp spacer that includes an *Xba* I restriction site (see figure below). An additional 13 bp repeat is found in most FRT sites, but is not required for cleavage (Andrews *et al.*, 1985). While Flp recombinase binds to all three of the 13 bp repeats, strand cleavage actually occurs at the boundaries of the 8 bp spacer region (see figure below) (Andrews *et al.*, 1985; Senecoff *et al.*, 1985).



CS = cleavage site

In the Flp-InTM System, the pFRT/lacZeo and pcDNA5/FRT vectors each contain a single FRT site. The pFRT/lacZeo plasmid is used to generate the Flp-InTM host cell line and the pcDNA5/FRT plasmid is used to express the gene of interest in the Flp-InTM host cell line. For more information about pFRT/lacZeo, pcDNA5/FRT, and the Flp-InTM System, refer to the Flp-InTM System manual.

Generating Stable Expression Cell Lines

You will cotransfect the pOG44 plasmid and your pcDNA5/FRT construct into your Flp-In[™] host cell line(s) to generate stable cell lines that express your protein of interest. Cotransfection of pOG44 and pcDNA5/FRT allows expression of Flp recombinase resulting in integration of the pcDNA5/FRT plasmid into the genome via the FRT sites. Once the pcDNA5/FRT construct has integrated into the genome, the Flp recombinase is no longer required. The continued presence of Flp recombinase would actually be detrimental to the cells because it could mediate excision of the pcDNA5/FRT construct. For this reason, the pOG44 plasmid lacks an antibiotic resistance marker for selection in mammalian cells. When generating stable expression cell lines, the pOG44 plasmid and, therefore, Flp recombinase expression, will gradually be lost from transfected cells as they are cultured and selected.

Using pOG44

Introduction

General guidelines to transform pOG44 into E. coli are provided in this section.

General Molecular Biology Techniques

For help with *E. coli* transformation, restriction enzyme analysis, and DNA biochemistry, refer to *Molecular Cloning: A Laboratory Manual* (Sambrook *et al.*, 1989) or *Current Protocols in Molecular Biology* (Ausubel *et al.*, 1994).

E. coli Strain

Many *E. coli* strains are suitable for the propagation and maintenance of the pOG44 vector including TOP10, DH5α, and JM109. We recommend that you propagate the vector in *E. coli* strains that are recombination deficient (*rec*A) and endonuclease A deficient (*end*A).

For your convenience, TOP10 is available as chemically competent or electrocompetent cells from Invitrogen.

Item	Quantity	Catalog no.
One Shot® TOP10F' (chemically competent cells)	21 x 50 μl	C4040-03
One Shot [®] TOP10 Electrocomp [™] (electrocompetent cells)	21 x 50 µl	C4040-52
Electrocomp [™] TOP10 (electrocompetent cells)	5 x 80 μl	C664-55

Transformation Method

You may use any method of your choice for transformation. Chemical transformation is the most convenient method for many researchers. Electroporation is the most efficient and the method of choice for large plasmids.

Maintenance of Plasmid

To propagate and maintain the pOG44 vector, we recommend resuspending the vector in 20 μ l sterile water to prepare a 1 μ g/ μ l stock solution. Store the stock solution at -20°C.

Use this stock solution to transform a recA, endA E. coli strain like TOP10, DH5 α , JM109, or equivalent. Select transformants on LB agar plates containing 50 to 100 μ g/ml ampicillin. Be sure to prepare a glycerol stock of the plasmid for long-term storage (see below).

Preparing a Glycerol Stock

Once you have identified the correct clone, purify the colony and make a glycerol stock for long-term storage. You should keep a DNA stock of your plasmid at -20°C.

- Streak the original colony out on an LB plate containing 50 μg/ml ampicillin. Incubate the plate at 37°C overnight.
- Isolate a single colony and inoculate into 1-2 ml of LB containing 50 μg/ml ampicillin.
- Grow the culture to mid-log phase ($OD_{600} = 0.5 0.7$).
- Mix 0.85 ml of culture with 0.15 ml of sterile glycerol and transfer to a cryovial.
- Store at -80°C.

Using pOG44, continued

Plasmid Preparation

Plasmid DNA for transfection into eukaryotic cells must be very clean and free from phenol and sodium chloride. Contaminants will kill the cells, and salt will interfere with lipid complexing, decreasing transfection efficiency. We recommend isolating plasmid DNA using the S.N.A.P. ™ MiniPrep Kit (10-15 µg DNA, Catalog no. K1900-01), the S.N.A.P. ™ MidiPrep Kit (10-200 µg DNA, Catalog no. K1910-01), or CsCl gradient centrifugation.



Several Flp-In[™] host cell lines which stably express the *lacZ-Zeocin*[™] fusion gene and contain a single integrated FRT site are available from Invitrogen (see page vi for ordering information). If you wish to express your gene of interest in 293, CV-1, CHO, 3T3, BHK, or Jurkat cells, you may want to use one of the Flp-In[™] host cell lines to establish your expression cell line.



We have observed down-regulation of the viral CMV promoter and subsequent loss of gene expression when pcDNA5/FRT-based expression constructs are introduced into 3T3 or BHK cells. This behavior is not observed with pEF5/FRT-based expression constructs. If you are generationg Flp-In[™] expression cell lines using a 3T3 or BHK host cell line, we recommend that you clone your gene of interest into a pEF5/FRT-based expression plasmid (*e.g.* pEF5/FRT/V5-D-TOPO[®] or pEF5/FRT/V5-DEST). For more information, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 7).



Because correct integration of your pcDNA5/FRT construct into the genome is dependent upon Flp recombinase, the expression levels of Flp recombinase in the cell will determine the efficiency of the recombination reaction. Flp recombinase levels must be sufficiently high to mediate recombination at the FRT sites (single recombination event) and overcome the low intrinsic activity of the enzyme (see page 1). We have varied the ratio of pOG44 and pcDNA5/FRT expression plasmid that we cotransfect into mammalian Flp-In host cells to optimize the recombination efficiency. We recommend that you cotransfect you Flp-In host cell line with a ratio of at least 9:1 (w/w) pOG44:pcDNA5/FRT plasmid. Note that this ratio may vary depending on the nature of the cell line. You may want to determine this ratio empirically for your cell line.



When transfecting your Flp- In^{TM} host cell line, be sure to use **supercoiled** pOG44 and pcDNA5/FRT plasmid DNA. Flp-mediated recombination between the FRT site on pcDNA5/FRT and the integrated FRT site in the Flp- In^{TM} host cell line will only occur if the pcDNA5/FRT plasmid is circularized. The pOG44 plasmid should be circularized to minimize the possibility of the plasmid integrating into the genome.

Cotransfection

Once you have cloned your gene of interest into pcDNA5/FRT and have prepared clean plasmid preparations of pOG44 and your pcDNA5/FRT construct, cotransfect the plasmids into your mammalian Flp-InTM host cell line to generate your stable Flp-InTM expression cell line. We recommend that you include the appropriate positive and negative controls to help you evaluate your results. Specific guidelines and protocols for generation of the Flp-InTM expression cell line can be found in the Flp-InTM System manual.

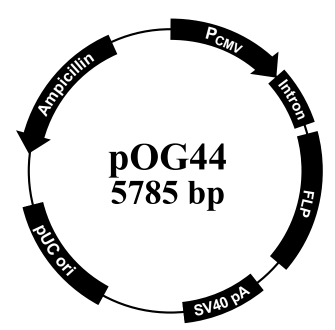
Reminder: The pcDNA5/FRT plasmid contains the hygromycin resistance gene to allow selection of transfectants using hygromycin. The pOG44 plasmid does not contain an antibiotic resistance gene for selection in mammalian cells (see pages 5-6).

Appendix

pOG44 Vector

Map of pOG44

pOG44 is a 5785 bp vector that expresses a temperature-sensitive Flp recombinase (flp-F70L) under the control of the human CMV promoter as previously described (O'Gorman *et al.*, 1991). The vector contains a synthetic intron to enhance expression of the *FLP* gene. Note that the vector does not contain an antibiotic resistance marker to allow stable selection in mammalian cells. The figure below summarizes the features of the pOG44 vector. The complete sequence for pOG44 is available for downloading from our World Wide Web site (www.invitrogen.com) or by contacting Technical Service (see page 7).



Comments for pOG44 5785 nucleotides

CMV promoter: bases 234-821 Synthetic intron: bases 871-1175 FLP ORF: bases 1202-2473

SV40 late polyadenylation signal: bases 2597-2732 pUC origin: bases 3327-3993 (complementary strand) *bla* promoter: bases 4999-5097 (complementary strand)

Ampicillin (bla) resistance gene: bases 4138-4998 (complementary strand)

pOG44 Vector, continued

Features of pOG44 The table below describes the relevant features of pOG44. All features have been functionally tested.

Feature	Benefit
Human cytomegalovirus (CMV) immediate early promoter	Allows high-level expression of the <i>FLP</i> gene (Andersson <i>et al.</i> , 1989; Boshart <i>et al.</i> , 1985; Nelson <i>et al.</i> , 1987)
Synthetic intron	Hybrid fragment which contains sequences derived from the adenovirus major late region and an IgG variable region (Huang and Gorman, 1990; O'Gorman <i>et al.</i> , 1991) and functions to enhance expression of the <i>FLP</i> gene
FLP ORF (flp-F70L)	Encodes a temperature-sensitive Flp recombinase (Buchholz <i>et al.</i> , 1996) that mediates conservative recombination via FRT sites (O'Gorman <i>et al.</i> , 1991)
SV40 late polyadenylation signal	Allows polyadenylation of mRNA
pUC origin	Allows high-copy number replication and growth in <i>E. coli</i>
bla promoter	Allows expression of the ampicillin (bla) resistance gene
Ampicillin (bla) resistance gene (β -lactamase)	Allows selection of transformants in <i>E. coli</i>

Technical Service

World Wide Web



Visit the Invitrogen Web Resource using your World Wide Web browser. At the site, you can:

- Get the scoop on our hot new products and special product offers
- View and download vector maps and sequences
- Download manuals in Adobe[®] Acrobat[®] (PDF) format
- Explore our catalog with full color graphics
- Obtain citations for Invitrogen products
- Request catalog and product literature

Once connected to the Internet, launch your Web browser (Internet Explorer 5.0 or newer or Netscape 4.0 or newer), then enter the following location (or URL):

http://www.invitrogen.com

...and the program will connect directly. Click on underlined text or outlined graphics to explore. Don't forget to put a bookmark at our site for easy reference!

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United States Headquarters:

Invitrogen Corporation 1600 Faraday Avenue Carlsbad, CA 92008 USA Tel: 1 760 603 7200

Tel (Toll Free): 1 800 955 6288

Fax: 1 760 602 6500

E-mail:

tech service@invitrogen.com

Japanese Headquarters

Invitrogen Japan K.K. Nihonbashi Hama-Cho Park Bldg. 4F 2-35-4, Hama-Cho, Nihonbashi

Tel: 81 3 3663 7972 Fax: 81 3 3663 8242

E-mail: jpinfo@invitrogen.com

European Headquarters:

Invitrogen Ltd 3 Fountain Drive Inchinnan Business Park Paisley PA4 9RF, UK

Tel (Free Phone Orders): 0800 269 210 Tel (General Enquiries): 0800 5345 5345

Fax: +44 (0) 141 814 6287

E-mail: eurotech@invitrogen.com

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- 1. On the home page, go to the left-hand column under 'Technical Resources' and select 'MSDS Requests'.
- 2. Follow instructions on the page and fill out all the required fields.
- 3. To request additional MSDSs, click the 'Add Another' button.
- 4. All requests will be faxed unless another method is selected.
- When you are finished entering information, click the 'Submit' button. Your MSDS will be sent within 24 hours.

Technical Service, continued

Emergency Information

In the event of an emergency, customers of Invitrogen can call the 3E Company, 24 hours a day, 7 days a week for disposal or spill information. The 3E Company can also connect the customer with poison control or with the University of California at San Diego Medical Center doctors.

3E Company

Voice: 1-760-602-8700

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- any use of the System or Expression Products to facilitate or advance any research or development program the results of which will be applied to the development of a Commercial Product.

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