Growth and Maintenance of Flp-In[™] Cell Lines

Catalog nos. R750-07, R752-07, R758-07, R760-07, R761-07, R762-07

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Important Information

Shipping/Storage	All cell lines are shi	pped on dry ice. Store	e in liquid nitrogen upon receipt.
Contents	This manual is supp	plied with the followi	ng cell lines:
	Cell Line	Catalog no.	
	Flp-In [™] -293	R750-07	
	Flp-In [™] -CV-1	R752-07	
	Flp-In [™] -CHO	R758-07	
	Flp-In [™] -BHK	R760-07	
	Flp-In [™] -3T3	R761-07	
	Flp-In [™] -Jurkat	R762-07	
Product	Freezing Medium.		ataining 3 x 10 ⁶ frozen cells in 1 ml of
Qualification	0	1 1	•
	 Each cell line is tested independently and certified to be free of mycoplasma. Prior to freezing, cells are greater than 95% viable. Forty-eight hours after thawing, cells are greater than 90% viable. Each cell line is tested for β-galactosidase activity by plating cells into 6-well plates in medium containing Zeocin[™]. After several days of growth, the cells are assayed for β-galactosidase activity using the β-Gal Staining Kit from Invitrogen. Each cell line must exhibit greater than 95% cells expressing 		
	β-galactosidase		

Accessory Products

IntroductionThe products listed in this section are intended for use with the Flp-In[™] Cell
Lines and the Flp-In[™] System. For more information, refer to our Web site
(www.invitrogen.com) or call Technical Service (see page 12).

Cell CultureA large variety of Gibco[™] cell culture products are available from Invitrogen to
facilitate growth and maintenance of the Flp-In[™] cell lines. For more information
about the products listed below, refer to our Web site (www.invitrogen.com) or
call Technical Service (see page 12). Note: Reagents are available in other sizes.

Item	Amount	Catalog no.
Dulbecco's Modified Eagle Medium (D-MEM)	500 ml	11965-092
Ham's F-12	500 ml	11765-054
RPMI Medium 1640	500 ml	11875-093
Fetal Bovine Serum	500 ml	16000-044
Donor Calf Serum	500 ml	16030-074
200 mM L-Glutamine	100 ml	25030-081
Penicillin-Streptomycin	100 ml	15070-063
Trypsin-EDTA	100 ml	25300-054

Additional Reagents

The products listed below may be used with the Flp-In[™] Cell Lines. Zeocin[™] is available for maintenance and growth of the Flp-In[™] cell lines. For more information, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 12).

Item	Amount	Catalog no.
Zeocin TM	1 g	R250-01
	5 g	R250-05
Hygromycin	1 g	R220-05
Lipofectamine [™] 2000 Reagent	0.75 ml	11668-027
	1.5 ml	11668-019
Lipofectamine [™] Reagent	1 ml	18324-012
Plus [™] Reagent	0.85 ml	11514-015
Phosphate-Buffered Saline (PBS) 7.4 (1X)	500 ml	10010-023

Flp-In[™] Products

The plasmids required to generate Flp-In[™] host cell lines and expression cell lines are available separately from Invitrogen. For more information about the features of each vector, refer to our Web site (www.invitrogen.com) or call Technical Service (see page 12). Ordering information is provided below.

Product	Amount	Catalog no.
pFRT/lacZeo	20 μg, lyophilized in TE	V6015-20
pFRT/lacZeo2	20 μg, lyophilized in TE	V6022-20
pOG44	20 μg, lyophilized in TE	V6005-20
pcDNA [™] 5/FRT	20 μg, lyophilized in TE	V6010-20
pcDNA [™] 5/FRT/V5-His TOPO [®] TA Expression Kit	20 reactions	K6020-01
pSecTag/FRT/V5-His TOPO® TA Expression Kit	20 reactions	K6025-01
pEF5/FRT/V5-DEST Gateway [®] Vector Pack	6 µg	V6020-20

Overview

designed for use wi Each cell line contai from pFRT/lacZeo of below and the next lines. For more infor the Flp-In [™] System is Technical Service (s downloading from of Generation of Flp-Ir cell line with a Flp-I Flp recombinase exp recombinase mediai genome at the integ (O'Gorman <i>et al.</i> , 19 interest from the Flp hygromycin B. For i	The Flp-In [™] cell lines stably express the <i>lacZ</i> -Zeocin [™] fusion gene and are designed for use with the Flp-In [™] System (Catalog nos. K6010-01 and K6010-02). Each cell line contains a single integrated Flp Recombination Target (FRT) site from pFRT/ <i>lac</i> Zeo or pFRT/ <i>lac</i> Zeo2 as confirmed by Southern blot analysis. See below and the next page for information about the generation of the Flp-In [™] cell lines. For more information about the Flp-In [™] System and its components, refer to the Flp-In [™] System manual, visit our Web site (www.invitrogen.com), or call Technical Service (see page 12). The Flp-In [™] System manual is also available for downloading from our Web site.
	Generation of Flp-In [™] expression cell lines requires cotransfection of the Flp-In [™] cell line with a Flp-In [™] expression vector containing your gene of interest and the Flp recombinase expression plasmid, pOG44 (O'Gorman <i>et al.</i> , 1991). Flp recombinase mediates insertion of your Flp-In [™] expression construct into the genome at the integrated FRT site through site-specific DNA recombination (O'Gorman <i>et al.</i> , 1991; Sauer, 1994). Stable cell lines expressing your gene of interest from the Flp-In [™] expression vector can be generated by selection using hygromycin B. For more information about FRT sites and Flp recombinase- mediated DNA recombination, refer to the Flp-In [™] System manual.
Parental Cell Lines	The table below provides a brief description of the source of the parental cell line used to generate each Flp-In [™] cell line. The parental cell lines were obtained from the American Type Culture Collection (ATCC). The ATCC number for each cell line is included. For further information about the parental cell lines, refer to the

Cell Line Characteristic Source ATCC Number Human embryonic kidney (Graham et al., 1977) 293 Adherent CRL-1573 CV-1 Adherent African Green Monkey kidney (Kit et al., 1965) CCL-70 CHO-K1 Adherent Chinese Hamster ovary (Kao and Puck, 1968) CCL-61 Adherent Baby hamster kidney (Talavera and Basilico, BHK CCL-10 1977) NIH/3T3 Adherent Mouse (NIH Swiss) embryonic fibroblast CRL-1658 (Jainchill *et al.*, 1969) Human T-cell leukemia (Weiss et al., 1984) TIB-152 Jurkat Suspension

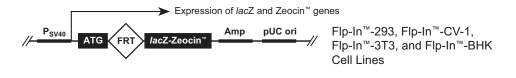
ATCC Web site (www.atcc.org).

Overview, continued

Description of Flp-In[™] Cell Lines

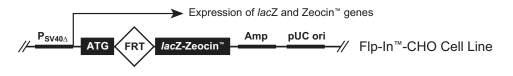
All of the Flp-In[™] cell lines (except Flp-In[™]-CHO; see below) contain a single integrated FRT site and stably express the *lacZ*-Zeocin[™] fusion gene from the pFRT/*lac*Zeo plasmid under the control of the SV40 early promoter (see diagram below). The location of the FRT site in each Flp-In[™] cell line has not been mapped, but is presumed to have integrated into a transcriptionally active genomic locus as determined by generation of a Flp-In[™] expression cell line containing the pcDNA[™]5/FRT/CAT or pEF5/FRT/GW-CAT control plasmid. The Flp-In[™] cell lines should be maintained in medium containing Zeocin[™] (see the next page).

For more information about pFRT/*lac*Zeo, refer to the Flp-In[™] System manual. For information about pcDNA[™]5/FRT/CAT or pEF5/FRT/GW-CAT, refer to the pcDNA[™]5/FRT or pEF5/FRT/V5-DEST manuals, respectively.



Flp-In[™]-CHO Cell Line

The Flp-InTM-CHO cell line contains a single integrated FRT site and stably expresses the *lacZ*-ZeocinTM fusion gene from the pFRT/*lac*Zeo2 plasmid. Note that pFRT/*lac*Zeo2 contains a mutated SV40 early promoter ($P_{SV40\Delta}$) which is severely abrogated in its activity. The SV40 Δ early promoter in pFRT/*lac*Zeo2 exhibits approximately 60-fold less activity than the wild-type SV40 early promoter in pFRT/*lac*Zeo. Because of the minimal activity of the SV40 Δ promoter, we expect that stable transfectants expressing the *lacZ*-ZeocinTM gene from pFRT/*lac*Zeo2 should contain FRT sites which have integrated into the most transcriptionally active genomic loci. The location of the FRT site in the Flp-InTM-CHO cell line has not been mapped, but has been demonstrated to have integrated into a highly transcriptionally active genomic locus as determined by generation of a Flp-InTM expression cell line containing the pcDNATM5/FRT/luc (luciferase-expressing) control plasmid. The Flp-InTM-CHO cell line should be maintained in medium containing ZeocinTM (see below). For more information about pFRT/*lac*Zeo2 and the SV40 Δ early promoter, refer to the pFRT/*lac*Zeo2 manual.



Overview, continued

Media for CellThe table below lists the recommended complete medium, freezing medium, and
antibiotic concentration required to maintain and culture each Flp-In[™] cell line.

Cell Line	Complete Medium	[Antibiotic]	Freezing Medium
Flp-In [™] -293	D-MEM (high glucose)	100 µg/ml Zeocin [™]	90% complete medium
	10% FBS*		10% DMSO
	2 mM L-glutamine		
	1% Pen-Strep (optional)		
Flp-In [™] -CV-1	D-MEM (high glucose)	100 µg/ml Zeocin [™]	90% complete medium
	10% FBS*		10% DMSO
	2 mM L-glutamine		
	1% Pen-Strep (optional)		
Flp-In [™] -CHO	Ham's F12	100 µg/ml Zeocin [™]	90% complete medium
	10% FBS*		10% DMSO
	2 mM L-glutamine		
	1% Pen-Strep (optional)		
Flp-In [™] -BHK	D-MEM (high glucose)	100 µg/ml Zeocin [™]	90% complete medium
	10% FBS*		10% DMSO
	2 mM L-glutamine		
	1% Pen-Strep (optional)		
Flp-In [™] -3T3	D-MEM (high glucose)	100 µg/ml Zeocin [™]	90% complete medium
	10% donor calf serum		10% DMSO
	2 mM L-glutamine		
	1% Pen-Strep (optional)		
Flp-In [™] -Jurkat	RPMI 1640	100 µg/ml Zeocin [™]	90% complete medium
	10% FBS*		10% DMSO
	2 mM L-glutamine		
	1% Pen-Strep (optional)		

*FBS = fetal bovine serum

Overview, continued

• FBS does not need to be heat inactivated for use with the	ese cell lines.
Guidelines • Cell lines should be maintained in medium containing 2 concentrations listed above.	Zeocin [™] at the
 If adherent cells (<i>e.g.</i> Flp-In[™]-293, Flp-In[™]-CV-1, Flp-In[™] Flp-In[™]-BHK) are split at a 1:5 to 1:10 dilution, they will 80-90% confluence in 3-4 days. 	
 Suspension Flp-In[™]-Jurkat cells will demonstrate optimal characteristics if maintained at a cell density between 1 1 x 10⁶ cells/ml. 	
 When maintaining Flp-In[™]-Jurkat cells in suspension cu the medium to turn yellow; this indicates that cells have density or that the medium is depleted of nutrients. If the fresh complete media to the cells or passage them. 	e reached too high a

Methods

Culturing Flp-In[™] Cell Lines

General Cell	Follow the guidelines below to successfully grow and maintain your cells.
Handling	• All solutions and equipment that come in contact with the cells must be sterile. Always use proper sterile technique and work in a laminar flow hood.
	• Before starting experiments, be sure to have cells established and also have some frozen stocks on hand. We recommend that you always use early- passage cells for your experiments. Upon receipt of the cells from Invitrogen, grow and freeze multiple vials of the particular cell line to ensure that you have any adequate supply of early-passage cells.
	• Cells should be at the appropriate confluence (approximately 60%) and >90% viability prior to transfection (see page 9).
	• For general maintenance of cells, pass all cell lines when they are $80-90\%$ confluent (for adherent cells) or when they reach a density of 2×10^6 to 4×10^6 cells/ml (for suspension cells).
	 Use trypan blue exclusion to determine cell viability. Log phase cultures should be >90% viable.
Before Starting	Be sure to have the following solutions and supplies available:
	• 15 ml sterile, conical tubes
	• 5, 10, and 25 ml sterile pipettes
	Cryovials
	Phosphate-Buffered Saline (see page v for ordering information)
	• 0.4% Trypan blue in PBS and hemacytometer (for counting cells)
	• Reagents to prepare the appropriate complete medium (see page 3)
	• Freezing Medium (see pages 3 and 8)
	Table-top centrifuge
	• 75 cm ² flasks, 175 cm ² flasks and other appropriately-sized tissue culture flasks or plates
	• Trypsin/versene (EDTA) solution or other trypsin solution

Culturing Flp-In[™] Cell Lines, continued

Thawing Adherent Cells	cul	e following protocol is designed to help you thaw adherent cells to initiate cell ture. All cell lines are supplied in vials containing 3×10^6 cells in 1 ml of pezing Medium.
	1.	Remove the vial of cells from the liquid nitrogen and thaw quickly in a 37°C water bath.
	2.	Just before the cells are completely thawed, decontaminate the outside of the vial with 70% ethanol, and transfer the cells to a T-75 flask containing 12 ml of complete medium without Zeocin [™] .
	3.	Incubate the flask at 37°C for 2-4 hours to allow the cells to attach to the bottom of the flask.
	4.	Aspirate off the medium and replace with 12 ml of fresh, complete medium without Zeocin [™] .
	5.	Incubate cells overnight at 37°C.
	6.	The next day, aspirate off the medium and replace with fresh, complete medium containing Zeocin [™] (at the recommended concentration listed on page 3).
	7.	Incubate the cells and check them daily until the cells are 80-90% confluent (2-7 days).
	8.	Proceed to Passaging Adherent Cells, next page.
Thawing Suspension Cells	cel	e following protocol is designed to help you thaw suspension cells to initiate l culture. All cell lines are supplied in vials containing 3×10^6 cells in 1 ml of eezing Medium.
	1.	Remove the vial of cells from the liquid nitrogen and thaw quickly in a 37°C water bath.
	2.	Just before the cells are completely thawed, decontaminate the outside of the vial with 70% ethanol, and transfer the cells to a sterile, conical tube containing 5 ml of complete medium without Zeocin [™] .
	3.	Centrifuge for 3 minutes at 750 x g at room temperature.
	4.	Aspirate off the medium and resuspend the cells in 12 ml of fresh, complete medium without $ ext{Zeocin}^{\mathbb{M}}$.
	5.	Transfer the cells to a T-75 flask and incubate cells overnight at 37°C.
	6.	The next day, add Zeocin ^{TM} to the cells (at the recommended concentration listed on page 3).
	7.	Incubate the cells and count them daily until the cells reach a density ranging from 2×10^6 cells/ml to 4×10^6 cells/ml (2-7 days). Note: You may add fresh, complete medium containing Zeocin TM to the cells every few days.
	8.	Proceed to Passaging Suspension Cells, next page.
		continued on next nage

Culturing Flp-In[™] Cell Lines, continued

Passaging	1.	When cells are ~80-90% confluent, remove all medium from the flask.
Adherent Cells	2.	Wash cells once with 10 ml PBS to remove excess medium and serum. Serum contains inhibitors of trypsin.
	3.	Add 5 ml of trypsin/versene (EDTA) solution to the monolayer and incubate 1 to 5 minutes at room temperature until cells detach. Check the cells under a microscope and confirm that most of the cells have detached. If cells are still attached, incubate a little longer until most of the cells have detached.
	4.	Add 5 ml of complete medium to stop trypsinization.
	5.	Briefly pipet the solution up and down to break up clumps of cells.
	6.	To maintain cells in 75 cm ² flasks, transfer 1 ml of the 10 ml cell suspension from Step 5 to a new 75 cm ² flask and add 15 ml fresh, complete containing Zeocin ^{M} . If you want the cells to reach confluency sooner, split the cells at a lower dilution (<i>e.g.</i> 1:4).
		Note: To expand cells into 175 cm ² flasks, add 28 ml of fresh, complete medium containing Zeocin [™] to each of three 175 cm ² flasks, then transfer 2 ml of the cell suspension to each flask to obtain a total volume of 30 ml.
	7.	Incubate flasks in a humidified, 37° C, 5% CO ₂ incubator.
	8.	Repeat Steps 1-7 as necessary to maintain or expand cells.
Passaging	1.	Passage suspension cells when they reach a density of 2-4 x 10^6 cells/ml.
Suspension Cells	2.	To maintain cells in 75 cm ² flasks, transfer 1-1.5 ml of cell suspension from Step 1 to a new 75 cm ² flask containing 13-14 ml of fresh, complete medium with Zeocin ^{TM} .
		Note: You may split the cells at a lower dilution (e.g. 1:4), if desired.
	3.	To expand cells into 175 cm² flasks, add 28 ml of fresh, complete medium containing Zeocin [™] to each of three 175 cm² flasks, then transfer 2 ml of the cell suspension to each flask to obtain a total volume of 30 ml.
		Note: You may also expand cells into a spinner flask, if desired.
	4.	Incubate flasks in a humidified, 37° C, CO ₂ incubator.
	Re	peat Steps 1-4 as necessary to maintain or expand cells.

Freezing Cells

Introduction	 When freezing the Flp-In[™] cell lines, we recommend the following: Freeze cells at a density of at least 3 x 10⁶ cells/ml. Use a freezing medium composed of 90% complete medium and 10% DMSO. Complete medium is medium containing serum. Guidelines to prepare freezing medium and freeze cells are provided in this section.
Preparing Freezing Medium	 Freezing medium should be prepared fresh immediately before use. 1. In a sterile, conical centrifuge tube, mix together the following reagents for every 1 ml of freezing medium needed: Fresh complete medium 0.9 ml DMSO 0.1 ml Place the tube on ice. Discard any remaining freezing medium after use.
Freezing the Cells	 Before starting, label cryovials and prepare freezing medium (see above). Keep the freezing medium on ice. 1. To collect cells, perform the following: For adherent cells, follow Steps 1-5 of Passaging Adherent Cells, page 7. For suspension cells, transfer cells to a sterile, conical centrifuge tube. 2. Count the cells. Pellet cells at 250 x g for 5 minutes in a table top centrifuge at room temperature and carefully aspirate off the medium. 4. Resuspend the cells at a density of at least 3 x10⁶ cells/ml in chilled freezing medium. 5. Place vials in a microcentrifuge rack and aliquot 1 ml of the cell suspension into each cryovial. 6. Freeze cells in an automated or manual, controlled-rate freezing apparatus following standard procedures. For ideal cryopreservation, the freezing rate should be a decrease of 1°C per minute. 7. Transfer vials to liquid nitrogen for long-term storage. Note: You may check the viability and recovery of frozen cells 24 hours after storing cryovials in liquid nitrogen by following the procedure outlined in Thawing Adherent Cells or Thawing Suspension Cells, page 6, as appropriate.

Transfecting Cells

Introduction	To generate stable Flp-In [™] expression cell lines, you will cotransfect your Flp-In [™] expression construct and the pOG44 plasmid into the Flp-In [™] cell line and select for stable transfectants using hygromycin. General guidelines and recommendations for transfection are provided in this section. We recommend that you read through this section before beginning.
Transfection Methods	The Flp-In [™] cell lines are generally amenable to transfection using standard methods including calcium phosphate precipitation (Chen and Okayama, 1987; Wigler <i>et al.</i> , 1977), lipid-mediated transfection (Felgner <i>et al.</i> , 1989; Felgner and Ringold, 1989), and electroporation (Chu <i>et al.</i> , 1987; Shigekawa and Dower, 1988). We typically use lipid-based transfection reagents to introduce Flp-In [™] expression constructs into the Flp-In cell lines. The table below lists the recommended transfection reagent for each Flp-In [™] cell line. Other transfection reagents may be suitable.

To transfect	Use
Flp-In [™] -293	Lipofectamine [™] 2000 Reagent
Flp-In [™] -CV-1	
Flp-In [™] -CHO	
Flp-In [™] -Jurkat	
Flp-In [™] -3T3	Lipofectamine [™] Reagent and
Flp-In [™] -BHK	Plus [™] Reagent

Note: LipofectamineTM 2000 Reagent, LipofectamineTM Reagent, and PlusTM Reagent are available from Invitrogen (see page v for ordering information).



We have observed down-regulation of the viral CMV promoter and subsequent loss of gene expression when pcDNA[™]5/FRT-based expression constructs are introduced into Flp-In[™]-3T3 or Flp-In[™]-BHK cells. This behavior is not observed with pEF5/FRT-based expression constructs. If you are generating Flp-In[™] expression cell lines using the Flp-In[™]-3T3 or Flp-In[™]-BHK cell line, we recommend that you express your gene of interest from a pEF5/FRT-based plasmid (*e.g.* pEF5/FRT/V5-DEST).

Transfecting Cells, continued

Generating Stable Expression Cell Lines

To generate Flp-In[™] expression cell lines, cotransfect your Flp-In[™] expression construct and the pOG44 plasmid into the Flp-In[™] cell line of choice, and select for stable transfectants using hygromycin B. Before transfection, you may want to test the sensitivity of the Flp-In[™] cell line to hygromycin B to more accurately determine the hygromycin B concentration to use for selection. A suggested range of hygromycin B concentrations to use for selection of your Flp-In[™] expression vector is listed below. For more information, refer to the Flp-In[™] System manual. Hygromycin B may be obtained from Invitrogen (see page v for ordering information).

Important: Following cotransfection, your Flp-In[™] expression clones should become sensitive to Zeocin[™]; therefore, your selection medium should NOT contain Zeocin[™].

Cell Line (After Transfection with Flp-In [™] Expression Vector)	Estimated Hygromycin B Concentration (µg/ml)
Flp-In [™] -293	100-200
Flp-In [™] -CV-1	100-200
Flp-In [™] -CHO	500-600
Flp-In [™] -BHK	100-200
Flp-In [™] -3T3	100-200
Flp-In [™] -Jurkat	200-400

When transfecting Flp-InTM-CHO cells, we recommend following these guidelines:

- 48 hours after transfection, split the cells directly into medium containing the appropriate concentration of hygromycin.
- Split the cells such that they are no more than 25% confluent. If the cells are too dense, the antibiotic will not kill the cells. Antibiotics work best on actively dividing cells.

Polyclonal Selection of Isogenic Cell Lines

Note

Because every Flp-In[™] cell line contains a single integrated FRT site, all of the hygromycin-resistant foci that you obtain after cotransfection with the Flp-In[™] expression vector and pOG44 should be isogenic (*i.e.* the Flp-In[™] expression vector should integrate into the same genomic locus in every clone; therefore, all clones should be identical). To obtain stable expression cell lines, you may perform "polyclonal" selection and screening of your hygromycin-resistant cells. After hygromycin selection, simply pool the hygromycin-resistant foci and screen the entire population of cells for the following phenotypes:

- Zeocin[™] sensitivity
- Lack of β-galactosidase activity
- Expression of the gene of interest

Transfecting Cells, continued

Selection of
Individual Cell
LinesIf desired, single hygromycin-resistant foci can be isolated and expanded to
generate individual clonal cell lines. To isolate individual clones, simply pick
5-20 hygromycin-resistant foci and expand the cells. You may verify that your
Flp-In[™] expression construct has integrated into the FRT site by testing each
clone for Zeocin[™] sensitivity and lack of β-galactosidase activity. Select those
clones that are hygromycin-resistant, Zeocin[™]-sensitive, and lack β-galactosidase
activity, and assay for expression of your gene of interest.NoteNote that in rare instances, it is possible to generate a Flp-In[™] expression cell
line in which the Flp-In[™] expression plasmid has undergone both Flp
recombinase-mediated integration into the FRT site and random integration
into a second genomic site. In this case, clones will still exhibit hygromycin

Note that in rare instances, it is possible to generate a FIp-InTM expression cell line in which the FIp-InTM expression plasmid has undergone both FIp recombinase-mediated integration into the FRT site and random integration into a second genomic site. In this case, clones will still exhibit hygromycin resistance. To test for these "second site integrants", transfect the cells with the pOG44 plasmid and select for ZeocinTM resistance. The Flp recombinase should mediate excision of the Flp-InTM expression plasmid at the FRT site and restore the *lacZ*-ZeocinTM fusion gene. The resulting cells should exhibit β -galactosidase activity, ZeocinTM resistance, and continued expression of the gene of interest. Alternatively, you may perform Southern blot analysis to identify second site integrants if suitable restriction enzymes are selected.

Appendix

Technical Service

World Wide Web	 site, you can: Get the scoop on our hot new processory View and download vector maps Download manuals in Adobe[®] Accossory Explore our catalog with full colors Obtain citations for Invitrogen processory Request catalog and product literations Once connected to the Internet, launch or newer or Netscape 4.0 or newer), the www.in and the program will connect direct 	and sequences robat [®] (PDF) format graphics oducts
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To request an MSDS, visit our Web site at www.invitrogen.com. On the home page, go to 'Technical Resources', select 'MSDS', and follow instructions on the page.

Technical Service, continued

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Purchaser Notification

Introduction	Use of the Flp-In [™] Cell Lines and the Flp-In [™] System ("System") is covered under a number of different licenses including those detailed below.
Information for European Customers	The Flp-In [™] cell lines are genetically modified and carry either the pUC-derived plasmid, pFRT/ <i>lacZ</i> eo or pFRT/ <i>lacZ</i> eo2. As a condition of sale, this product must be in accordance with all applicable local legislation and guidelines including EC Directive 90/219/EEC on the contained use of genetically modified organisms.
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