Cell-Specific Ecdysone-Inducible Expression of FLP Recombinase in Mammalian Cells


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INTRODUCTION

The ability to manipulate genomes of higher organisms using site-directed recombinases such as FLP, from *Saccharomyces cerevisiae*, and Cre, from bacteriophage P1, is widely recognized as an important way to explore the molecular basis for developmental processes. The simplicity by which these recombinases work to catalyze excisions, integrations, inversions or translocations at specific target sites in DNA also suggests that with the development of strategies to regulate the recombination events they mediate, these enzymes will be important tools for introducing specific genetic alterations required by gene therapy.

Several studies have shown that recombinase-mediated DNA alterations can be regulated either (i) by controlling the expression of the recombinase with inducible promoters (3,5–7,11), (ii) by introducing recombinase mRNA or protein into cells (1,8) or (iii) by regulating the activity of an expressed recombinase (9). When applied to whole animals, however, each of these approaches is either impractical or has the disadvantage that cell-specific regulation cannot be attained.

Recently, the insect steroid hormone ecdysone has been shown to be an efficient and potent inducer of gene expression in cultured cells and transgenic mice (12). In this report, we place the FLP coding sequence under the regulation of an ecdysone-responsive promoter, we temporally induce FLP-mediated, site-specific recombination events in specific cells and their descendants.

MATERIALS AND METHODS

Plasmid Constructs

pNEOβGAL, pFRTβGAL and pOG44 were purchased from Stratagene (La Jolla, CA, USA). The pVgRXR and pIND/lacZ plasmids are components of the Ecdysone-Inducible Mammalian Expression System (Invitrogen, Carlsbad, CA, USA). More information on these plasmids is available on the Invitrogen Web site (http://www.invitrogen.com). The pIND/FLP plasmid, which contains the ecdysone-responsive minimal heat shock promoter (IND), was constructed by ligating the 2030-bp XbaI/ApaI fragment containing the FLP sequence of pOG44 into the XbaI and ApaI sites in the multiple cloning site of pIND. pK19/VgRXR was constructed in two steps. First, three fragments were ligated into a HindIII/PstI-digested pUC19 vector sequence as follows: (i) a PstI/ApaI fragment containing the VgEcR and TKpA sequences of pVgRXR, (ii) a RXR-BGHpA sequence amplified from pVgRXR using two oligonucleotides; 5′-CCATCGATATATCCACATGGACAC-3′ and 5′-TCGTAAAGCTTCCGTCATTATACGACACAGA-3′ (underlined nucleotides correspond to Clal and HindIII sites) and (iii) a murine K19 promoter sequence (10) amplified from...
a plasmid pK19βgal using two oligonucleotides; 5'-AGTGTGTCAGTGCCAGTAGCTTT-3' and 5'-CCATCTGATGATGAGGAGGAGGAGACC-3' (underlined nucleotides correspond to ApaLI and ClaI sites). The resulting 8070-bp plasmid was digested with PstI and NdeI. To introduce a second K19 promoter sequence into the construct 5' to the VgEcR sequence, the resulting 7815-bp fragment was ligated to another fragment containing the same K19 promoter sequence as above, this time amplified from pK19βgal using two different oligonucleotides, 5'-AGTCTCATATGTCAGTGCCAGTAGCTTT-3' and 5'-TCGGCTGACAGTATGAGGAGGAGGAGACC-3' (underlined nucleotides correspond to NdeI and PstI sites).

Cell Culture and Transfection

NIH 3T3 cells (7.5 × 10^6) (Accession No. CRL-1658; ATCC, Rockville, MD, USA) were electroporated in phosphate-buffered saline (PBS), pH 7.3, using a Gene Pulser (960 µF, 0.3 kV, 0.4-cm electrode) (Bio-Rad, Hercules, CA, USA). Ten micrograms of each plasmid DNA were used. Electroporated cells were seeded into 15-mm tissue culture dishes. Twenty-four hours after electroporation, the culture medium was replaced with fresh culture medium or medium containing muristerone A for 24 h. Following fixation with acetone for 10 min at -20°C, cells were incubated overnight at 4°C in a humidified chamber with a monoclonal antibody to K19 (1:10) (Catalog No. RPN1165; Amersham Pharmacia Biotech, Piscataway, NJ, USA) and a rabbit polyclonal antibody to β-gal (1:100) (Catalog No. 7-063100; 5 Prime → 3 Prime, Boulder, CO, USA). The secondary antibodies used were a CY3-conjugated goat anti-mouse IgG and a FITC-conjugated goat anti-rabbit IgG (Jackson ImmunoResearch, West Grove, PA, USA).

RESULTS

Figure 1 shows the DNA constructs used in this study. In the plasmid pNEOβGAL, the β-gal coding sequence has been disrupted by a neo gene that is flanked by FLP recognition target (FRT) sites in direct orientation. The FRT-flanked neo cassette is inserted after the first codon (AUG) of the β-gal coding sequence; therefore, not even a truncated β-gal protein is made from the disrupted β-gal sequence. FLP-
mediated recombination between these FRTs results in the excision of neo, thereby reconstituting an intact and functional β-gal sequence. The plasmid pFRTβGAL contains an intact β-gal coding sequence preceded by a single FRT. Elements required for ec dysone induction are housed in the plasmids pVgRXR, pIND/FLP and pIND/lacZ. pVgRXR contains genes encoding a heterodimer of the ec dysone receptor (VgEcR; derived from Drosophila and modified to contain the VP16 transactivation domain) and the retinoid-X receptor (RXR; derived from mammalian cells). This heterodimer binds to a modified ec dysone response element (E/GRE) in the presence of ec dysone (or a natural analog, muristerone A). In pIND/FLP, the FLP recombinase sequence has been placed under the control of the inducible promoter sequence, IND, containing the modified E/GRE linked to a minimal heat shock promoter. The same promoter controls activation of the β-gal encoding lacZ gene in pND/lacZ.

Figure 2 outlines the method for combining the genetic elements required for ec dysone-induction with those required for FLP-mediated recombination, which can introduce site-
specific changes in DNA in an inducible manner. When cells that express the ecdysone receptor-RXR heterodimer are treated with muristerone, FLP recombinase under the control of the IND promoter will be expressed. Site-directed, FLP-mediated recombination events will then occur between DNA sequences containing FRTs. Figure 2 shows a FLP-mediated recombination event that results in the excision of a neo sequence, thereby reconstituting an intact lacZ gene.

To test whether FLP-mediated recombination can actually be induced by ecdysone, NIH-3T3 cells were transiently co-transfected with the plasmids pVgRXR, pIND/FLP and pneoβgal. Twenty-four hours after electroporation, muristerone A was added to the culture medium at various concentrations. After another 24 h, cells were fixed and histochemically stained for β-gal activity (cells having β-gal activity stain blue; Figure 3A).

Several control transfections (C1–C5) were also done. Table 1 summarizes the number of blue cells in each transfection, with and without muristerone A induction. For each transfection, in addition to the plasmids indicated in Table I, cells were co-transfected with a plasmid, pCX/EGFP, encoding a variant green fluorescent protein (GFP). The number of blue cells was normalized to the number of cells expressing GFP (Figure 3B). Our ability to transfact cells at a reasonable rate is demonstrated by the high number of cells that stain blue in C1, i.e., cells transfected with pFRTβGAL. No blue cells were observed in cells transfected with pIND/lacZ (C2), either in the absence or presence of muristerone A, indicating that IND is not leaky and is not activated in the absence of the VgEcR-RXR heterodimer. In cells transfected with pVgRXR and pIND/lacZ (C3), the presence of a small number of blue cells in the absence of muristerone indicates that, in the presence of the VgEcR-RXR heterodimer, the IND promoter is slightly leaky. Activation of this promoter is shown to be responsive to muristerone in a dose-dependent manner, as the number of blue cells increased with increased concentrations of muristerone (also see Figure 3C). No blue cells were observed when cells were transfected with pNEOβGAL (C4) indicating that there is little, if any, spontaneous excision of the FRT-flanked neo sequence. The absence of blue cells when cells were electroporated with PBS lacking DNA (C5) confirmed the specificity of the histochemical stain for lacZ-encoded β-gal activity.

When cells were transfected with pVgRXR, pIND/FLP and pNEOβGAL, very few cells stained blue in the absence of muristerone. The number of blue cells increased significantly in a dose-dependent manner when cells were treated with muristerone (Figure 3C). In the absence or presence of muristerone, the number of blue cells is lower than in C3, probably reflecting the requirement of two events for β-gal activity; namely induction of FLP and FLP-mediated excision of neo. These observations, together with the results of the control transfections, lead us to conclude that FLP-mediated, site-specific recombination events can be tightly regulated by the induced activation of an ecdysone-responsive promoter.

As mentioned, the expression of FLP recombinase or the activation of its enzyme activity has previously been successfully placed under the regulation of other inducible promoters. Using cell-specific promoters, its expression has also been activated in specific cells (4). In principle, the two-tiered regulatory system outlined here has the added advantage that judicious selection of a promoter to regulate the expression of the VgEcR and RXR elements should allow temporal induction of FLP-mediated changes in DNA in specific cells in response to muristerone A. To test whether this is in fact the case, we transfected ES with pK19VgRXR, pIND/FLP and pNEOβGAL. In pK19VgRXR, the expression of VgEcR and RXR is regulated by the promoter of the murine keratin 19 (K19) gene. K19 is known to be expressed upon the differentiation of ES cells into organized structures known as embryoid bodies (13). Following transfection, ES cells were grown under conditions that induce differentiation and treated with muristerone A. Twenty-four hours later, fixed cells were immunostained for both β-gal and K19. β-gal protein, the consequence of FLP recombinase mediated reconstitution of
a functional lacZ sequence, was only observed in cells that also expressed K19 (Figure 4). β-gal protein was not observed in ES cells that were not treated with muristerone A.

**DISCUSSION**

These studies demonstrate that by combining the genetic elements required for ecdysone induction and for FLP recombination, it is possible to temporally change gene expression in specific cells. This system potentially has several practical applications in cultured cells, transgenic animals (excluding insects) and possibly in transgenic plants as well. Similar to a simpler system in which the sequence encoding FLP recombinase is placed under the control of a cell-specific promoter, the system described here is useful for the analysis of cell lineages in developing systems both in vitro and in vivo. However, the ecdysone/FLP system has the added advantage that if a given promoter is active in specific cells throughout development, the starting point at which one begins to track a cell lineage can be regulated by the administration of ecdysone. This added capability increases the potential for studying the progeny of specific cells in tissues that are in advanced stages of differentiation as well as in self-renewing tissues in the adult organism. The conditional activation of genes that is possible using this system facilitates the study of transgenes whose expression during embryogenesis is lethal, e.g., overexpressed oncogenes or dominant negatives. Using this system, it should also be possible to identify the cells from which primary tumors arise and to track metastatic foci deriving from a primary tumor. The results of these experiments also suggest applications of this system to gene therapy strategies that are designed either to activate functional gene sequences in specific cell lineages at specified times or to ablate cells of a specified lineage.

Spatio-temporal expression of Cre recombinase has recently been achieved using tamoxifen induction of a modified ligand-binding domain of the human es-

### Table I. Number of Cells Expressing β-gal Activity after Transfection

<table>
<thead>
<tr>
<th>Plasmid DNA(s)</th>
<th>Concentration of muristerone A</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>none</td>
</tr>
<tr>
<td>pVgRXR + pIND/FLP + pNEOβGAL</td>
<td>24 ± 6^b</td>
</tr>
<tr>
<td>pFRTβGAL (Control 1)</td>
<td>3481 ± 905</td>
</tr>
<tr>
<td>pIND/LacZ (Control 2)</td>
<td>0</td>
</tr>
<tr>
<td>pVgRXR + pIND/LacZ (Control 3)</td>
<td>42 ± 11</td>
</tr>
<tr>
<td>pNEOβGAL (Control 4)</td>
<td>0</td>
</tr>
<tr>
<td>no DNA (Control 5)</td>
<td>0</td>
</tr>
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</table>

^a_Cells were electroporated with the indicated plasmid DNA(s) plus the plasmid pCX-EGFP. They were then cultured in the presence of 0, 1 or 5 µM muristerone A. Blue cells were counted and normalized to the number of cells expressing GFP.

^b_The results represent the mean ± SD of three independent experiments performed in triplicate.
trogen receptor (2). Regulatory systems such as this one and the ecdysone-based system described here clearly empower the utility of site-specific recombinases.

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REFERENCES


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