

peTurboYFP-PRL-dest1 vector

The vector sequence has been compiled using the informa- tion from sequence databases, published literature, and other sources, together with partial sequences obtained by Evrogen. This vector has not been completely sequenced.	Product	Cat.#	Size
	peTurboYFP-PRL-dest1 vector	FP616	20 µg
HSV TK poly A peTurboYFP-PRL- dest1 vector, 5.4kb Fl ori Psv40 SV40 poly A fl ori	Vector type Reporter Reporter codon usage Promoter for TurboYFP Host cells Selection Replication	promoterless expression TurboYFP mammalian NO mammalian prokaryotic - kanamycin eukaryotic - neomycin (G4 prokaryotic - pUC ori	vector 418)
For vector sequence, please visit our Web site at http://www.evrogen.com/products/vectors.shtml	Use	eukaryotic - SV40 ori Monitoring of activity of different promoters and promoter/enhancer combinations	
Multiple cloning site (MCS)			

* - not unique sites.

Location of features

MCS: 12-89 TurboYFP-dest1 Kozak consensus translation initiation site: 90-100 Start codon (ATG): 97-99 Last amino acid in TurboYFP: 823-825 Amino acid residues of mouse ornithine decarboxylase (MODC) PEST sequence: 847-966 Stop codon: 967-969 Fragment of human beta globin (HBB) gene Last 35 bp of HBB exon 2 : 978-1012 HBB intron 2: 974-1859 First 233 bp of HBB exon 3: 1860-2096 SV40 early mRNA polyadenylation signal Polyadenylation signals: 2238-2243 & 2267-2272 mRNA 3' ends: 2276 & 2288 f1 single-strand DNA origin: 2335-2790 Bacterial promoter for expression of Kan^r gene -35 region: 2852-2857; -10 region: 2875-2880 Transcription start point: 2887 SV40 origin of replication: 3131-3266 SV40 early promoter Enhancer (72-bp tandem repeats): 2964-3035 & 3036-3107 21-bp repeats: 3111-3131, 3132-3152 & 3154-3174 Early promoter element: 3187-3193 Major transcription start points: 3183, 3221, 3227 &

3232 Kanamycin/neomycin resistance gene

Neomycin phosphotransferase coding sequences: Start codon (ATG): 3315-3317; Stop codon: 4107-4109

G->A mutation to remove Pst I site: 3497 C->A (Arg to Ser) mutation to remove BssH II site: 3843 Herpes simplex virus (HSV) thymidine kinase (TK)

polyadenylation signal Polyadenylation signals: 4345-4350 & 4358-4363

pUC plasmid replication origin: 4694-5337

References

Haas, J. et al. (1996) "Codon usage limitation in the expression of HIV-1 envelope glycoprotein." Curr Biol, 6 (3): 315–324 / pmid: 8805248

Kozak, M. (1987) "An analysis of 5'-noncoding sequences from 699 vertebrate messenger RNAs." Nucleic Acids Res, 15 (20): 8125–8148 / pmid: 3313277

Li, X. et al. (1998) "Generation of destabilized green fluorescent protein as a transcription reporter." J Biol Chem, 273 (52): 34970–34975 / pmid: 9857028

Vector description

peTurboYFP-PRL-dest1 is a promoterless vector encoding destabilized variant of the yellow fluorescent protein TurboYFP, which can be used as *in vivo* reporter of promoter activity. To generate TurboYFP-dest1 variant, residues 422-461 of mouse ornithine decarboxylase (MODC) were fused to the TurboYFP C-terminus. This MODC region contains a PEST amino acid sequence that targets the protein for degradation and provides for rapid protein turnover [Li et al. 1998]. TurboYFP-dest1 retains fluorescent properties of the native protein and has a half-life of approximately 1-1.5 hours, as measured by fluorescence intensity of cells treated with the protein synthesis inhibitor, cycloheximide. Rapid TurboYFP-dest1 turnover allows accurate analysis of changes in gene regulation.

TurboYFP-dest1 codon usage is optimized for high expression in mammalian cells (humanized) [Haas et al. 1996]. To increase mRNA translation efficiency, Kozak consensus translation initiation site is generated upstream of the TurboYFP-dest1 coding sequence [Kozak 1987]. Fragments of exons 2 and 3 and intron 2 of human beta globin gene are added in the 3' UTR of TurboYFP-dest1 coding sequence in order to increase the protein expression level.

Multiple cloning site (MCS) is located upstream of the Kozak consensus translation initiation site and can be used to clone a promoter or a promoter/enchancer combination of interest. Without the addition of a functional promoter, this vector will not express TurboYFP-dest1.

The vector backbone contains SV40 origin for replication in mammalian cells expressing SV40 T-antigen, pUC origin of replication for propagation in *E. coli* and f1 origin for single-stranded DNA production. SV40 polyadenylation signals (SV40 poly A) direct proper processing of the 3'-end of the reporter mRNA.

SV40 early promoter (P_{SV40}) provides neomycin resistance gene (Neo^r) expression to select stably transfected eukaryotic cells using G418. Bacterial promoter (P) provides kanamycin resistance gene expression (Kan^r) in *E. coli*. Kan^r/Neo^r gene is linked with herpes simplex virus (HSV) thymidine kinase (TK) polyadenylation signals.

Note: The plasmid DNA was isolated from dam⁺-methylated *E.coli*. Therefore some restriction sites are blocked by methylation. If you wish to digest the vector using such sites you will need to transform the vector into a dam⁺ host and make fresh DNA.

Propagation in E. coli

Suitable host strains for propagation in *E. coli* include DH5alpha, HB101, XL1-Blue, and other general purpose strains. Plasmid incompatibility group is pMB1/ColE1. The vector confers resistance to kanamycin ($30 \mu g/ml$) to *E. coli* hosts. Copy number in *E. coli* is about 500.

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