miRNAs are initially transcribed as long precursor transcripts known as primary microRNAs (pri-miRNAs). Within these transcripts, the mature miRNA sequences are found in ~60–80 nucleotide hairpin structures. Mature miRNAs are generated from pri-miRNAs by sequential processing (Fig. 1). Pri-miRNAs are initially recognized in the nucleus by the microprocessor complex which includes as core components the RNase-III enzyme Drosha and its obligate partner DGCR8. This complex excises the hairpin structure containing the mature miRNA sequence. The liberated hairpins, referred to as precursor miRNAs (pre-miRNAs), are recognized by the nuclear export factor exportin 5 which transports them to the cytoplasm. There, the RNase-III enzyme Dicer performs a second cleavage to generate a double-stranded 18–24 nucleotide RNA molecule. The RISC then associates with this RNA duplex and unwinds it. Generally, only one strand is stably incorporated into the RISC; the other is discarded and rapidly degraded. miRNAs guide the RISC to target messages that are subsequently cleaved or translationally silenced.

**Figure 1. miRNA Biogenesis and function**
Synthetic miRNA molecules based on predicted mature miRNA sequence are sometimes used. Despite their optimized design criteria, synthetic miRNAs underscore the importance of primary miRNA in its native expressed form. The primary miRNA contains critical biological components involved in mature miRNA expression and cellular processing, and is often processed into several mature miRNA molecules.

Cell Biolabs’ miRNASelect™ precursor expression vectors express each individual human miRNA precursor in its native context while preserving putative hairpin structures to ensure biologically relevant interactions with endogenous processing machinery and regulatory partners, and that leads to properly cleaved microRNAs. Each individual miRNA precursor is cloned between BamHI and Nhe I sites.

The miRNASelect™ pEP-mir expression vectors contain the following features:
- **miRNA precursor** – 100 bp stem loop precursor in its native context flanked by a human intron sequence to preserve the putative hairpin structure and proper endogenous processing
- **EF-1α Promoter** - ensures a high level of expression in mammalian cells
- **Puromycin resistance marker** - to monitor cells positive for expression and stable selection
- **SV40 polyadenylation signal** - enables efficient termination of transcription
- **pUC origin** - for high copy replication and maintenance of the plasmid in *E. coli*
- **Ampicillin resistance gene** - for selection in *E. coli*

![Figure 2](image.png)

**Figure 2.** Schematic representation of the miRNASelect™ pEP-miR expression vector.

Cell Biolabs’ miRNASelect™ pEP-miR Null Control Vector is similar to the pEP-miR vector except it does not contain any miRNA precursor or the BamHI and NheI cloning sites (Figure 3).
Figure 3. Schematic representation of the miRNASElect™ pEP-miR null control vector.

Methods

1) Bacterial culture: the pEP-miR null control vector is provided as bacterial glycerol stock. Individual colonies can be obtained by culturing in an LB-ampicillin plate.
2) Plasmid isolation: we recommend EndoFree Plasmid Kits (QIAGEN).
3) Transfection into target cells: we recommend Lipofectamine 2000 (Invitrogen).
4) Stable selection: 48 hrs post-transfection, select stable clones in 1-10 µg/mL Puromycin-containing medium.

References

1. microRNA sequences listed in Sanger’s miRBase (http://microrna.sanger.ac.uk/sequences/).

Recent Product Citations

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