

## **Product datasheet for SR417835**

## OriGene Technologies, Inc.

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## **Clasrp Mouse siRNA Oligo Duplex (Locus ID 53609)**

## **Product data:**

**Guaranteed:** 

**Product Type:** siRNA Oligo Duplexes

Purity: HPLC purified

**Quality Control:** Tested by ESI-MS

Sequences: Available with shipment

**Stability:** One year from date of shipment when stored at -20°C.

# of transfections: Approximately 330 transfections/2nmol in 24-well plate under optimized conditions (final

conc. 10 nM).

**Note:** Single siRNA duplex (10nmol) can be ordered.

RefSeq: NM 016680, NR 125832, NR 125833, NR 125834, NR 125835, NR 125836

UniProt ID: Q8CFC7

Synonyms: Clasp; Sfr; Sfrs16; Srs; Srsf16; SW; Swap2

Components: Clasrp (Mouse) - 3 unique 27mer siRNA duplexes - 2 nmol each (Locus ID 53609)

Included - SR30004, Trilencer-27 Universal Scrambled Negative Control siRNA Duplex - 2 nmol

Included - SR30005, RNAse free siRNA Duplex Resuspension Buffer - 2 ml

Summary: The protein encoded by this gene contains serine/arginine (SR) dipeptide repeat domains,

and is thought to be involved in the regulation of alternative splicing. This protein is thought to interact with, and be phosphorylated by, Clk4. Alternative splicing results in multiple

transcript variants. [provided by RefSeq, Aug 2014]

**Performance** OriGene guarantees that at least two of the three Dicer-Substrate duplexes in the kit will

provide at least 70% or more knockdown of the target mRNA when used at 10 nM

concentration by quantitative RT-PCR when the TYE-563 fluorescent transfection control duplex (cat# SR30002) indicates that >90% of the cells have been transfected and the HPRT

positive control (cat# SR30003) provides 90% knockdown efficiency.

For non-conforming siRNA, requests for replacement product must be made within ninety (90) days from the date of delivery of the siRNA kit. To arrange for a free replacement with newly designed duplexes, please contact Technical Services at techsupport@origene.com. Please provide your data indicating the transfection efficiency and measurement of gene expression knockdown compared to the scrambled siRNA control (quantitative RT-PCR data

required).

