

## **Product datasheet for SC204905**

## OriGene Technologies, Inc.

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## Filamin A (FLNA) (NM 001456) Human 3' UTR Clone

**Product data:** 

**Product Type:** 3' UTR Clones

Product Name: Filamin A (FLNA) (NM\_001456) Human 3' UTR Clone

**Vector:** pMirTarget (PS100062)

Symbol: FLNA

Synonyms: ABP-280; ABPX; CSBS; CVD1; FGS2; FLN; FLN-A; FLN1; FMD; MNS; NHBP; OPD; OPD1; OPD2;

XLVD; XMVD

**ACCN:** NM\_001456

**Insert Size:** 348 bp

Insert Sequence: >SC204905 3'UTR clone of NM\_001456

The sequence shown below is from the reference sequence of NM\_001456. The complete

sequence of this clone may contain minor differences, such as SNPs.

Blue=Stop Codon Red=Cloning site

GGCAAGTTGGACGCCCGCAAGATCCGCGAGATTCTCATTAAGGCCAAGAAGGGCGGAAAGATCGCCGTG

TAACAATTGGCAGAGCTCAGAATTCAAGCGATCGCC

HC

CGAGATTTCGATTCCACCGCCGCCTTCTATGAAAGG

**Restriction Sites:** Sgfl-Mlul

**OTI Disclaimer:** Our molecular clone sequence data has been matched to the sequence identifier above as a

point of reference. Note that the complete sequence of this clone is largely the same as the

reference sequence but may contain minor differences, e.g., single nucleotide

polymorphisms (SNPs).

**Components:** The cDNA clone is shipped in a 2-D bar-coded Matrix tube as 10 ug dried plasmid DNA. The

package also includes 100 pmols of both the corresponding 5' and 3' vector primers in

separate vials.

RefSeq: <u>NM 001456.4</u>





## Filamin A (FLNA) (NM\_001456) Human 3' UTR Clone - SC204905

**Summary:** The protein encoded by this gene is an actin-binding protein that crosslinks actin filaments

and links actin filaments to membrane glycoproteins. The encoded protein is involved in remodeling the cytoskeleton to effect changes in cell shape and migration. This protein interacts with integrins, transmembrane receptor complexes, and second messengers. Defects in this gene are a cause of several syndromes, including periventricular nodular heterotopias (PVNH1, PVNH4), otopalatodigital syndromes (OPD1, OPD2), frontometaphyseal dysplasia (FMD), Melnick-Needles syndrome (MNS), and X-linked congenital idiopathic

intestinal pseudoobstruction (CIIPX). Two transcript variants encoding different isoforms have

been found for this gene.[provided by RefSeq, Mar 2009]

**Locus ID:** 2316

MW: 12.3