

Product datasheet for SC311618

OriGene Technologies, Inc. 9620 Medical Center Drive, Ste 200

Rockville, MD 20850, US
Phone: +1-888-267-4436
https://www.origene.com
techsupport@origene.com
EU: info-de@origene.com
CN: techsupport@origene.cn

SH3TC2 (AL832651) Human Untagged Clone

Product data:

Product Type: Expression Plasmids

Product Name: SH3TC2 (AL832651) Human Untagged Clone

Tag: Tag Free Symbol: SH3TC2

Synonyms: CMT4C; MNMN

Vector: pCMV6 series

Fully Sequenced ORF: >NCBI ORF sequence for AL832651, the custom clone sequence may differ by one or more

nucleotides

Restriction Sites: Please inquire

ACCN: AL832651

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

point of reference. Note that the complete sequence of our molecular clones may differ from the sequence published for this corresponding reference, e.g., by representing an alternative

RNA splicing form or single nucleotide polymorphism (SNP).

OTI Annotation: This TrueClone is provided through our Custom Cloning Process that includes sub-cloning

into OriGene's pCMV6 vector and full sequencing to provide a non-variant match to the expected reference without frameshifts, and is delivered as lyophilized plasmid DNA.

Components: The ORF clone is ion-exchange column purified and shipped in a 2D barcoded Matrix tube

containing 10ug of transfection-ready, dried plasmid DNA (reconstitute with 100 ul of water).

Reconstitution Method: 1. Centrifuge at 5,000xg for 5min.

2. Carefully open the tube and add 100ul of sterile water to dissolve the DNA.

3. Close the tube and incubate for 10 minutes at room temperature.

4. Briefly vortex the tube and then do a quick spin (less than 5000xg) to concentrate the liquid

at the bottom.

5. Store the suspended plasmid at -20°C. The DNA is stable for at least one year from date of

shipping when stored at -20°C.

 RefSeq:
 AL832651.2

 RefSeq Size:
 4525 bp





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 RefSeq ORF:
 4525 bp

 Locus ID:
 79628

 Cytogenetics:
 5q32

Gene Summary: This gene encodes a protein with two N-terminal Src homology 3 (SH3) domains and 10

tetratricopeptide repeat (TPR) motifs, and is a member of a small gene family. The gene product has been proposed to be an adapter or docking molecule. Mutations in this gene result in autosomal recessive Charcot-Marie-Tooth disease type 4C, a childhood-onset neurodegenerative disease characterized by demyelination of motor and sensory neurons.

[provided by RefSeq, Jul 2008]