

## Product datasheet for RC223206L1V

## OriGene Technologies, Inc.

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## DNMT3B (NM 006892) Human Tagged ORF Clone Lentiviral Particle

**Product data:** 

**Product Type:** Lentiviral Particles

**Product Name:** DNMT3B (NM\_006892) Human Tagged ORF Clone Lentiviral Particle

Symbol:

ICF; ICF1; M.HsallIB Synonyms:

**Mammalian Cell** 

Selection:

None

Vector: pLenti-C-Myc-DDK (PS100064)

Myc-DDK Tag: NM 006892 ACCN:

**ORF Size:** 2559 bp

**ORF Nucleotide** 

OTI Disclaimer:

Sequence:

The ORF insert of this clone is exactly the same as(RC223206).

The molecular sequence of this clone aligns with the gene accession number as a point of reference only. However, individual transcript sequences of the same gene can differ through

naturally occurring variations (e.g. polymorphisms), each with its own valid existence. This clone is substantially in agreement with the reference, but a complete review of all prevailing

variants is recommended prior to use. More info

**OTI Annotation:** This clone was engineered to express the complete ORF with an expression tag. Expression

varies depending on the nature of the gene.

RefSeq: NM 006892.3

RefSeq Size: 4353 bp RefSeq ORF: 2562 bp Locus ID: 1789 **UniProt ID:** Q9UBC3

20q11.21 Cytogenetics:

**Domains:** PWWP, DNA\_methylase



## DNMT3B (NM\_006892) Human Tagged ORF Clone Lentiviral Particle - RC223206L1V

Protein Families: Druggable Genome, Embryonic stem cells, Induced pluripotent stem cells, Stem cell -

Pluripotency

**Protein Pathways:** Cysteine and methionine metabolism, Metabolic pathways

**MW:** 95.8 kDa

**Gene Summary:** CpG methylation is an epigenetic modification that is important for embryonic development,

imprinting, and X-chromosome inactivation. Studies in mice have demonstrated that DNA

methylation is required for mammalian development. This gene encodes a DNA methyltransferase which is thought to function in de novo methylation, rather than

maintenance methylation. The protein localizes primarily to the nucleus and its expression is developmentally regulated. Mutations in this gene cause the immunodeficiency-centromeric instability-facial anomalies (ICF) syndrome. Eight alternatively spliced transcript variants have been described. The full length sequences of variants 4 and 5 have not been determined.

[provided by RefSeq, May 2011]