

Product datasheet for KN222919BN

WRN Human Gene Knockout Kit (CRISPR)

Product data:

Product Type: Knockout Kits (CRISPR)

Format: 2 gRNA vectors, 1 mBFP-Neo donor, 1 scramble control

Donor DNA: mBFP-Neo

Symbol: WRN 7486 Locus ID:

KN222919G1, WRN gRNA vector 1 in pCas-Guide CRISPR vector (GE100002) Components:

KN222919G2, WRN gRNA vector 2 in pCas-Guide CRISPR vector (GE100002)

KN222919BND, donor DNA containing left and right homologous arms and mBFP-Neo

functional cassette.

GE100003, scramble sequence in pCas-Guide vector

Disclaimer: These products are manufactured and supplied by OriGene under license from ERS. The kit is

> designed based on the best knowledge of CRISPR technology. The system has been functionally validated for knocking-in the cassette downstream the native promoter. The efficiency of the knock-out varies due to the nature of the biology and the complexity of the

experimental process.

RefSeq: NM 000553

UniProt ID: Q14191

Synonyms: RECQ3; RECQL2; RECQL3

Summary: This gene encodes a member of the RecQ subfamily of DNA helicase proteins. The encoded

> nuclear protein is important in the maintenance of genome stability and plays a role in DNA repair, replication, transcription and telomere maintenance. This protein contains a Nterminal 3' to 5' exonuclease domain, an ATP-dependent helicase domain and RQC (RecQ helicase conserved region) domain in its central region, and a C-terminal HRDC (helicase RNase D C-terminal) domain and nuclear localization signal. Defects in this gene are the cause of Werner syndrome, an autosomal recessive disorder characterized by accelerated

aging and an elevated risk for certain cancers. [provided by RefSeq, Aug 2017]



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Product images:

Donor Vector Edited Chromosome



RFP, Luc, and mBFP will be under native gene promoter