

Product datasheet for KN222919RB

WRN Human Gene Knockout Kit (CRISPR)

Product data:

OriGene Technologies, Inc.

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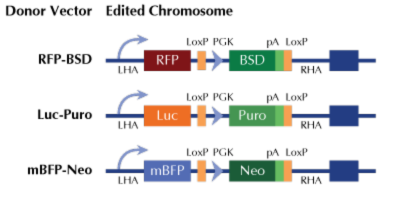
Product Type:	Knockout Kits (CRISPR)
Format:	2 gRNA vectors, 1 RFP-BSD donor, 1 scramble control
Donor DNA:	RFP-BSD
Symbol:	WRN
Locus ID:	7486
Components:	 KN222919G1, WRN gRNA vector 1 in pCas-Guide CRISPR vector (GE100002) KN222919G2, WRN gRNA vector 2 in pCas-Guide CRISPR vector (GE100002) KN222919RBD, donor DNA containing left and right homologous arms and RFP-BSD 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:	<u>NM 000553</u>
UniProt ID:	<u>Q14191</u>
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 N-terminal 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:



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

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