

Rabbit Anti-FAM91A1 antibody

SL8210R

Product Name:	FAM91A1
Chinese Name:	FAM91A1蛋白抗体
Alias:	FAM91A1; Family with sequence similarity 91 member A1; FLJ23790; Hypothetical
	protein LOC157769; LOC157769; F91A1_HUMAN.
Organism Species:	Rabbit
Clonality:	Polyclonal
React Species:	Human, Mouse, Rat, Chicken,
Applications:	WB=1:500-2000ELISA=1:500-1000IHC-P=1:400-800IHC-F=1:400-800IF=1:100-
	500 (Paraffin sections need antigen repair)
	not yet tested in other applications.
	optimal dilutions/concentrations should be determined by the end user.
Molecular weight:	94kDa
Cellular localization:	cytoplasmic
Form:	Lyophilized or Liquid
Concentration:	1mg/ml
immunogen:	KLH conjugated synthetic peptide derived from human FAM91A1:335-440/838
Lsotype:	IgG
Purification:	affinity purified by Protein A
Storage Buffer:	0.01M TBS(pH7.4) with 1% BSA, 0.03% Proclin300 and 50% Glycerol.
Storage:	Store at -20 °C for one year. Avoid repeated freeze/thaw cycles. The lyophilized
	antibody is stable at room temperature for at least one month and for greater than a year
	when kept at -20°C. When reconstituted in sterile pH 7.4 0.01M PBS or diluent of
	antibody the antibody is stable for at least two weeks at 2-4 °C.
PubMed:	<u>PubMed</u>
Product Detail:	Made up of nearly 146 million bases, chromosome 8 encodes about 800 genes.
	Translocation of portions of chromosome 8 with amplifications of the c-Myc gene are
	found in some leukemias and lymphomas, and typically associated with a poor
	prognosis. Portions of chromosome 8 have been linked to schizophrenia and bipolar
	disorder. Trisomy 8, also known as Warkany syndrome 2, most often results in early
	miscarriage but is occasionally seen in a mosaic form in surviving patients who suffer

to a varying degree from a number of symptoms including retarded mental and motor development, and certain facial and developmental defects. WRN is a DNA helicase encoded by chromosome 8 and shown defective in those with the early aging disorder Werner syndrome. Chromosome 8 is also associated with Pfeiffer syndrome, congenital hypothyroidism and Waardenburg syndrome. The FAM91A1 gene product has been provisionally designated FAM91A1 pending further characterization.

Similarity:

Belongs to the FAM91 family.

SWISS:

Q658Y4

Gene ID:

157769

Database links:

Entrez Gene: 157769Human

SwissProt: Q658Y4Human

<u>Unigene: 459174</u>Human

Important Note:

This product as supplied is intended for research use only, not for use in human, therapeutic or diagnostic applications.