

# Rabbit Anti-C6orf191 antibody

## SL15233R

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Product Name:	C6orf191
Chinese Name:	6号染色体开放阅读框191抗体
Alias:	bA174C7.4; C6orf191; TM244_HUMAN; Chromosome 6 open reading frame 191;
	Putative transmembrane protein C6orf191.
Organism Species:	Rabbit
Clonality:	Polyclonal
React Species:	Human,Pig,Cow,Horse,Sheep,
Applications:	WB=1:500-2000ELISA=1:500-1000IHC-P=1:400-800IHC-F=1:400-800ICC=1:100-
	500IF=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:	15kDa
Cellular localization:	The cell membrane
Form:	Lyophilized or Liquid
Concentration:	1mg/ml
immunogen:	KLH conjugated synthetic peptide derived from human C6orf191:31-128/128
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:	Making up nearly 6% of the human genome, chromosome 6 contains around 1,200
	genes within 170 million base pairs of sequence. Deletion of a portion of the q arm of
	chromosome 6 is associated with early onset intestinal cancer suggesting the presence
	of a cancer susceptibility locus. Porphyria cutanea tarda is associated with chromosome
	6 through the HFE gene which, when mutated, predisposes an individual to developing
	this porphyria. Notably, the PARK2 gene, which is associated with Parkinson's disease,

and the genes encoding the major histocompatibility complex proteins, which are key molecular components of the immune system and determine predisposition to rheumatic diseases, are also located on chromosome 6. Stickler syndrome, 21-hydroxylase deficiency and maple syrup urine disease are also associated with genes on chromosome 6. A bipolar disorder susceptibility locus has been identified on the q arm of chromosome 6. The C6orf191 gene product has been provisionally designated C6orf191 pending further characterization.

#### **Subcellular Location:**

Membrane; Multi-pass membrane protein (Potential).

SWISS: Q5VVB8

Gene ID: 253582

#### Database links:

Entrez Gene: 253582Human

SwissProt: Q5VVB8Human

Unigene: 448372Human

### **Important Note:**

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