## bs-15051R

## [ Primary Antibody ]

## C1orf192 Rabbit pAb



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Host: Rabbit Clonality: Polyclonal GenelD: 257177	Isotype: IgG SWISS: Q5VTH2	Applications: IHC-P (1:100-500) IHC-F (1:100-500) IF (1:100-500) ICC/IF (1:100-500) ELISA (1:5000-10000)
Target: Clorf192 Immunogen: KLH conjugated synthetic peptide derived from human Clorf192: 11-80/177. Purification: affinity purified by Protein A		
Glycerol. Shipped at 4°C. freeze/thaw cyc Background: Chromosome 1 260 million bas There are abou the great numb diseases associ disease Hutchin gene which enc product can bu nuclear blebs. T and is a topic o located on chrc adenomatous p disease and Us 1. A breakpoint gene and is link are found in a v malignant mela	is the largest human chromosome spanning about e pairs and making up 8% of the human genome. t 3,000 genes on chromosome 1, and considering wer of genes there are also a large number of ated with chromosome 1. Notably, the rare aging hson-Gilford progeria is associated with the LMNA codes lamin A. When defective, the LMNA gene ild up in the nucleus and cause characteristic The mechanism of rapidly enhanced aging is uncle f continuing exploration. The MUTYH gene is bomosome 1 and is partially responsible for familia polyposis. Stickler syndrome, Parkinsons, Gauche her syndrome are also associated with chromosome has been identified in 1q which disrupts the DISC ted to schizophrenia. Aberrations in chromosome ariety of cancers including head and neck cancer, anoma and multiple myeloma. The C1orf192 gene en provisionally designated C1orf192 pending	ear I r me 1