

# MFRP Rabbit pAb

CatalogNo: YT6216

# **Key Features**

Host Species Reactivity Applications
• Rabbit • Human, Mouse • IHC, IF, WB

MW Isotype • 62kD (Observed) IgG

### **Recommended Dilution Ratios**

IHC 1:50-200 WB 1:500-2000 IF 1:50-200

# Storage

Storage\* -15°C to -25°C/1 year(Do not lower than -25°C)

**Formulation** Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

# **Basic Information**

**Clonality** Polyclonal

# Immunogen Information

Immunogen Synthesized peptide derived from human MFRP

**Specificity** This antibody detects endogenous levels of human MFRP

# | Target Information

Gene name MFRP

#### **Protein Name**

**MFRP** 

**Gene ID** UniProt ID **Organism** Human 114902: Q9BY79;

### Cellular Localization

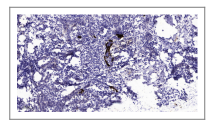
Apical cell membrane; Single-pass type II membrane protein.

**Tissue specificity** Specifically expressed in brain. Strongly expressed in medulla oblongata and to a lower extent in hippocampus and corpus callosum. Expressed in keratinocytes.

#### **Function**

developmental stage: Expressed in fetal brain., Disease: Defects in C1QTNF5 are a cause of late-onset retinal degeneration (LORD) [MIM:605670]. LORD is an autosomal dominant disorder characterized by onset in the fifth to sixth decade with night blindness and punctate yellow-white deposits in the retinal fundus, progressing to severe central and peripheral degeneration, with choroidal neovascularization and chorioretinal atrophy., Disease: Defects in MFRP are the cause of microphthalmia MFRP-related (MCOPMFRP) [MIM:611040]. Microphthalmia is a clinically heterogeneous disorder of eye formation, ranging from small size of a single eye to complete bilateral absence of ocular tissues. Ocular abnormalities like opacities of the cornea and lens, scaring of the retina and choroid, cataract and other abnormalities like cataract may also be present. MCOPMFRP is characterized by posterior microphthalmia, retinitis pigmentosa, foveoschisis and optic disc drusen., Disease: Defects in MFRP are the cause of nanophthalmos 2 (NNO2) [MIM:609549]. NNO2 is a rare autosomal recessive disorder of eye development characterized by extreme hyperopia and small functional eyes., Function: May play a role in eye development., similarity: Contains 1 C1q domain., similarity: Contains 1 collagen-like domain., similarity: Contains 1 FZ (frizzled) domain., similarity: Contains 2 CUB domains., similarity: Contains 2 LDL-receptor class A domains., tissue specificity: Specifically expressed in brain. Strongly expressed in medulla oblongata and to a lower extent in hippocampus and corpus callosum. Expressed in keratinocytes.,

### **| Validation Data**



Immunohistochemical analysis of paraffin-embedded human Breast cancer. 1, Antibody was diluted at 1:200(4° overnight). 2, Tris-EDTA,pH9.0 was used for antigen retrieval. 3, Secondary antibody was diluted at 1:200 (room temperature, 45min).

## | Contact information

Orders: order@immunoway.com Support: tech@immunoway.com

Telephone: 877-594-3616 (Toll Free), 408-747-0185

Website: http://www.immunoway.com

Address: 2200 Ringwood Ave San Jose, CA 95131 USA



Please scan the QR code to access additional product information: MFRP Rabbit pAb

For Research Use Only. Not for Use in Diagnostic Procedures.	Antibody   ELISA Kits   Protein   Reagents
	Immunoway - 3 / 3