



## GPR143 rabbit pAb

**Cat#: orb769312 (Manual)** 

For research use only. Not intended for diagnostic use.

**Product Name** GPR143 rabbit pAb

**Host species** Rabbit

**Applications** IF;ELISA

**Species Cross-Reactivity** Human; Mouse

**Recommended dilutions** Immunofluorescence: 1/200 - 1/1000. ELISA: 1/10000. Not yet tested in

other applications.

**Immunogen** The antiserum was produced against synthesized peptide derived from

human GPR143. AA range:151-200

GPR143 Polyclonal Antibody detects endogenous levels of GPR143 protein. **Specificity** 

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

azide..

Store at -20°C. Avoid repeated freeze-thaw cycles. **Storage** 

**Protein Name** G-protein coupled receptor 143

**GPR143** Gene Name

Cellular localization Melanosome membrane; Multi-pass membrane protein . Lysosome

membrane; Multi-pass membrane protein. Apical cell membrane; Multi-pass membrane protein. Distributed throughout the endo-melanosomal system but most of endogenous protein is localized in unpigmented stage II melanosomes. Its expression on the apical cell membrane is sensitive to tyrosine (PubMed:18828673).



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Purification The antibody was affinity-purified from rabbit antiserum by affinity-

epitope-specific immunogen. chromatography using

**Clonality** Polyclonal

Concentration 1 mg/ml

**Observed band** 

**Human Gene ID** 4935

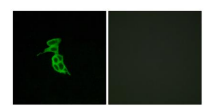
**Human Swiss-Prot Number** P51810

**Alternative Names** GPR143; OA1; G-protein coupled receptor 143; Ocular albinism type 1

protein

**Background** This gene encodes a protein that binds to heterotrimeric G proteins and is

targeted to melanosomes in pigment cells. This protein is thought to be involved in intracellular signal transduction mechanisms. Mutations in this gene cause ocular albinism type 1, also referred to as Nettleship-Falls type ocular albinism, a severe visual disorder. A related pseudogene has been identified on chromosome Y. [provided by RefSeq, Dec 2009],



Immunofluorescence analysis of LOVO cells, using GPR143 Antibody. The picture on the right is blocked with the synthesized peptide.