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# Niemann Pick C1 (YD14389) Rabbit mAb

货号: **AYD15316**

## 产品信息

反应	Human,Mouse,Rat
宿主	Rabbit
克隆性	Monoclonal
预测反应	
应用	WB IHC-P ICC/IF FC
推荐浓度	
理论分子量	142kDa
实测分子量	
形式	Liquid
保存条件	Store at -20°C. Avoid freeze / thaw cycles. Buffer: PBS with 0.75% BSA,50% glycerol,pH7.3.
偶联物	Unconjugated
阳性对照	A-549,Mouse liver,Rat lung
细胞定位	Late endosome membrane, Lysosome membrane
纯化	亲和纯化

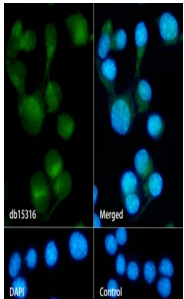
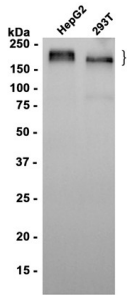
## 抗原信息

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## 靶点信息

研究背景	This gene encodes a large protein that resides in the limiting membrane of endosomes and lysosomes and mediates intracellular cholesterol trafficking via binding of cholesterol to its N-terminal domain. It is predicted to have a cytoplasmic C-terminus, 13 transmembrane domains, and 3 large loops in the lumen of the endosome - the last loop being at the N-terminus. This protein transports low-density lipoproteins to late endosomal/lysosomal compartments where they are hydrolyzed and released as free cholesterol. Defects in this gene cause Niemann-Pick type C disease, a rare autosomal recessive neurodegenerative disorder characterized by over accumulation of cholesterol and glycosphingolipids in late endosomal/lysosomal compartments.
基因ID	4864
基因名	NPC1
Swiss	O15118 ( <a href="https://www.uniprot.org/uniprotkb/O15118/entry">https://www.uniprot.org/uniprotkb/O15118/entry</a> )
别名	Niemann Pick C1 (YD14389),Niemann Pick C1 (YD14389) Rabbit mAb,NPC1,Niemann-Pick C1 protein

## 产品验证



## 实验步骤

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