

# ACADM (YD18875) Rabbit mAb

货号: **AYD12189**

## 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 IP
推荐浓度	
理论分子量	47kDa
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	HT-29,HepG2,U-251MG,K-562,Mouse liver,Mouse kidney,Mouse brain,Mouse heart,Rat liver
细胞定位	Mitochondrion matrix
纯化	

## 抗原信息

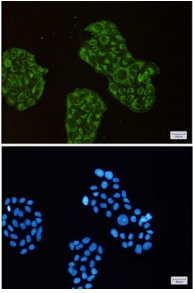
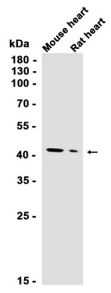
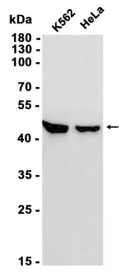
抗原信息	
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## 靶点信息

研究背景	This gene encodes the medium-chain specific (C4 to C12 straight chain) acyl-Coenzyme A dehydrogenase . The homotetramer enzyme catalyzes the initial step of the mitochondrial fatty acid beta-oxidation path way. Defects in this gene cause medium-chain acyl-CoA dehydrogenase deficiency, a disease characterized by hepatic dysfunction, fasting hypoglycemia, and encephalopathy, which can result in infantile death. Alternatively spliced transcript variants encoding different isoforms have been found for this gene.
基因ID	34
基因名	ACADM
Swiss	P11310

别名	ACADM (YD18875)
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## 产品验证



## 实验步骤

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