

SPRTN Rabbit pAb

货号: **AYP19941**

产品信息

反应	Human,Mouse,Rat
宿主	Rabbit
克隆性	Polyclonal
预测反应	
应用	WB
推荐浓度	WB: 1:500 - 1:2000
理论分子量	24kDa/29kDa/55kDa
实测分子量	60kDa
形式	Liquid
保存条件	Store at -20°C. Avoid freeze / thaw cycles. Buffer: PBS with 0.01% thiomersal,50% glycerol,pH7.3.
偶联物	Unconjugated
阳性对照	U-87MG
细胞定位	Chromosome,Nucleus
纯化	Affinity purification

抗原信息

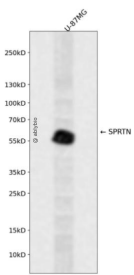
抗原信息	Recombinant fusion protein containing a sequence corresponding to amino acids 1-240 of human SPRTN (NP_001010984.1).
序列	MDDDLMLALRLQEEWNLQEAERDHAQESLSLVDASWELVDPTDQLQALFVQFNDQFFWGWQLEAVEVKWSVRMTLCAG ICSYEGKGGMCSIRLSEPLLKLRPRKDLVETLLHEMIHAYLFTNNDKDREGHGPEFCKHMRINSLTGANITVYHTFHDEV DEYRRHWRCNGPCQHRPPYYGYVKRATNREPSAHDYWWAEHQKTCGGTYIKIKEPENYSKKGKAKLGKEPVLAAE NKG

靶点信息

研究背景	The protein encoded by this gene may play a role in DNA repair during replication of damaged DNA. This protein recruits valosin containing protein (p97) to stalled DNA replication forks where it may prevent excessive translesional DNA synthesis and limit the number of DNA-damage induced mutations. It may also be involved in replication-related G2/M-checkpoint regulation. Deficiency of a similar protein in mouse causes chromosomal instability and progeroid phenotypes. Mutations in this gene have been associated with Ruijs-Aalfs syndrome (RJALS). Alternatively spliced transcript variants have been identified.
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基因ID	83932
基因名	SPRTN
Swiss	Q9H040
别名	SPRTN;C1orf124;DVC1;PRO4323;spartan

产品验证



Western blot analysis of SPRTN expressed in U-87MG using SPRTN Rabbit pAb at 1:1000. Secondary antibody: HRP Goat Anti-Rabbit IgG (H+L) at 1:5000. Lysates/proteins: 30ug per lane. Blocking buffer: 5% non-fat dry milk in TBST. Detection: ECL Enhanced Kit. Exposure time: 120s.

实验步骤

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