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Anti-Human/Mouse/Rat OCLN Polyclonal Antibody

多克隆抗体

产品货号: PA12522

第三版

描述:This gene encodes an integral membrane protein that is required for cytokine-induced regulation of the tight junction paracellular permeability barrier. Mutations in this gene are thought to be a cause of band-like calcification with simplified gyration and polymicrogyria (BLC-PMG), an autosomal recessive neurologic disorder that is also known as pseudo-TORCH syndrome. Alternative splicing results in multiple transcript variants. A related pseudogene is present 1.5 Mb downstream on the garm of chromosome 5.

抗原:Recombinant protein of human OCLN

配方:

如何使用:加1ml超纯水重溶

稳定性: -20°C保存条件下,冻干粉,保质期为五年;液体,保质期为两年。

稀释液:PBS (pH7.4) , 1% BSA

应用:WB1~5 μ g/ml.

特异性:Localized at tight junctions of both epithelial and endothelial cells. Highly expressed in kidney. Not detected in testis.

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