

anti- Alpha galactosidase A antibody

Product Information

Catalog No.:	FNab00330
Size:	100μ g
Form:	liquid
Purification:	Protein A+G purification
Purity:	≥95% as determined by SDS-PAGE
Host:	Mouse
IsoType:	IgG2a
Storage:	PBS with 0.02% sodium azide and 50% glycerol pH 7.3 , -20°C for 12 months (Avoid repeated freeze / thaw cycles.)

Background

GLA, also named as Melibiase, Agalsidase and Alpha-galactosidase A, belongs to the glycosyl hydrolase 27 family. It hydrolyzes terminal, non-reducing alpha-D-galactose residues in alpha-D-galactosides, including galactose oligosaccharides, galactomannans and galactolipids. Fabry disease is an X-linked lysosomal storage disorder resulting from the deficient activity of GLA. Enzyme replacement therapy (ERT) with GLA is currently the most effective therapeutic strategy for patients with Fabry disease, a lysosomal storage disease.

Immunogen information

Immunogen:	galactosidase, alpha
Synonyms:	Alpha galactosidase A, galactosidase, alpha
Calculated MW:	49kDa
Uniprot ID :	P06280

Application

Reactivity:	Human
Tested Application:	ELISA, IHC, WB, IF
Recommended dilution:	WB: 1:500-1:2000; IHC: 1:20-1:200; IF: 1:20-1:200

Wuhan Fine Biotech Co., Ltd.

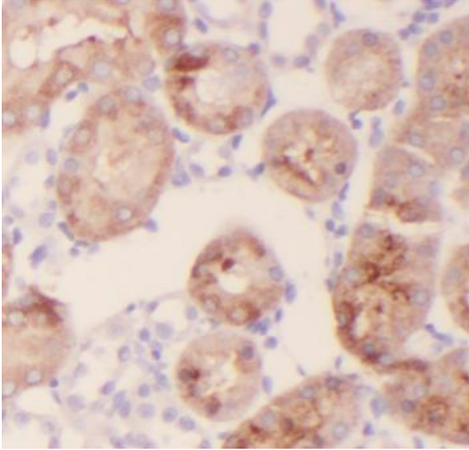
B9 Bld, High-Tech Medical Devices Park, No. 818 Gaoxin Ave. East Lake High-Tech Development Zone. Wuhan, Hubei, China(430206)

Tel : (0086)027-87384275

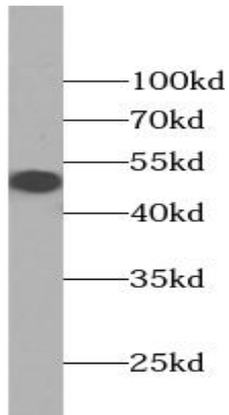
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Image:



Immunohistochemistry of paraffin-embedded human kidney tissue slide using FNab00330(GLA Antibody) at dilution of 1:50



HeLa cells were subjected to SDS PAGE followed by western blot with FNab00330(GLA antibody) at dilution of 1:1000

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