

Plxdc2 antibody

Cat. No. GTX85419

Host	Rabbit
Clonality	Polyclonal
Isotype	IgG
Applications	WB, IHC-P, ELISA
Reactivity	Human, Mouse, Rat

Package
100 µg

Applications

Application Note

*Optimal dilutions/concentrations should be determined by the researcher.

Suggested dilution	Recommended dilution
WB	0.5 - 1 µg/mL
IHC-P	2.5 µg/mL
ELISA	Assay dependent

Not tested in other applications.

Calculated MW	60 kDa. (Note)
Product Note	This antibody is specific for Plxdc2 C-Terminus

Properties

Form	Liquid
Buffer	PBS
Preservative	0.02% Sodium azide
Storage	Store as concentrated solution. Centrifuge briefly prior to opening vial. For short-term storage (1-2 weeks), store at 4°C. For long-term storage, aliquot and store at -20°C or below. Avoid multiple freeze-thaw cycles.
Concentration	1 mg/ml (Please refer to the vial label for the specific concentration.)
Immunogen	Plxdc2 antibody was raised against a 17 amino acid synthetic peptide from near the carboxy terminus of human Plxdc2. The immunogen is located within the last 50 amino acids of Plxdc2.
Purification	Purified by antigen-affinity chromatography
Conjugation	Unconjugated



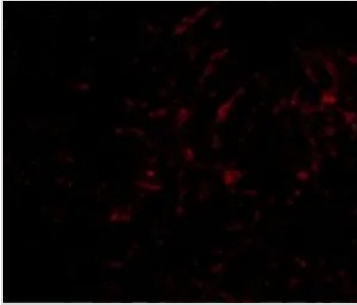
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Note

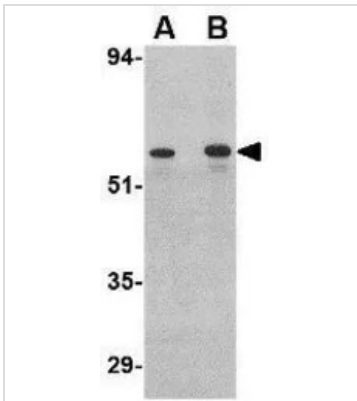
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DATA IMAGES



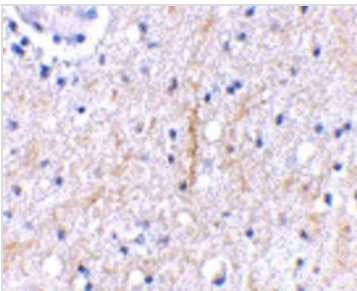
GTx85419 IHC-P Image

IHC-P analysis of human brain tissue using GTx85419 Plxdc2 antibody.
Working concentration : 50 µg/ml



GTx85419 WB Image

WB analysis of human colon tissue lysate using GTx85419 Plxdc2 antibody.
Working concentration : (A) 0.5 and (B) 1 µg/ml



GTx85419 IHC-P Image

IHC-P analysis of human brain tissue using GTx85419 Plxdc2 antibody.
Working concentration : 2.5 µg/ml



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