

Product Data Sheet

Anti-gfap Antibody

Catalog #	Source	Reactivity	Applications		
CQA9902	Rabbit	Z	WB, IH		
Description	Ra	bbit polyclonal antibod	/ to gfap		
Immunogen	KL	H-conjugated synthetic	peptide encompassing a sequence within the N-terminal		
	re	region of zebrafish gfap. The exact sequence is proprietary.			
Purification		The antibody was purified by immunogen affinity chromatography.			
Specificity Recognizes endogenous levels of gfap protein.					
Clonality	Ро	Polyclonal			
Conjugation					
Form		Liquid in 0.42% Potassium phosphate, 0.87% Sodium chloride, pH 7.3, 30% glycerol,			
	an	d 0.01% sodium azide.			
Dilution	W	B (1/500 - 1/1000), IH (1/	50 - 1/200)		
Gene Symbol GFA		AP			
Alternative Na	ames				
Entrez Gene		30646 (Human)			
SwissProt	Q	58EE9 (Human)			
Storage/Stabi	lity Sh	ipped at 4°C. Upon deliv	very aliquot and store at -20°C for one year. Avoid		
	fre	eeze/thaw cycles.			
Alternative Na Entrez Gene SwissProt	ames 30 Q5	9646 (Human) 58EE9 (Human)	very aliquet and store at 20°C for one year. Avoid		

Application key: E- ELISA, WB- Western blot, IH- Immunohistochemistry, IF- Immunofluorescence, FC- Flow cytometry, IC-Immunocytochemistry, IP- Immunoprecipitation, ChIP- Chromatin Immunoprecipitation, EMSA- Electrophoretic Mobility Shift Assay, BL- Blocking, SE- Sandwich ELISA, CBE- Cell-based ELISA, RNAi- RNA interference Species reactivity key: H- Human, M- Mouse, R- Rat, B- Bovine, C- Chicken, D- Dog, G- Goat, Mk- Monkey, P- Pig, Rb-Rabbit, S- Sheep, Z- Zebrafish

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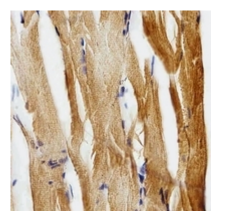
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For research purposes only, not for human use

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Western blot analysis of gfap expression in zebrafish brain (A) whole cell lysates. (Predicted band size: 51 kD; Observed band size: 50 kD)



Immunohistochemical analysis of gfap staining in zebrafish muscle formalin fixed paraffin embedded tissue section. The section was pre-treated using heat mediated antigen retrieval with sodium citrate buffer (pH 6.0). The section was then incubated with the antibody at room temperature and detected using an HRP conjugated compact polymer system. DAB was used as the chromogen. The section was then counterstained with haematoxylin and mounted with DPX.

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