

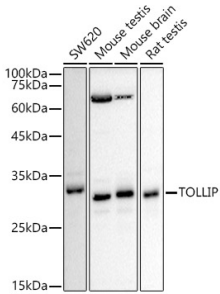
Anti-Tollip Antibody (A307074)

Specifications:

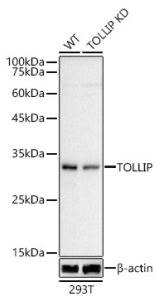
Name:	Anti-Tollip Antibody
Description:	Rabbit polyclonal antibody to Tollip.
Applications:	WB
Recommended Dilutions:	WB: 1:100-1:500
Reactivity:	Human, Mouse, Rat
Immunogen:	Recombinant fusion protein containing a sequence corresponding to amino acids 1-274 of human TOLLIP (NP_061882.2).
Sequence:	MATTVSTQRGPVYIGELPQDFLRITPTQQQRQVQLDAQAAQQLQYGGAVGTVGRLNIT VVQAKLAKNYGMTRMDPYCRLRLGYAVYETPTAHNGAKNPRWNKVIHCTVPPGVDSFY LEIFDERAFSMDDRIAWTHITIPESLRQGKVEDKWYSLSGRQGDDKEGMINLVMSYAL LPAAMVMPPQPVVLMP TVYQQGVGYVPITGMPAVCSPGMVPVALPPAAVNAQPRCSEE DLKAIQDMFPNMDQEVIRSVLEAQRGNKDAAINSLLQMGEEP
Host:	Rabbit
Clonality:	Polyclonal
Isotype:	IgG
Conjugate:	Unconjugated
Purification:	Affinity purification.
Molecular Weight:	30 kDa
Product Form:	Liquid
Formulation:	Supplied in Phosphate Buffered Saline, pH 7.3, with 50% Glycerol and 0.05% Proclin 300.
Storage:	Shipped at 4°C. Upon delivery aliquot and store at -20°C. Avoid freeze / thaw cycles.
Disclaimer:	This product is for research use only. It is not intended for diagnostic or therapeutic use.

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Images:



Western blot analysis of extracts of various cell lines, using Anti-Tollip Antibody (A307074) at 1:500 dilution. The secondary antibody was Goat Anti-Rabbit IgG H&L Antibody (HRP) at 1:10,000 dilution. Lysates/proteins were present at 25 μ g per lane. The blocking buffer used was 3% non-fat dry milk in TBST. Detection was with a ECL Basic Kit. Exposure time: 3S.



Western blot analysis of extracts from wild type (WT) and TOLLIP knockdown (KD) 293T cells, using Anti-Tollip Antibody (A307074) at 1:500 dilution. The secondary antibody was Goat Anti-Rabbit IgG H&L Antibody (HRP) at 1:10,000 dilution. Lysates/proteins were present at 25 μ g per lane. The blocking buffer used was 3% non-fat dry milk in TBST. Detection was with a ECL Basic Kit. Exposure time: 3S.