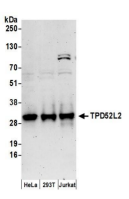
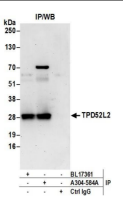
TPD52L2 Antibody

Rabbit Polyclonal Antigen Affinity Purifi Catalog No. A304- Lot No. A304-		Protein ID GenelD	NP_003279.2 7165	BETHYL LABORATORIES, INC
APPLICATIONS	WB, IP			
SPECIES REACTIVITY Human				
PRESUMED REACTIVITY	Based on 100% sequence identity, this antibody is predicted to react with Orangutan			
AMOUNT	100 µl			
CONCENTRATION	1000 μg/ml			
STORAGE/SHELF LIFE	2 - 8° C / 1 year from date of receipt			
PHYSICAL STATE	Liquid			
BUFFER	Tris-citrate/phosphate buffer, pH 7 to 8 containing 0.09% Sodium Azide			
ISOTYPE	IgG			
ORIGIN	USA			
PRODUCTION PROCEDURES	Antibody was affinity purified using an epitope specific to TPD52L2 immobilized on solid support.			
	The epitope recognized by A304–584A maps to a region between residue 156 to 206 of human Tumor Protein D52-like 2 using the numbering given in entry NP_003279.2 (GeneID 7165).			
	Antibody concentration was determined by extinction coefficient: absorbance at 280 nm of 1.4 equals 1.0 mg of IgG.			
APPLICATIONS	Centrifuge tube to remove product from lid. Optimal working dilutions should be determined experimentally by the investigator. Prepare working dilution immediately before use.			
	Western Blot	1:1	,000 - 1:5,000	
	Immunoprecipita	tion 2 –	10 µg/mg lysate	
APPLICATION NOTES	Western blot of immunoprecipitates performed using Normal Pig Serum (Cat. No. S100–020), Goat anti-Rabbit Light Chain HRP Conjugate (Cat. No. A120–113P) and 4–20% SDS–PAGE (link to IP-western blot protocol in Additional Info section below).			
	Western blot of lysates performed using standard western blot reagents and 4-20% SDS-PAGE.			
ADDITIONAL INFO	https://www.bethyl.com/product/A304–584A			
			S, a current list of citations s://www.bethyl.com/conte	, and other product specific information. nt/protocol_IP_WB

This document certifies that this product has met all of the quality control standards defined by Bethyl Laboratories, Inc. Eric McIntush, PhD | Chief Scientific Officer Date: June 21, 2019

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Detection of human TPD52L2 by western blot. Samples: Whole cell lysate (50 μ g) from HeLa, HEK293T, and Jurkat cells prepared using NETN lysis buffer. Antibodies: Affinity purified rabbit anti-TPD52L2 antibody A304-584A (lot A304-584A-1) used for WB at 0.4 μ g/ml. Detection: Chemiluminescence with an exposure time of 3 minutes. Detection of human TPD52L2 by western blot of immunoprecipitates. *Samples:* Whole cell lysate (0.5 or 1.0 mg per IP reaction; 20% of IP loaded) from 293T cells prepared using NETN lysis buffer. *Antibodies:* Affinity purified rabbit anti-TPD52L2 antibody A304-584A (lot A304-584A-1) used for IP at 6 µg per reaction. TPD52L2 was also immunoprecipitated by rabbit anti-TPD52L2 antibody BL17361. For blotting immunoprecipitated TPD52L2, A304-584A was used at 1 µg/ml. *Detection:* Chemiluminescence with an exposure time of 3 minutes.