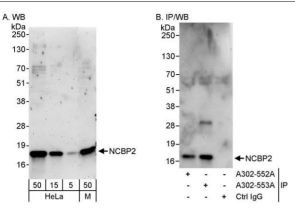
NCBP2 Antibody

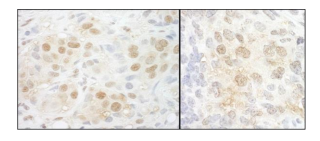
Rabbit Polyclonal			Protein ID	NP_031388.2			
Antigen Affinity Purified Catalog No. A302–553A			GenelD	22916			
Lot No.	A302-1		Geneid	22910		BETHYL LABORATORIES, INC	
APPLICATIONS		WB, IP, IHC					
SPECIES REACTIVITY		Human, Mouse					
PRESUMED REACTIVITY		Based on 100% sequence identity, this antibody is predicted to react with Bovine					
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 NCBP2 immobilized on solid support.					
Incolocites		The epitope recognized by A302-553A maps to a region between residue 106 to 156 of human nuclear cap binding protein subunit 2 using the numbering given in entry NP_031388.2 (GeneID 22916).					
		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	1:2	,000 - 1:10,000			
		Immunoprecipit	ation 5 –	15 µg/mg lysate			
		Immunohistoche		,000 – 1:5,000. Epitope ommended for FFPE tiss		e buffer pH 6.0 is	
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 4–12% SDS–PAGE (link to IP–western blot protocol in Additional Info section below).					
IHC HUMAN CONTROLS		Western blot of lysates performed using standard western blot reagents and 4–12% SDS-PAGE. Breast Carcinoma, Testicular Seminoma					
IHC MOUSE CONTROLS Terate		Teratoma	eratoma				
ADDITIONAL INFO		https://www.bethyl.com/product/A302-553A Use the link above to view SDS, a current list of citations, and other product specific information. IP-western blot protocol: https://www.bethyl.com/content/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 and mouse NCBP2 by western blot (h&m) and immunoprecipitation (h). Samples: Whole cell lysate from HeLa (5, 15 and 50 μ g for WB; 1 mg for IP, 20% of IP loaded) and mouse NIH 3T3 (M; 50 μ g) cells. Antibodies: Affinity purified rabbit anti-NCBP2 antibody A302-553A used for WB at 0.4 μ g/ml (A) and 1 μ g/ml (B) and used for IP at 10 μ g/mg lysate. NCBP2 was also immunoprecipitated by rabbit anti-NCBP2 antibody A302-552A, which recognizes an upstream epitope. Detection: Chemiluminescence with exposure times of 3 minutes (A) and 30 seconds (B).



Detection of human and mouse NCBP2 by

immunohistochemistry. *Sample:* FFPE section of human breast carcinoma (left) and mouse teratoma (right). *Antibody:* Affinity purified rabbit anti- NCBP2 (Cat. No. A302-553A Lot1) used at a dilution of 1:5,000 (0.2µg/ml) and 1:1,000 (1 µg/ml). *Detection:* DAB