

A Reliable Research Partner in Life Science and Medicine

Human THP Antibody Pair Set

Catalog No.E-KAB-0453ApplicationsELISASynonymsUromodulin;UMOD;ADMCKD2;FJHN;HNFJ1;MCKD2;THGP

Kit components & Storage

Title	Specifications	Storage
Human THP Capture Antibody	1 vial, 100 μ g	Store at -20°C for one year. Avoid
		freeze/thaw cycles.
Human THP Detection Antibody (Biotin)	1 vial, 50 μL	Store at -20°C for one year. Avoid
		freeze/thaw cycles.

Note: Centrifuge before opening to ensure complete recovery of vial contents.

Product Information

Items		Characteristic (E-KAB-0453)	
		Human THP Capture Antibody	Human THP Detection Antibody
			(Biotin)
Immunogen	Immunogen	Recombinant Human THP protien	Recombinant Human THP protien
Information	Swissprot	P07911	
Product details	Reactivity	Human	Human
	Host	Mouse	Sheep
	Conjugation	Unconjugated	Biotin
	Concentration	0.5 mg/mL	/
	Buffer	PBS with 0.04% Proclin 300; 50%	PBS with 0.04% Proclin 300; 1%
		glycerol; pH 7.5	protective protein; 50% glycerol; pH
			7.5
	Purify	Protein A or G	Antigen Affinity
	Specificity	Detects Human THP in ELISAs.	

For Research Use Only

Toll-free: 1-888-852-8623 Tel: 1-832-243-6086 Fax: 1-832-243-6017 Web: www.elabscience.com Email: techsupport@elabscience.com



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Applications

Human THP Sandwich ELISA Assay

	Recommended	Reagent	Images
	Concentration/Dilution		
ELISA	0.5-4 μg/mL	Human THP Capture	
Capture		Antibody	10
			b 1
			o priceal Density
ELISA	1:1000-1:10000	Human THP Detection	do Object
Detection		Antibody (Biotin)	0.01
			0.1 1 10 100 1000
			Human THP Concentration(ng/mL)

Note: This standard curve is only for demonstration purposes. A standard curve should be generated for each assay!

Background

The protein encoded by this gene is the most abundant protein in mammalian urine under physiological conditions. Its excretion in urine follows proteolytic cleavage of the ectodomain of its glycosyl phosphatidylinosital-anchored counterpart that is situated on the luminal cell surface of the loop of Henle. This protein may act as a constitutive inhibitor of calcium crystallization in renal fluids. Excretion of this protein in urine may provide defense against urinary tract infections caused by uropathogenic bacteria. Defects in this gene are associated with the renal disorders medullary cystic kidney disease-2 (MCKD2), glomerulocystic kidney disease with hyperuricemia and isosthenuria (GCKDHI), and familial juvenile hyperuricemic nephropathy (FJHN). Alternative splicing of this gene results in multiple transcript variants.

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