

Cytokeratin 1 (CK1) (ABT157R) Rabbit mAb (Ready to Use)

CatalogNo: YM7092R Recombinant 💦

Key Features

Host Species

Rabbit

Reactivity
• Human,

Applications
• IHC

IsotypeIgG1,Kappa

Recommended Dilution Ratios

Ready to use for IHC

Storage

Storage*	2°C to 8°C/1 year
Formulation	The prediluted ready-to-use antibody is diluted in phosphate buffer saline containing stabilizing protein and 0.05% Proclin 300

Basic Information

Clonality	Monoclonal
Clone Number	ABT157R

Immunogen Information

ImmunogenSynthesized peptide derived from human Cytokeratin 1 AA range:200-300SpecificityThis antibody detects endogenous levels of Cytokeratin 1

Target Information

Gene name KRT1 KRTA

Protein NameKeratin, type II cytoskeletal 1 (67 kDa cytokeratin) (Cytokeratin-1) (CK-1) (Hair alpha
protein) (Keratin-1) (K1) (Type-II keratin Kb1)

	Organism	Gene ID	UniProt ID
	Human	<u>3848;</u>	<u>P04264;</u>
Cellular Localization	Cytoplasmic, Membranous		
Tissue specificity	The source of this protein is neonatal terminally differentiating epidermis.	foreskin. The 67-kDa type II k	eratins are expressed in
Function	Disease:Defects in KRT1 are a cause (BCIE) [MIM:113800]; also known as a erythroderma ichthyosiformis congen disorder characterized by widespread that persist into adulthood. Histologic the lower spinous layer of the epidern blister formation diminish and hyperk of ichthyosis annular epidermolytic (A epidermolytic hyperkeratosis. AEI is a ichthyosiform erythroderma. Affected childhood and hyperkeratotic lichenif at later ages. The feature that disting annular polycyclic plaques with scale and can persist for several weeks or a palmoplantar keratoderma non-epide dermatological disorder characterized and follicular lesions.,Disease:Defects Macklin type (IHCM) [MIM:146590]. IF hyperkeratosis. Affected individuals n neck, and limbs with truncal erythem lips, ears, nipples and buttocks.,Disea keratoderma striate type 3 (SPPK3) [I striata III. SPPK3 is a dermatological of corneum and epidermal layers are th skin, and both hair and nails are norn as PKC and SRC via binding to integri kinase C (RACK1/GNB2L1).,miscelland microfibrillar keratin: I (acidic; 40-55 information:Keratin-1 entry,polymorp allele 1A and allele 1B with allelic free compared to allele 1A.,PTM:Undergoe (citrullination).,similarity:Belongs to t location:Located on plasma membrar cells.,subunit:Heterotetramer of two ta associated with keratin-10. Interacts with GNB2L1.,tissue specificity:The so	epidermolytic hyperkeratosis (ita of Brocq. BCIE is an autoso blistering and an ichthyotic e ally there is a diffuse epiderm nis. Within a few weeks from k eratoses develop.,Disease:De KEI) [MIM:607602]; also known skin disorder resembling bull individuals present with bullo ied plaques in the flexural are uishes AEI from BCIE is drama which coalesce to involve mo even months.,Disease:Defects rmolytic (NEPPK) [MIM:600962 by focal palmoplantar kerato is in KRT1 are the cause of icht ICM is a genodermatosis with nanifest congenital verrucous a, palmoplantar keratoderma ise:Defects in KRT1 are the ca MIM:607654]; also known as k lisorder affecting palm and so ickened. There is no involvem hal.,Function:May regulate the n beta-1 (ITB1) and the recept cous:There are two types of cy kDa) and II (neutral to basic; 5 hism:There are two size variar quencies of 0.61 and 0.39. Alle is deimination of some arginin he intermediate filament famil te of neuroblastoma NMB7 type I and two type II keratins. with ITGB1 in the presence of ource of this protein is neonata	EHK) or bullous omal dominant skin rythroderma at birth objytic degeneration in birth, erythroderma and fects in KRT1 are a cause a scyclic ichthyosis with ous congenital ous ichthyosis in early as and extensor surfaces atic episodes of flares of ost of the body surface in KRT1 are a cause of 2]. NEPKK is a oderma with oral, genital, thyosis hystrix Curth- severe verrucous black scale on the scalp, and keratoses on the suse of palmoplantar eratosis palmoplantar eratosis palmoplantar eactivity of kinases such tor of activated protein vtoskeletal and 66-70 kDa).,online nts of KRT1, termed ele 1B lacks 7 residues the residues ly.,subcellular

Validation Data



Human skin squamous cell carcinoma was stained with anti-Cytokeratin 1 (CK1) (ABT157R) rabbit mAb

Contact information

Orders:	order.cn@immunoway.com
Support:	support.cn@immunoway.com
Telephone:	400-8787-807(China)
Website:	http://www.immunoway.com.cn
Address:	2200 Ringwood Ave San Jose, CA 95131 USA



Please scan the QR code to access additional product information: **Cytokeratin 1 (CK1)** (ABT157R) Rabbit mAb (Ready to Use)

For Research Use Only. Not for Use in Diagnostic Procedures.

Antibody | ELISA Kits | Protein | Reagents