

Rabbit Anti-ABCA12 antibody

SL11906R

ABCA12
腺苷三磷酸结合盒转运体12抗体
ABC transporter A family member 12; ABC transporter ABCA.12; ABC12; ABCA12; ABCAC_HUMAN; AtABCA12; ATH16; ATP binding cassette 12; ATP binding cassette sub family A (ABC1) member 12; ATP binding cassette sub family A member 12; ATP binding cassette transporter 12; ATP-binding cassette 12; ATP-binding cassette sub-family A member 12; ATP-binding cassette transporter 12; Ichthyosis congenita II lamellar ichthyosis B; ICR2B; LI2; Putative ABC2 homolog 16; ABCAC_HUMAN.
Rabbit
Polyclonal
Human,Mouse,Rat,Dog,Pig,Rabbit,
ELISA=1:500-1000IHC-P=1:400-800IHC-F=1:400-800ICC=1:100-500IF=1:100- 500 (Paraffin sections need antigen repair) not yet tested in other applications. optimal dilutions/concentrations should be determined by the end user.
293kDa
The cell membrane
Lyophilized or Liquid
1mg/ml
KLH conjugated synthetic peptide derived from human ABCA12:2051-2200/2595
IgG
affinity purified by Protein A
0.01M TBS(pH7.4) with 1% BSA, 0.03% Proclin300 and 50% Glycerol.
Store at -20 °C for one year. Avoid repeated freeze/thaw cycles. The lyophilized antibody is stable at room temperature for at least one month and for greater than a year when kept at -20°C. When reconstituted in sterile pH 7.4 0.01M PBS or diluent of antibody the antibody is stable for at least two weeks at 2-4 °C.
PubMed
The ATP-binding cassette (ABC) transporters, or traffic ATPases, constitute an

expansive family of proteins accountable for the transport of a wide variety of substrates across cell membranes in both prokaryotic and eukaryotic cells. They also aid in the regulation of lipid transport and membrane trafficking. ABCA12 (ATP-Binding Cassette, Subfamily A, Member 12) contains two transmembrane (TM) domains, each with six membrane-spanning segments, and two nucleotide-binding domains (NBDs), which are located in the cytoplasm. ABCA12 is expressed in normal human keratinocytes (RT-PCR reveals expression in placenta, testis, fetal brain, and skin) and is upregulated during keratinization. Immunoelectron microscopy reveals that the ABCA12 protein is located in lamellar granules in the upper epidermal keratinocytes of human skin. The ABCA12 gene, which synthesizes a 2,595-amino acid protein, may produce an alternative splice variant with an in-frame deletion leading to truncation of 79 amino acids.

Function:

Probable transporter involved in lipid homeostasis.

Subcellular Location: Membrane.

Tissue Specificity: Mainly expressed in the stomach, placenta, testis and fetal brain.

DISEASE:

Defects in ABCA12 are the cause of ichthyosis harlequin (HI) [MIM:242500]; also known as harlequin fetus. HI is a very severe skin disorder in which the neonate is born with a thick covering of armor-like scales. The skin dries out to form hard diamond-shaped plaques separated by fissures, resembling 'armor plating'. The normal facial features are severely affected, with distortion of the lips (eclabion), eyelids (ectropion), ears, and nostrils. Affected babies are often born prematurely and rarely survive the perinatal period.

Defects in ABCA12 are the cause of ichthyosis lamellar type 2 (LI2) [MIM:601277]; also known as ichthyosis congenita IIB (ICR2B). LI is a non-bullous ichthyosis, a skin disorder characterized by abnormal cornification of the epidermis. It is one the most severe forms of ichthyoses apparent at birth and persisting throughout life. LI patients are born encased in a tight, shiny, translucent covering called collodion membrane. Over the first weeks of life, the collodion membrane is gradually replaced by generalized large, dark brown, plate-like scales with minimal to no erythroderma. Tautness of facial skin commonly results in ectropion, eclabium and scarring alopecia of the scalp. Common complications are severe heat intolerance and recurrent ear infections.

Similarity:

Belongs to the ABC transporter superfamily. ABCA family. Contains 2 ABC transporter domains.

SWISS:

Q86UK0
Gene ID:
26154
Database links:
Entrez Gene: 26154Human
Entrez Gene: 74591Mouse
Entrez Gene: 301482Rat
<u>Omim: 607800</u> Human
SwissProt: Q86UK0Human
Unigene: 134585Human
SwissProt: Q86UK0Human Unigene: 134585Human Important Note:
Important Note:
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