## **TSPEAR Antibody**



## PACO15728

Reactivity:

## **Product Information**

Size: Protein Background:

50ul This gene encodes a protein that contains a N-terminal thrombospondin-type laminin

G domain and several tandem arranged epilepsy-associated repeats (EARs). A mutation in this gene is the cause of autosomal recessive deafness-98. Alternate splicing results

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Human in multiple transcript variants.

Source: Gene ID:

Rabbit TSPEAR

Isotype: Uniprot

IgG Q8WU66

Applications: Synonyms:

ELISA, IHC thrombospondin-type laminin G domain and EAR repeats

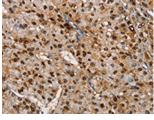
Recommended dilutions: Immunogen:

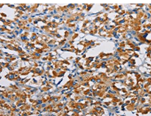
ELISA:1:1000-1:5000, IHC:1:50-1:200 Fusion protein of human TSPEAR.

Storage:

-20° C, pH7.4 PBS, 0.05% NaN3, 40% Glycerol

## **Product Images**





The image on the left is immunohistochemistry of paraffin-embedded Human liver cancer tissue using PACO15728(TSPEAR Antibody) at dilution 1/50, on the right is treated with fusion protein. (Original magnification: x—200).

The image on the left is immunohistochemistry of paraffin-embedded Human thyroid cancer tissue using PACO15728(TSPEAR Antibody) at dilution 1/50, on the right is treated with fusion protein. (Original magnification: x—200).