For Research Use Only

androgen receptor Polyclonal antibody

Antibodies | ELISA kits | Proteins WWW.ptglab.com

Catalog Number:22576-1-AP

2 Publications

Catalog Number:

Basic Information

- 22576-1-AP Size: 400 µ g/ml Source: Rabbit Isotype: IgG Immunogen Catalog Number: AG17385
- GenBank Accession Number: BC132975 GeneID (NCBI): 367 UNIPROT ID: P10275 Full Name: androgen receptor Calculated MW: 914 aa, 99 kDa Observed MW: 110 kDa

Purification Method: Antigen affinity purification

Applications

Tested Applications: ELISA Cited Applications: WB, IF, IHC Species Specificity: human, monkey, pig Cited Species: human

Background Information

Androgen receptor (AR) also konwn as Dihydrotestosterone receptor (DHTR), Nuclear receptor subfamily 3 group C member 4 (NR3C4). It is one of steriod hormoen receptors, which are ligand-activated transcription factors that regulate eukaryotic gene expression and affect cellular proliferation and differentiation in target tissues. Transcription factor activity is modulated by bound coactivator and corepressor proteins. Transcription activation is down-regulated by NR0B2. Activated, but not phosphorylated, by HIPK3 and ZIPK/DAPK3.Defects in AR are the cause of androgen insensitivity syndrome (AIS). Affected males have female external genitalia, female breast development, blind vagina, absent uterus and female adnexa, and abdominal or inguinal testes, despite a normal 46,XY karvotype Defects in AR are the cause of spinal and bulbar muscular atrophy X-linked type 1 (SMAX1). In SMAX1 patients the number of Gln ranges from 38 to 62. Longer expansions result in earlier onset and more severe clinical manifestations of the disease. Defects in AR may play a role in metastatic prostate cancer. The mutated receptor stimulates prostate growth and metastases development despite of androgen ablation. This treatment can reduce primary and metastatic lesions probably by inducing apoptosis of tumor cells when they express the wildtype receptor. Defects in AR are the cause of androgen insensitivity syndrome partial (PAIS). PAIS is characterized by hypospadias, hypogonadism, gynecomastia, genital ambiguity, normal XY karyotype, and a pedigree pattern consistent with X-linked recessive inheritance. Some patients present azoospermia or severe oligospermia without other clinical manifestations. This antibody is a rabbit polyclonal antibody. It can specifically recognize the 110kd AR protein.

Notable Publications

| Author | Pubmed ID | Journal | Application |
|-------------|-----------|-------------|-------------|
| Kejun Cheng | 29904891 | Med Oncol | WB |
| Qingfu Deng | 30664187 | Mol Med Rep | WB,IHC,IF |

Storage

Storage:

Store at -20°C. Stable for one year after shipment. Storage Buffer: PBS with 0.02% sodium azide and 50% glycerol pH 7.3. Aliquoting is unnecessary for -20°C storage

This product is exclusively available under Proteintech Group brand and is not available to purchase from any other manufacturer. Selected Validation Data