bs-1472R

[Primary Antibody]

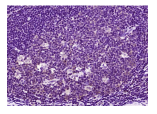
IFI16 Rabbit pAb



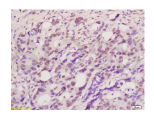
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– DATASHEET –		400-901-9800
Host: Rabbit	Isotype: IgG	Applications: IHC-P (1:100-500)
Clonality: Polyclonal		IHC-F (1:100-500) IF (1:100-500)
GenelD: 3428	SWISS: Q16666	Doo stivity Userson
Target: IFI16		Reactivity: Human
Immunogen: KLH conjugated 201-350/785.	synthetic peptide derived from human IFI16:	
Purification: affinity purified by Protein A		Predicted MW.: ^{86 kDa}
Concentration: 1mg/ml		
Storage: 0.01M TBS (pH7.4) with 1% BSA, 0.02% Proclin300 and 50% Glycerol. Shipped at 4°C. Store at -20°C for one year. Avoid repeated freeze/thaw cycles.		Subcellular Location: ^{Cytoplasm} ,Nucleus
HIN 200 human a comtaining regu transcriptional re has three isotype mRNA alternativ threonine residu is restricted to th epithelial cells. II lineage is tightly and proliferation regulation of pro assumed that IFI differentiation. In	nducible 16 (IFI16) protein belongs to a family of and mouse proteins. IFI16 is a nuclear protein latory domains such as DNA binding domain, egulatory domain and DAPIN/PAAD domain. IFI16 es A, B, and C (85-95 kDa), which arise as a result of e splicing. All are phosphorylated on serine and es and can homo and heterodimerize. Expression ie nuclei of hematopoietic cells, fibroblasts and F116 expression in hematopoietic cells of myeloid regulated and highly induced in the differentiation of the cell. Due to its localization in the nucleus, tein expression, and ability to bind DNA, it is 16 has a role in transcription regulation of cell n addition, it was found that IFI16 can act as a epressor and is involved in regulation and in cancer cells.	

– VALIDATION IMAGES



Paraformaldehyde-fixed, paraffin embedded (human tonsil); Antigen retrieval by boiling in sodium citrate buffer (pH6.0) for 15min; Block endogenous peroxidase by 3% hydrogen peroxide for 20 minutes; Blocking buffer (normal goat serum) at 37°C for 30min; Antibody incubation with (IFI16) Polyclonal Antibody, Unconjugated (bs-1472R) at 1:200 overnight at 4°C, followed by operating according to SP Kit(Rabbit) (sp-0023) instructionsand DAB staining.



Tissue/cell: human colon carcinoma; 4% Paraformaldehyde-fixed and paraffinembedded; Antigen retrieval: citrate buffer (0.01M, pH 6.0), Boiling bathing for 15min; Block endogenous peroxidase by 3% Hydrogen peroxide for 30min; Blocking buffer (normal goat serum,C-0005) at 37°C for 20 min; Incubation: Anti-IFI16 Polyclonal Antibody, Unconjugated(bs-1472R) 1:200, overnight at 4°C, followed by conjugation to the secondary antibody(SP-0023) and DAB(C-0010) staining

- SELECTED CITATIONS -

• [IF=1.69] Li et al. Histopathology of melanosis coli and determination of its associated genes by comparative analysis of expression microarrays. (2015) Mol.Med.Re. 12:5807-15 IF,WB ;Human. 26238215