

## HNF1α (Phospho Ser247) rabbit pAb

| Catalog No :            | YP1354  |
|-------------------------|---|
| Reactivity :            | Human;Rat;Mouse;  |
| Applications :          | WB  |
| Target :                | HNF1A   |
| Fields :                | >>Maturity onset diabetes of the young  |
| Gene Name :             | HNF1A TCF1  |
| Protein Name :          | HNF1a (Ser247)  |
| Human Gene Id :         | 6927  |
| Human Swiss Prot        | P20823  |
| No :<br>Mouse Gene Id : | 21405   |
| Mouse Swiss Prot        | P22361  |
| No :<br>Rat Gene Id :   | 24817   |
| Rat Swiss Prot No :     | P15257  |
| Immunogen :             | Synthesized phosho peptide around human HNF1a (Ser247)                          |
| Specificity :           | This antibody detects endogenous levels of Human HNF1a (phospho-Ser247)         |
| Formulation :           | Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.         |
| Source :                | Polyclonal, Rabbit,IgG  |
| Dilution :              | WB 1:1000-2000  |
| Purification :          | The antibody was affinity-purified from rabbit serum by affinity-chromatography |



|                        | using specific immunogen.  |
|------------------------|--|
|                        |  |
| <b>Concentration :</b> | 1 mg/ml  |
| Storage Stability :    | -15°C to -25°C/1 year(Do not lower than -25°C)   |
| Observed Band :        | 69kD   |
| Cell Pathway :         | Maturity onset diabetes of the young;  |
| Background :           | The protein encoded by this gene is a transcription factor required for the expression of several liver-specific genes. The encoded protein functions as a homodimer and binds to the inverted palindrome 5'-GTTAATNATTAAC-3'. Defects in this gene are a cause of maturity onset diabetes of the young type 3 (MODY3) and also can result in the appearance of hepatic adenomas. Alternative splicing results in multiple transcript variants encoding different isoforms. [provided by RefSeq, Apr 2015],  |
| Function :             | disease:Defects in HNF1A are a cause of susceptibility to insulin-dependent<br>diabetes mellitus (IDDM) [MIM:222100].,disease:Defects in HNF1A are the cause<br>of maturity onset diabetes of the young type 3 (MODY3) [MIM:600496]; also<br>symbolized MODY-3. MODY [MIM:606391] is a form of diabetes characterized<br>by an autosomal dominant mode of inheritance, age of onset of 25 years or<br>younger and a primary defect in insulin secretion. The clinical phenotype of<br>MODY3 is characterized by severe insulin secretory defects, and by major<br>hyperglycemia associated with microvascular complications.,disease:Defects in<br>HNF1A may predispose to hepatic adenomas [MIM:142330]. Hepatic adenomas<br>are benign tumors at risk of malignant transformation. Bi-allelic inactivation of<br>HNF1A, whether sporadic or associated with MODY3, may be an early step in the<br>developmant of some hepatocellular carcinomas.,function:Required |
| Subcellular            | Nucleus .  |
| Location :             |  |
| Expression :           | Liver.   |

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