# **Product Name: Transferrin(10G3)Mouse Monoclonal**

**Antibody** 

Catalog #: AMM19206



### Summary

Transferrin(10G3)Mouse Monoclonal Antibody **Production Name** 

Description Mouse Monoclonal Antibody

Host Mouse **Application** WB,IHC, Reactivity Human

#### **Performance**

Conjugation Unconjugated Modification Unmodified

Isotype IgG

**Clonality** Monoclonal **Form** Liquid

Store at 4°C short term. Aliquot and store at -20°C long term. Avoid freeze/thaw Storage

cycles.

Buffer Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% New type preservative N.

**Purification** Affinity purification

#### **Immunogen**

**Gene Name** TF

**Alternative Names** TF; Serotransferrin; Transferrin; Beta-1 metal-binding globulin; Siderophilin

Gene ID 7018.0

P02787.Protein SwissProt ID

# **Application**

**Dilution Ratio** WB 1:2000-5000, IHC 1:100-200.

**Molecular Weight** 77kD

# **Background**

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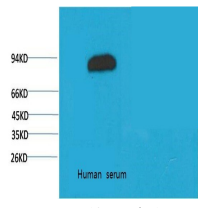
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transferrin(TF) Homo sapiens This gene encodes a glycoprotein with an approximate molecular weight of 76.5 kDa. It is thought to have been created as a result of an ancient gene duplication event that led to generation of homologous C and N-terminal domains each of which binds one ion of ferric iron. The function of this protein is to transport iron from the intestine, reticuloendothelial system, and liver parenchymal cells to all proliferating cells in the body. This protein may also have a physiologic role as granulocyte/pollen-binding protein (GPBP) involved in the removal of certain organic matter and allergens from serum. [provided by RefSeq, Sep 2009], disease:Defects in TF are the cause of atransferrinemia [MIM:209300]. Atransferrinemia is rare autosomal recessive disorder characterized by iron overload and hypochromic anemia., function:Transferrins are iron binding transport proteins which can bind two Fe(3+) ions in association with the binding of an anion, usually bicarbonate. It is responsible for the transport of iron from sites of absorption and heme degradation to those of storage and utilization. Serum transferrin may also have a further role in stimulating cell proliferation., online information:Transferrin entry, polymorphism:Different polymorphic variants of transferrin are known. The sequence shown is the predominant electrophoretic variant (C1 or TF\*C1)., similarity:Belongs to the transferrin family., similarity:Contains 2 transferrin-like domains., subunit:Monomer., tissue specificity:Expressed by the liver and secreted in plasma.,

#### Research Area

### **Image Data**



Western blot analysis of Human Serum with Transferrin Mouse mAb diluted at 1:2,000.

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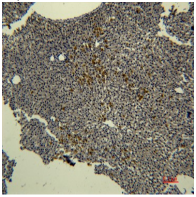
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Immunohistochemical analysis of paraffin-embedded human-liver using antibody diluted at 1:50.



Immunohistochemical analysis of paraffin-embedded rat-liver using antibody diluted at 1:50.

## Note

For research use only.

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