

# GC Antibody(Center) (Ascites)

Mouse Monoclonal Antibody (Mab)
Catalog # AM2180a

## **Specification**

# GC Antibody(Center) (Ascites) - Product Information

Application WB,E
Primary Accession P04062

Other Accession Q70KH2, Q2KHZ8, NP 000148.2

Reactivity
Predicted
Bovine, Pig
Host
Clonality
Human
Bovine, Pig
Mouse
Monoclonal

Isotype IgM
Calculated MW 59716
Antigen Region 337-365

# GC Antibody(Center) (Ascites) - Additional Information

#### **Gene ID 2629**

### **Other Names**

Glucosylceramidase, Acid beta-glucosidase, Alglucerase, Beta-glucocerebrosidase, Beta-GC, D-glucosyl-N-acylsphingosine glucohydrolase, Imiglucerase, GBA, GC, GLUC

# Target/Specificity

This GC antibody is generated from mice immunized with a KLH conjugated synthetic peptide between 337-365 amino acids from the Central region of human GC.

#### **Dilution**

WB~~1:100~1600

#### **Format**

Mouse monoclonal antibody supplied in crude ascites with 0.09% (W/V) sodium azide.

#### **Storage**

Maintain refrigerated at 2-8°C for up to 2 weeks. For long term storage store at -20°C in small aliquots to prevent freeze-thaw cycles.

# **Precautions**

GC Antibody(Center) (Ascites) is for research use only and not for use in diagnostic or therapeutic procedures.

## GC Antibody(Center) (Ascites) - Protein Information

Name GBA1 (HGNC:4177)

Synonyms GBA, GC, GLUC



Function Glucosylceramidase that catalyzes, within the lysosomal compartment, the hydrolysis of glucosylceramides/GlcCers (such as beta- D-glucosyl-(11')-N-acylsphing-4-enine) into free ceramides (such as N-acylsphing-4-enine) and glucose (PubMed:15916907, PubMed:24211208, PubMed:32144204, PubMed:9201993). Plays a central role in the degradation of complex lipids and the turnover of cellular membranes (PubMed: 27378698). Through the production of ceramides, participates in the PKC-activated salvage pathway of ceramide formation (PubMed: 19279011). Catalyzes the glucosylation of cholesterol, through a transglucosylation reaction where glucose is transferred from GlcCer to cholesterol (PubMed: 24211208, PubMed: <u>26724485</u>, PubMed: <u>32144204</u>). GlcCer containing mono-unsaturated fatty acids (such as beta-D- glucosyl-N-(9Z-octadecenoyl)-sphing-4-enine) are preferred as glucose donors for cholesterol glucosylation when compared with GlcCer containing same chain length of saturated fatty acids (such as beta-D- glucosyl-N-octadecanoyl-sphing-4-enine) (PubMed: 24211208). Under specific conditions, may alternatively catalyze the reverse reaction, transferring glucose from cholesteryl 3-beta-D-glucoside to ceramide (Probable) (PubMed: 26724485). Can also hydrolyze cholesteryl 3-beta-D- glucoside producing glucose and cholesterol (PubMed: 24211208, PubMed: 26724485). Catalyzes the hydrolysis of galactosylceramides/GalCers (such as beta-D-galactosyl-(11')-N- acylsphing-4-enine), as well as the transfer of galactose between GalCers and cholesterol in vitro, but with lower activity than with GlcCers (PubMed: 32144204). Contrary to GlcCer and GalCer, xylosylceramide/XylCer (such as beta-D-xyosyl-(11')-N-acylsphing-4- enine) is not a good substrate for hydrolysis, however it is a good xylose donor for transxylosylation activity to form cholesteryl 3-beta- D-xyloside (PubMed: 33361282).

#### **Cellular Location**

Lysosome membrane; Peripheral membrane protein; Lumenal side. Note=Interaction with saposin-C promotes membrane association (PubMed:10781797). Targeting to lysosomes occurs through an alternative MPR-independent mechanism via SCARB2 (PubMed:18022370).

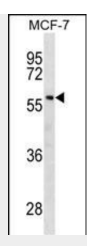
## GC Antibody(Center) (Ascites) - Protocols

Provided below are standard protocols that you may find useful for product applications.

- Western Blot
- Blocking Peptides
- Dot Blot
- <u>Immunohistochemistry</u>
- Immunofluorescence
- Immunoprecipitation
- Flow Cytomety
- Cell Culture

# GC Antibody(Center) (Ascites) - Images





GC Antibody(Center) (Cat. #AM2180a) western blot analysis in MCF-7 cell line lysates (35µg/lane). This demonstrates the GC antibody detected the GC protein (arrow).

# GC Antibody(Center) (Ascites) - Background

This gene encodes a lysosomal membrane protein that cleaves the beta-glucosidic linkage of glycosylceramide, an intermediate in glycolipid metabolism. Mutations in this gene cause Gaucher disease, a lysosomal storage disease characterized by an accumulation of glucocerebrosides. A related pseudogene is approximately 12 kb downstream of this gene on chromosome 1. Alternative splicing results in multiple transcript variants.

# GC Antibody(Center) (Ascites) - References

Dos Santos, A.V., et al. Neurosci. Lett. 485(2):121-124(2010) Bailey, S.D., et al. Diabetes Care 33(10):2250-2253(2010) Jeong, S.Y., et al. Blood Cells Mol. Dis. (2010) In press: Hu, F.Y., et al. Eur. J. Neurol. (2010) In press: Velayati, A., et al. Curr Neurol Neurosci Rep 10(3):190-198(2010)