

## HEXA DNAxPab

Catalog # H00003073-W01P      Size 200 ug

### Specification

<b>Product Description</b>	Rabbit polyclonal antibody raised against a full-length human HEXA DNA using DNAx™ Immune technology.
<b>Technology</b>	<a href="#">DNAx™ Immune</a>
<b>Immunogen</b>	Full-length human DNA
<b>Sequence</b>	MTSSRLWFSLLLAAAFAGRATALWPWPQNFQTSDQRYVLYPNNFQFYDVSSAAQPGCSVLD AFQRYRDLLFGSGSWPRPYLTGKRHTLEKNVLVVSVTPGCNQLPTLESVENYTLTINDDQCCLLS ETVWGALRGLETFSQLVWKSAEGTFFINKTEIEDFPRFPHRGLLLDTSRHYPPLSSILDLDVMAYN KLNVFHWHLVDDPSFPYESFTPELMRKGSYNPVTHIYTAQDVKEVIEYARLRGIRVLAEDTPGH TLSWGPGPGLLTPCYSGSEPSGTFGPVNPSLNNTYEFMSTFFLEVSSVFPDFYLHLGGDEVDF CWKSNPEIQDFMRKKKGFGEDFKQLESFYIQTLLDIVSYGKGYVVWQEVDNKVKIQPDIIQVWR EDIPVNMYMKELELVTKAGFRALLSAPWYLNRISYGPDWKDFYVVEPLAFEGTPEQKALVIGGEAC MWGEYVDNTNLVPRWLPRAGAVAERLWSNKLTSDLTFAYERLSHFRCELLRRGVQAQPLNVGF CEQEFEQT
<b>Host</b>	Rabbit
<b>Reactivity</b>	Human
<b>Purification</b>	Protein A
<b>Quality Control Testing</b>	Antibody reactive against mammalian transfected lysate.
<b>Storage Buffer</b>	In 1x PBS, pH 7.4
<b>Storage Instruction</b>	Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

### Applications

- Western Blot (Transfected lysate)

[Protocol Download](#)

- Immunofluorescence (Transfected cell)
- Flow Cytometry (Transfected cell)

## Gene Info — HEXA

Entrez GenelD	<a href="#">3073</a>
GeneBank Accession#	<a href="#">NM_000520.2</a>
Protein Accession#	<a href="#">NP_000511.1</a>
Gene Name	HEXA
Gene Alias	MGC99608, TSD
Gene Description	hexosaminidase A (alpha polypeptide)
Omim ID	<a href="#">272800 606869</a>
Gene Ontology	<a href="#">Hyperlink</a>
Gene Summary	This gene encodes the alpha subunit of the lysosomal enzyme beta-hexosaminidase that, together with the cofactor GM2 activator protein, catalyzes the degradation of the ganglioside GM2, and other molecules containing terminal N-acetyl hexosamines. Beta-hexosaminidase is composed of two subunits, alpha and beta, which are encoded by separate genes. Both beta-hexosaminidase alpha and beta subunits are members of family 20 of glycosyl hydrolases. Mutations in the alpha or beta subunit genes lead to an accumulation of GM2 ganglioside in neurons and neurodegenerative disorders termed the GM2 gangliosidoses. Alpha subunit gene mutations lead to Tay-Sachs disease (GM2-gangliosidosis type I). [provided by RefSeq]
Other Designations	GM2 gangliosidosis N-acetyl-beta-glucosaminidase Tay Sachs disease beta-N-acetylhexosaminidase hexosaminidase A

## Pathway

- [Amino sugar and nucleotide sugar metabolism](#)
- [Glycosaminoglycan degradation](#)
- [Glycosphingolipid biosynthesis - ganglio series](#)
- [Glycosphingolipid biosynthesis - globo series](#)
- [Lysosome](#)

- [Metabolic pathways](#)
- [Other glycan degradation](#)

## Disease

- [Genetic Predisposition to Disease](#)
- [Sandhoff Disease](#)
- [Tay-Sachs disease](#)