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Datasheet for ABIN1402387  
**anti-MID1 antibody (AA 200-250) (Biotin)**

### Overview

Quantity:	100 µL
Target:	MID1
Binding Specificity:	AA 200-250
Reactivity:	Human, Mouse, Rat
Host:	Rabbit
Clonality:	Polyclonal
Conjugate:	This MID1 antibody is conjugated to Biotin
Application:	Western Blotting (WB), Immunohistochemistry (Paraffin-embedded Sections) (IHC (p))

### Product Details

Immunogen:	KLH conjugated synthetic peptide derived from human Midline-1/RNF59
Isotype:	IgG
Cross-Reactivity:	Human, Mouse, Rat
Purification:	Purified by Protein A.

### Target Details

Target:	MID1
Alternative Name:	Midline-1 ( <a href="#">MID1 Products</a> )
Background:	Synonyms: BBBG1, Finger on X and Y mouse homolog of antibody, FXY, GBBB1, MID-1, Mid1, Midin, Midline 1 Opitz/BBB syndrome, Midline 1, Midline 1 ring finger, Midline 1 RING finger

## Target Details

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protein, Midline-1, Midline1, OGS1, OSX, Putative transcription factor XPRF, RING finger protein 59, RNF59, TRI18, TRI18\_HUMAN, TRIM18, Tripartite motif containing protein 18, Tripartite motif protein TRIM18, Tripartite motif-containing protein 18, XPRF, Zinc finger X and Y antibody, ZNFXY.

Background: Midline-1 (Tripartite motif-containing protein 18, Putative transcription factor XPRF, RING finger protein 59) is a 667 amino acid protein encoded by the human gene MID1. Midline-1 belongs to the TRIM/RBCC family and contains two B box-type zinc fingers, one B30.2/SPRY domain, one COS domain, one fibronectin type-III domain and one RING-type zinc finger. Midline-1 is believed to have E3 ubiquitin ligase activity which targets the catalytic subunit of protein phosphatase 2 for degradation. It is a cytoplasmic protein found as a homodimer or heterodimer with Midline-2. It also interacts with IGBP1 (Lymphocyte signaling protein A4). Defects in MID1 are the cause of Opitz syndrome type I (OS-I). OS-I is an X-linked recessive disorder characterized by hypertelorism, genital-urinary defects such as hypospadias in males and splayed labia in females, lip-palate-laryngotracheal clefts, imperforate anus, developmental delay and congenital heart defects. OS-I mutations produce proteins with a decreased affinity for microtubules.

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Gene ID: 4281

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UniProt: [O15344](#)

## Application Details

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Application Notes: WB 1:300-5000  
IHC-P 1:200-400

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Restrictions: For Research Use only

## Handling

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Format: Liquid

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Concentration: 1 µg/µL

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Buffer: Aqueous buffered solution containing 0.01M TBS (pH 7.4) with 1 % BSA, 0.03 % Proclin300 and 50 % Glycerol.

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Preservative: ProClin

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Precaution of Use: This product contains ProClin: a POISONOUS AND HAZARDOUS SUBSTANCE, which should be handled by trained staff only.

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## Handling

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Storage: -20 °C

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Storage Comment: Store at -20°C for 12 months.

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Expiry Date: 12 months