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Diagnostik & molekulare Diagnostik



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Lieferung & Zahlungsart

siehe unsere [Liefer- und Versandbedingungen](#)

Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

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NDN (Human) Matched Antibody Pair

Catalog # : H00004692-AP11

規格 : [1 Set]

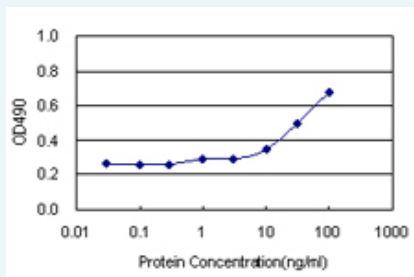
[List All](#)

Specification

Product Description: This antibody pair set comes with matched antibody pair to detect and quantify protein level of human NDN.

Reactivity: Human

Quality Control Testing: Standard curve using recombinant protein (H00004692-P01) as an analyte.



Sandwich ELISA detection sensitivity ranging from 3 ng/ml to 100 ng/ml.

Supplied Product: Antibody pair set content:
 1. Capture antibody: rabbit MaxPab® affinity purified polyclonal anti-NDN (100 ug)
 2. Detection antibody: mouse monoclonal anti-NDN, IgG2a Kappa (20 ug)
 *Reagents are sufficient for at least 1-2 x 96 well plates using recommended protocols.

Storage Instruction: Store reagents of the antibody pair set at -20°C or lower. Please aliquot to avoid repeated freeze thaw cycle. Reagents should be returned to -20°C storage immediately after use.

MSDS:  [Download](#)

Applications

ELISA Pair (Recombinant protein)

 [Protocol Download](#)

Gene Information

Entrez GeneID: [4692](#)

Gene Name: NDN

Gene Alias: HsT16328, PWCR

Gene Description: necdin homolog (mouse)

Omim ID: [176270](#), [602117](#)

Gene Ontology: [Hyperlink](#)

Gene Summary: This intronless gene is located in the Prader-Willi syndrome deletion region. It is an imprinted gene and is expressed exclusively from the paternal allele. Studies in mouse suggest that the protein encoded by this gene may suppress growth in postmitotic neurons. [provided by RefSeq]

Other OTTHUMP00000159437,necdin

Designations:

Related Disease

Attention Deficit Disorder with Hyperactivity Autistic Disorder Body Weight NARP Obesity Obesity, Morbid Prader-Willi Syndrome Prader-Willi syndrome Sleep Apnea, Obstructive

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