



# SZABO SCANDIC

Part of Europa Biosite

## Produktinformation



Forschungsprodukte & Biochemikalien



Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



Laborgeräte & Service

Weitere Information auf den folgenden Seiten!  
See the following pages for more information!



### Lieferung & Zahlungsart

siehe unsere [Liefer- und Versandbedingungen](#)

### Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

### SZABO-SCANDIC HandelsgmbH

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## MFN2 FISH Probe

Catalog # : FA0012

規格 : [ 200 uL ]

List All

### Specification

|                             |   |
|-----------------------------|---|
| <b>Product Description:</b> | Made to order FISH probes for identification of gene amplification using Fluorescent In Situ Hybridization Technique. ( <a href="#">Technology</a> )  |
| <b>Supplied Product:</b>    | DAPI Counterstain (1500 ng/mL ) 250 uL  |
| <b>Storage Instruction:</b> | Store at 4°C in the dark.   |
| <b>Origin:</b>              | Human   |
| <b>Source:</b>              | Genomic DNA   |
| <b>Notice:</b>              | We <b>strongly recommend</b> the customer to use FFPE FISH PreTreatment Kit 1 (Catalog #: <a href="#">KA2375</a> or <a href="#">KA2691</a> ) for the pretreatment of Formalin-Fixed Paraffin-Embedded (FFPE) tissue sections. |
| <b>Regulation Status:</b>   | For research use only (RUO)   |

### Application Image

Fluorescent In Situ Hybridization (Cell)

### Applications

Fluorescent In Situ Hybridization (Cell)

 [Protocol Download](#)

### Gene Information

**Entrez GeneID:** [9927](#)

**Gene Name:** MFN2

**Gene Alias:** CMT2A,CMT2A2,CPRP1,HSG,KIAA0214,MARF

**Gene Description:** mitofusin 2

**Omim ID:** [601152](#), [608507](#), [609260](#)

**Gene Ontology:** [Hyperlink](#)

**Gene Summary:** This gene encodes a mitochondrial membrane protein that participates in mitochondrial fusion and contributes to the maintenance and operation of the mitochondrial network. This protein is involved in the regulation of vascular smooth muscle cell proliferation, and it may play a role in the pathophysiology of obesity. Mutations in this gene cause Charcot-Marie-Tooth disease type 2A2, and hereditary motor and sensory neuropathy VI, which are both disorders of the peripheral nervous system. Defects in this gene have also been associated with early-onset stroke. Two transcript variants encoding the same protein have been identified. [provided by RefSeq]

**Other** OTTHUMP00000002509,hyperplasia suppressor,mitochondrial

**Designations:** assembly regulatory factor,mitofusin-2,transmembrane GTPase MFN2

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#### **Related Disease**

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[Charcot-Marie-Tooth Disease](#) [Genetic Predisposition to Disease](#) [Glaucoma, Open-Angle](#)  
[Hereditary Sensory and Motor Neuropathy](#)

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