



# SZABO SCANDIC

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## Produktinformation



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Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



Laborgeräte & Service

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### Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

### SZABO-SCANDIC HandelsgmbH

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## ATN1 Pre-design Chimera RNAi

Catalog # : H00001822-R01

規格 : [ 10 nmol ] [ 20 nmol ]

List All

### Specification

**Product Description:** Homo sapiens atrophin 1 (ATN1), transcript variant 1, mRNA.

**Reactivity:** Human

**Supplied Product:** DEPC water

**Target Refseq:** NM\_001007026

**Storage Instruction:** Store at -20°C, do not exceed 4 - 5 freeze-thaw cycles to ensure product integrity.

**Note:** Position of the Chimera RNAi.



### Application Image

RNAi Knockdown

### Publication Reference

- [dsCheck: highly sensitive off-target search software for double-stranded RNA-mediated RNA interference.](#)  
Naito Y, Yamada T, Matsumiya T, Ui-Tei K, Saigo K, Morishita S. *Nucleic Acids Res.* 2005 Jul 1;33(Web Server issue):W589-91.
- [Functional dissection of siRNA sequence by systematic DNA substitution: modified siRNA with a DNA seed arm is a powerful tool for mammalian gene silencing with significantly reduced off-target effect.](#)  
Ui-Tei K, Naito Y, Zenno S, Nishi K, Yamato K, Takahashi F, Juni A, Saigo K. *Nucleic Acids Res.* 2008 Apr;36(7):2136-51. Epub 2008 Feb 11.
- [Guidelines for the selection of highly effective siRNA sequences for mammalian and chick RNA interference.](#)  
Ui-Tei K, Naito Y, Takahashi F, Haraguchi T, Ohki-Hamazaki H, Juni A, Ueda R, Saigo K. *Nucleic Acids Res.* 2004 Feb 9;32(3):936-48. Print 2004.
- [siDirect: highly effective, target-specific siRNA design software for mammalian RNA interference.](#)  
Naito Y, Yamada T, Ui-Tei K, Morishita S, Saigo K. *Nucleic Acids Res.* 2004 Jul 1;32(Web Server issue):W124-9.

### Applications

RNAi Knockdown

### Gene Information

**Entrez GeneID:** [1822](#)

**Gene Name:** ATN1

**Gene Alias:** B37,D12S755E,DRPLA,NOD

**Gene** atrophin 1

**Description:**

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**Omim ID:** [125370](#), [607462](#)

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

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**Gene Summary:** Dentatorubral pallidoluysian atrophy is a rare neurodegenerative disorder characterized by cerebellar ataxia, myoclonic epilepsy, choreoathetosis, and dementia. The disorder is related to the expansion of a trinucleotide repeat within this gene. The encoded protein includes a serine repeat and a region of alternating acidic and basic amino acids, as well as the variable glutamine repeat. Alternative splicing results in two transcripts variants that encode the same protein.  
[provided by RefSeq]

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**Other** atrophin-1

**Designations:**

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**Related Disease**

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[Epilepsy](#) [Fragile X Syndrome](#) [Fragile X syndrome](#) [Friedreich Ataxia](#) [Friedreich's ataxia](#)  
[Genetic Predisposition to Disease](#) [Genomic Instability](#) [Huntington Disease](#)  
[Huntington disease](#) [Muscular Atrophy, Spinal](#) [Myoclonic Cerebellar Dyssynergia](#)  
[Myoclonic Epilepsies, Progressive](#) [Myotonic Dystrophy](#) [Myotonic dystrophy](#)  
[Parkinson Disease](#) [Parkinson disease](#) [Schizophrenia](#) [Schizophrenia](#)  
[Spinal muscular atrophy](#)

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