# атуюю вета Protein



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## Product Name

Amyloid Beta Protein

## Description

Human Synthetic Amyloid Beta Peptide 1-42 (HFIP treated)

## Applications

WB, In vivo Assay, In vitro Assay

#### Concentration

N/A - dried peptide film

#### Conjugates

No tag

### Nature

Synthetic

#### Species

Human

# **Expression System**

N/A

# Amino Acid Sequence

DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGVVIA

#### Purity

>95%

#### **Other Resources**

Protocol for re-suspension of Amyloid Beta Peptide 1-42

# **Protein Length**

42 amino acids

# **Field Of Use**

Not for use in humans. Not for use in diagnostics or therapeutics. For in vitro research use only.

# **Properties**

#### **Storage Buffer**

Dry powder. See "Other Resources" for re-suspension instructions/protocol.

## Storage Temperature

-80°C

#### Shipping Temperature

Blue Ice or 4°C

# Purification

N/A

## **Cite This Product**

Human Synthetic Amyloid Beta Peptide (StressMarq Biosciences Inc., Victoria BC CANADA, Catalog # SPR-485)

#### **Certificate Of Analysis**

Certified >95% pure using mass spec and HPLC.

# **Biological Description**

#### **Alternative Names**

Abeta Protein, Abeta peptide, Amyloid beta peptide, Beta amyloid peptide, amyloid beta precursor protein peptide, APP

# **Research Areas**

Alzheimer's Disease, Amyloid, Neurodegeneration, Neuroscience

#### **Cellular Localization**

Cell membrane, Intracellular Vesicles

# Gene ID 351 Swiss Prot 905067

#### **Scientific Background**

Our amyloid beta peptide 1-42 (Aβ42) is produced synthetically and treated with 1,1,1,3,3,3-Hexafluoro-2-propanol (HFIP) prior to drying which breaks down pre-formed fibrils and monomerizes the peptide, as previously published (1,2). Upon resuspension in DMSO/dH2O, our Aβ42 presents as a monomeric peptide without fibrils when observed under TEM, AFM and on a Western Blot with an anti-amyloid beta antibody. In contrast to AB42 oligomer and fibril constructs, our Aβ42 monomers were not toxic to primary rat cortical neurons. In the brain, amyloid beta peptide (Aβ) is generated by protease cleavage of amyloid precursor protein (APP), which aggregates into oligomers, protofibrils, fibrils and ultimately plaques in neurodegenerative diseases. The accumulation of Aβ plaques in the brain is considered a hallmark of Alzheimer's disease (AD), and most of the drugs tested for AD in the past 20 years have targeted amyloid beta accumulation (3). Soluble Aβ oligomers generated in vitro were toxic to PC12 cells (2) and SH-SY5Y cells (5). Aβ was demonstrated to interact with tauopathies to affect neurodegeneration in AD patients (6) and accumulations of Aβ were shown to be associated with lower survival rates in Parkinson's disease patients with dementia (7).

#### References

- 1. Stine et al. 2003. JBC. 278(13):11612-22. doi: 10.1074/jbc.M210207200
- 2. Chromy et al. 2003. Biochemistry. 42:12749-12760. doi: 10.1021/bi030029q
- 3. Panza et al. 2019. Nat Rev Neurol. 15:73-88 https://doi.org/10.1038/s41582-018-0116-6
- 4. Shankar et al. 2008. Nat Med. 14(8):837-842. doi: 10.1038/nm1782
- 5. Kayed et al. 2003. Science. 300(5618): 486-489. doi: 10.1126/science.1079469
- 6. Want et al. 2016. JAMA Neurol. 73(9):1070-7. doi: 10.1001/jamaneurol.2016.2078
- 7. Kotzbauer et al. 2012. Arch Neurol. 69(10): 1326-1331. doi: 10.1001/archneurol.2012.1608

# Product Images



TEM of amyloid beta 1-42 monomers (SPR-485). Negative stain transmission electron microscopy images acquired at 80 Kv on carbon coated 400 mesh copper grids using phosphotungstic acid and uranyl acetate stain. Scale bar = 100 nm.



AFM of amyloid beta 1-42 monomers (SPR-485). Atomic force microscopy analysis of 1.0 mg/mL samples diluted to 0.1 mg/mL in dH2O, mounted on freshly cleaved mica, washed, dried and analyzed with tapping mode. Representative images are 2.5 x 2.5  $\mu$ m x-y with a z-range of 10 nm.



Western blot of amyloid beta 1-42 monomers (SPR-485, left), oligomers (SPR-488, middle) and fibrils (SPR-487, right) using antiamyloid beta 6E10 antibody. Amyloid beta constructs at 160 pmol were run on 4-12% Bis-Tris SDS-PAGE, transferred to nitrocellulose in the presence of 0.02% v/v Tween-20, and blotted with 1:1000 mouse 6E10 primary antibody (Biolegend). Oligomers observed under TEM/AFM show distinct dimer/trimer bands as well as a signal from ~37-75 kDa (middle). Fibrils observed under TEM/AFM show a signal greater than 100 kDa and a distinct signal in the stacking gel (right).

#### Amyloid Beta Peptide (SPR-485)



Amyloid beta 1-42 oligomers (SPR-488) and fibrils (SPR-487) show a dose-dependent toxicity to primary rat cortical neurons, but not monomers (SPR-485). Survival of rat primary cortical neurons 14 days after treatment with different concentrations of (A) monomers, (B) oligomers or (C) fibrils quantified by MAP2 positive neurons and expressed as a percentage of control. Fibrils and respective vehicle controls were initially sonicated in a Bioruptor. Test conditions were run in the same plate as untreated control and vehicle controls, which consisted of buffer without amyloid beta 1-42 protein. Data expressed as mean +/- s.e.m. (n=6). A global analysis of the data was performed using a one-way ANOVA followed by Dunnett's test; \*\* p<0.01 stats vs control; ## p<0.01, #### p<0.0001 stats vs vehicle control. § represents untreated control condition.



TEM of amyloid beta 1-42 monomers (SPR-485, left), oligomers (SPR-488, middle) and fibrils (SPR-487, right). Negative stain transmission electron microscopy images acquired at 80 Kv on carbon coated 400 mesh copper grids using phosphotungstic acid and uranyl acetate stain. Scale bar = 100 nm.



AFM of amyloid beta 1-42 monomers (SPR-485, left), oligomers (SPR-488, middle) and fibrils (SPR-487, right). Atomic force microscopy analysis of 1.0 mg/mL samples diluted to 0.1 mg/mL in dH2O, mounted on freshly cleaved mica, washed, dried and analyzed with tapping mode. Representative images are 2.5 x 2.5 µm x-y with a z-range of 10 nm.

# **Product Citations (0)**

Currently there are no citations for this product.

# Reviews

There are no reviews yet.

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