Spinal Muscular Atrophy (SMA)

Novartis and AveXis combine, aiming to transform the care of SMA, the #1 genetic cause of infant mortality¹

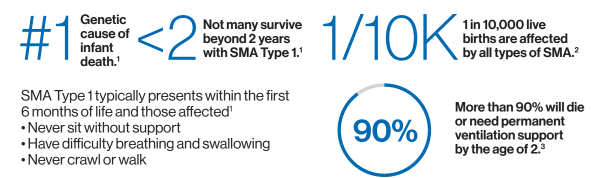
Novartis has completed the acquisition of the clinical-stage gene therapy company AveXis, Inc.

- The acquisition accelerates the Novartis strategy to pursue high-efficacy, first-in-class therapies and to deliver transformative innovation in areas of high unmet medical need.
- It provides another gene therapy platform, in addition to our CAR-T platform for cancer, and broadens our leadership in neuroscience.
- Together, we aim to advance a growing pipeline of gene therapies across therapeutic areas, including other SMA types, cancer, and blindness.

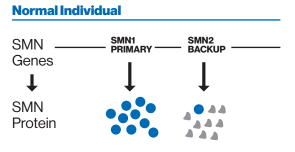
We are delighted to add the AveXis leading gene therapy technology to our company and to welcome our AveXis colleagues to Novartis. Together, we now have the potential to bring to children the first one-time gene-based treatment for the devastating disease, spinal muscular atrophy. - Vas Narasimhan, CEO Novartis

What is Spinal Muscular Atrophy (SMA)?

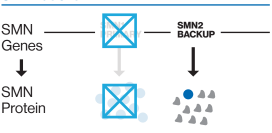
SMA is an inherited neurodegenerative disease caused by a defect in a single gene, the survival motor neuron (SMN1). It leads to a loss of motor neurons, resulting in progressive muscle weakness and paralysis.¹



Mechanism of disease and therapeutic strategies

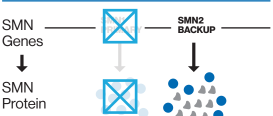


SMA Patient

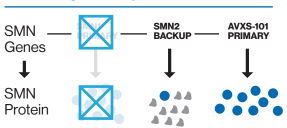


Therapeutic Strategies in SMA-Afflicted Individual





Introducing SMN1 expression



Time = neurons

Newborn screening could transform SMA care

Key Opportunity		Specialist referral	Geneticist / Genetic Counselor (varies)	
The Journey Begins	Symptoms	Diagnosis	Management & Treatment	The Journey Continues
Prenatal diagnosis Newborn screening	Pediatrician visit First symptoms	Neuromuscular Referral Genetic Testing	Diagnosis confirmed	Approved therap or Clinical trial enrollment

- All available treatments work better when started earlier; lost function unlikely to be regained.
- Clear progress in US Federal and State newborn screening for SMA. EU and other countries are also evaluating.

How does the gene therapy work?

The major steps of gene therapy are^{4,5}

Step 2 Step 1 Step 3 A modified noninfectious The virus enters neurons and The episome is transcribed and and nonpathogenic forms a DNA satelite called an translated to produce the missing adeno-associated Virus is episome that is delivered into SMN1 protein that is required for the the nucleus and does not health and survival of the neurons. administered to the patient integrate into the normal either intravenously or into DNA of the cell. the spinal cord. MODIFIED VIRAL DNA SMN1 GENE

Looking ahead

AveXis has expanded its clinical research to include additional types of SMA and other novel potential gene therapy treatment approaches.

VISION	ESTABLISH →	EXPAND	BUILD →
Leader in rare and life-threatening neurological genetic diseases	Foundational presence in SMA Type 1	Development of AVXS-101 into SMA Type 2	Pipeline of gene therapy treatments

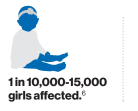
AveXis has built a team with exceptional depth of expertise and experience, a clinically proven gene delivery platform, manufacturing and R&D capabilities, while Novartis has been for 70 years a leader in Neuroscience, building on a global footprint and its extensive experience in bringing transformational medicines to the clinic stage. - Paul Hudson, CEO Novartis Pharmaceuticals

In addition to SMA, AveXis is also exploring novel gene therapy treatment approaches for Rett syndrome and a genetic form of amyotrophic lateral sclerosis (ALS) caused by mutations in the superoxide dismutase 1 (SOD1) gene.

RETT SYNDROME (RTT)

Rare, neurodevelopmental disorder predominantly affecting girls characterized by slowed growth,

loss of normal movement and coordination, and loss of communication skills.6





Typically results in significant disability that can include autistic-like behaviors, breathing irregularities, feeding and swallowing difficulties, growth retardation, and seizures.⁶

SOD1 AMYOTROPHIC LATERAL SCLEROSIS (ALS)

ALS (Lou Gehrig's Disease) is a progressive disease leading to significant disability, including muscle weakness, that results in loss of the ability to speak, eat, move, and eventually breathe.9



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Caused by mutations in the gene that produces the copper zinc superoxide dismutase 1 (SOD1) enzyme.7



40-70 Onset usually occurs in people between 40 and 70 years of age with a median age of diagnosis of 55.8,9



Genetic ALS affects 1,000-2,000 people in the U.S., 12%-20% caused by mutations in SOD1.7

vears

Typically results in death within 3-5 years of diagnosis.7,9

1. Farrar M., et al. Annals of Neurology. Emerging Therapies and Challenges in Spinal Muscular Atrophy. December 2016. 2. National Organization for Rare Disorders (NORD). Spinal Muscular Atrophy. http://rarediseases.org/rare-diseases/spinal-muscular-atrophy/ Accessed March 7, 2018. 3. Darras B.T., et al. Spinal Muscular Atrophy. Chapter 25 - Natural History of Spinal Muscular Atrophy. October 2016. 4. Global Genes. A Guide to Gene Therapy. https://globalgenes.org/ wp-content/uploads/2016/03/Guide-to-Gene Therapy_DIGITAL_spread-1.pdf Accessed April 19, 2018. 5. NIH. How does gene therapy work? https://ghr.nlm.nih.gov/primer/therapy/procedures Accessed April 19, 2018. 6. National Institute of Neurological Disorders and Stroke. Rett Syndrome Fact Sheet. https://www.ninds.nih.gov/Disorders/Patient-Caregiver-Education/Fact-Sheets/Rett-Syndrome-Fact-Sheet Accessed April 9, 2018. 7. National Institute of Neurological Disorders and Stroke. Amyotrophic Lateral Sclerosis (ALS) Fact Sheet. https://www.ninds.nih.gov/Disorders/Patient-Caregiver-Education/ Fact-Sheets/Amyotrophic-Lateral-Sclerosis-ALS-Fact-Sheet. Accessed April 25, 2018. 8. ALS Association. Facts You Should Know. http://www.alsa.org/ about-als/facts-you-should-know.html. Accessed April 25, 2018. 9. Chen S., et al. Molecular Neurodegeneration. Genetics of amyotrophic lateral sclerosis: an update. August 2013.



