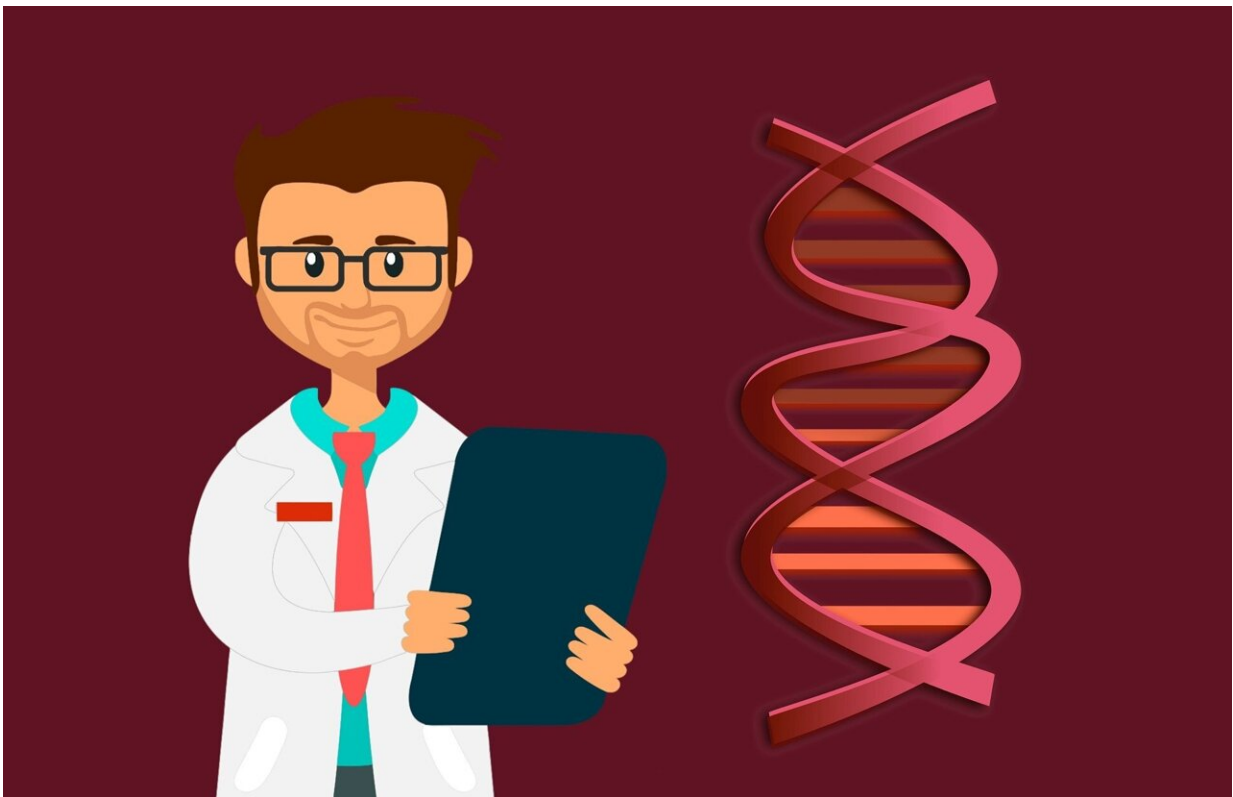


New components for antisense gene therapy show promise in treating spinal muscular atrophy

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Skoltech researchers and their colleagues from Russia and the UK investigated the safety and efficacy of new chemistry in antisense

oligonucleotides used to treat spinal muscular atrophy (SMA), a debilitating genetic disease. Their results may lead to the development of drugs with less toxicity and fewer injections needed thanks to prolonged action. The paper was published in the journal *Nucleic Acid Therapeutics*.

Antisense oligonucleotides are single stranded chemically modified fragments of DNA that target pre-messenger RNA, short bits of genetic information a ribosome reads to make a protein. Depending on how a particular antisense oligonucleotide works, the target mRNA can either be destroyed or undergo subtle changes in how it's spliced, i.e. how exons, the coding regions, are excluded or included in the final mRNA.

Antisense oligonucleotides are good at targeting so-called monogenic disorders, where the cause of the disease stems from one particular gene/protein. A common example of such disease is [spinal muscular atrophy](#) (SMA); people with this disease lose a functional protein encoded by gene SMN1, and even though the [human genome](#) contains a nearly identical copy, SMN2, the mRNAs transcribed from this gene lack just one necessary exon, which leads to a poorly functioning protein.

To help the cells successfully use SMN2 instead of SMN1, an antisense oligonucleotide can interfere with splicing of the precursor to mature mRNA that includes a particular exon. That is how nusinersen, the clinically approved antisense oligonucleotide against SMA marketed as Spinraza, works.

Timofei Zatsepin, Associate Professor at the Skoltech Center of Life Sciences, CLS Senior Research Scientist Olga Sergeeva and their colleagues studied alternatives to phosphorothioate groups in splice switching oligonucleotides. Previously, Dr. Stetsenko from Novosibirsk state university had developed oligonucleotides with methanesulfonyl (mesyl, μ) or 1-butanesulfonyl (busyl, β) phosphoramidate groups.

"Phosphorothioate is the key chemical modification of nucleic acids developed by Prof. Fritz Eckstein in late 1960s that is present in almost all oligonucleotide drugs approved so far. It improves stability, pharmacodynamics, and pharmacokinetics of oligonucleotides, but demonstrates significant toxicity that limits applications of [oligonucleotide](#) drugs. During the last 30 years, many alternatives were developed, but we do believe that mesyl phosphoramidates are superior to other phosphate mimics in therapeutic oligonucleotides," Zatspein said.

The new compounds are a bit unusual as their structure is quite confusing when one sees the formula for the first time. "You expect that such a bulk group should strongly interfere with all intracellular interactions. However, our colleagues from Novosibirsk, led by Dr. Dmitry Stetsenko, previously demonstrated that μ -oligonucleotides are much less toxic than PS oligonucleotides, while duplexes of μ -oligonucleotides with DNA are good substrates of RNase H—a key enzyme for mRNA degradation via antisense mechanism," Zatspein noted.

The researchers were inspired by these results and looked, among other things, for splice switching oligonucleotides with potentially prolonged action. Nusinersen is administered several times a year via an injection into the spinal canal, so fewer injections would improve the quality of life for patients with SMA.

The team was able to test the activity of novel [antisense oligonucleotides](#) in vitro in SMA patient-derived fibroblasts and in vivo in a neonatal mouse model of SMA. "In our study we found that μ -oligonucleotides were active in vitro, while in vivo the efficacy was lower in comparison to nusinersen at the same dose. As μ -oligos are more stable and less toxic in vivo than PS oligos, we propose that μ -oligos used in higher doses can provide the same efficacy together with more prolonged action—this

study is under development now," Zatsepin explained.

More information: Suzan M. Hammond et al, Mesyl Phosphoramidate Oligonucleotides as Potential Splice-Switching Agents: Impact of Backbone Structure on Activity and Intracellular Localization, *Nucleic Acid Therapeutics* (2021). [DOI: 10.1089/nat.2020.0860](https://doi.org/10.1089/nat.2020.0860)

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