

Shooting the messenger with single-stranded RNA gene silencing



New 'single-stranded RNA' gene silencing drugs could be safer & more effective to treat Huntington's disease

By Dr Nayana Lahiri on September 24, 2012

Edited by Dr Ed Wild

After huge leaps forward in recent years, we're edging ever closer to human trials of huntingtin lowering or 'gene silencing' as a potential treatment for Huntington's Disease. Newer, better and safer techniques are always welcome and the announcement of 'single-stranded RNA' silencing is causing quite a hubbub. What's it all about?

Gene silencing drugs work by telling cells not to make the huntingtin protein, the cause of damage in Huntington's disease. They do this by interfering with the cells' protein-making systems.

There are two main types of gene silencing drug: **anti-sense oligonucleotides** (ASOs) and **RNA interference** (RNAi). This article is about RNA interference.

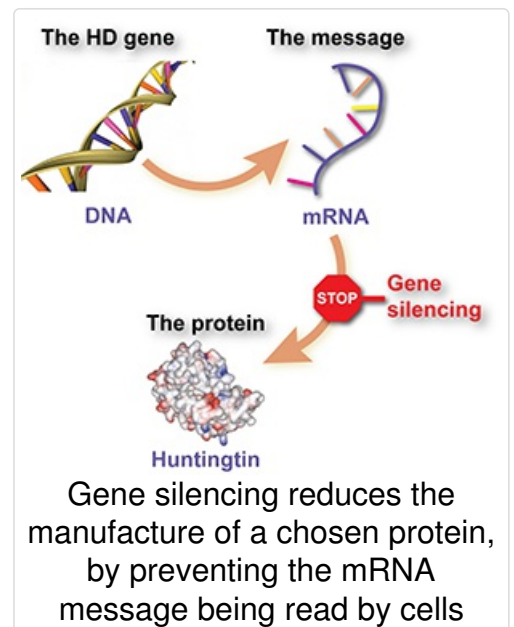
While pushing towards clinical trials as rapidly as possible, scientists have also been working on developing a new, and hopefully improved, RNA interference technique. To explain the differences from existing techniques we need to explain a bit about how genes are read to make proteins. Stick with us - it'll be worth it!

What are DNA and RNA?

DNA is the blueprint for making humans. It is a long molecule made up of stuck-together bits called **bases** which come in 4 'flavours' - C A G and T. These are the letters in which our genetic code is written.

The DNA 'double helix' is made up of two strands that line up side by side. Each strand is a string of bases, and the two strands are held together by chemical bonds between the bases on opposite strands. A pairs with T, and C pairs with G.

To get from DNA to a protein, a 'working copy' of the DNA needs to be made. The copy is called 'messenger' RNA, or mRNA. RNA is closely related to DNA but looks a little different.



mRNA is the template which tells a cell how to put a protein together. Whenever a cell makes any protein - including the huntingtin protein - it does so by reading the instructions in the mRNA that corresponds to that protein.

Interfering with huntingtin mRNA - or 'shooting the messenger' - prevents the huntingtin protein from being put together, and is the basis of gene silencing in RNA interference.

OK - back to gene silencing

So far, RNA-based gene silencing techniques have used double-stranded RNA molecules, called **siRNA**. Double-stranded RNA was needed until now, because single-stranded RNA is destroyed by the cells' own garbage-clearing mechanisms before it can do any silencing.

Once inside cells, the double-stranded RNA needs to separate into single strands to let it bind to the huntingtin mRNA message molecule. Then an enzyme in cells chews up the mRNA, so that the mutant protein never gets made.

We know that this way of doing things can dramatically decrease the amount of the mutant HD protein made in cells. What we are less sure about is whether the unwanted strand that separates off has any bad side effects on cells. There is a possibility that the body might launch an attack on the leftover strand and cause harm. Another possibility is that the leftover strand might bind to other mRNA, and prevent other important proteins from being made.

Double-stranded siRNA molecules don't spread very far within the brain, making treating large brain areas difficult.

A final difficulty of double-stranded siRNAs is that they need to be packaged up in a complicated way in order to get them to the right tissues for them to do their job.

There have been studies in mice and monkey animal models of Huntington's disease which showed that siRNAs are safe and effective, but scientists are a cautious bunch because we have to be really sure about safety before giving any treatments to precious humans. The last thing we want to do is make HD worse.

Single-stranded siRNA

One idea for reducing the risk of bad effects of RNA interference is to produce a single-stranded siRNA. But how could we overcome the stability problem - that annoying habit of cells of chopping up the single-stranded RNA drug?

After a lot of hard work, a group of researchers headed by David Corey in Dallas, in collaboration with ISIS pharmaceuticals have just announced they've finally cracked it. To give you an idea of the excitement this has caused, it was announced in not one, but two 'back-to-back' articles in the same edition of top journal Cell. Quite a scoop!

By making some chemical modifications to previous attempts, Corey's team have made a single-stranded siRNA, and were able to package it in a simple salt water solution. They successfully injected the single-stranded siRNA into the fluid spaces which surround the brain in

a mouse model of HD, and were able to show that it binds to huntingtin mRNA and prevents the HD protein from being made.

Win-win-win?

What's more, unlike double-stranded siRNA drugs that have been tested before, the effects of single-stranded siRNA spread throughout the brain rather than being confined to the small area near the injection. In these mice at least, single-stranded siRNA produced a win-win: it was stable and it spread further.

Not content with that, they took things further. By slightly altering the structure of their molecule, they could also create a single-stranded siRNA which only blocked production of the **mutant** HD protein and didn't prevent the normal version of the HD protein from being made. They did this by targeting the siRNA to the abnormally long 'CAG' stretch in the HD gene.

With a single-stranded siRNA we don't need to worry about what a second strand might be doing, and by targeting just the mutant huntingtin mRNA there's less concern about the possible effects of stopping production of the normal HD protein.

What next?




So this new method looks safe and effective in mouse models. Now we need to make sure it is safe and effective in other, larger animal models before even considering going into human trials. That's happening now!

Remaining questions

A few questions will need to be answered before single-stranded RNA interference can be tested in people.

First, there are some other genes with a stretch of CAGs in them. We don't yet know if introducing single-stranded siRNAs targeting those CAG stretches might inadvertently switch off other important genes.

Second, the delivery problem. How are we going to reach the important parts of the brain of Huntington's disease patients? Luckily, researchers in HD and other diseases are already working on this. A trial of a similar drug made from single-stranded DNA is being trialled in motor neuron disease (ALS) right now.

	Stable in cells?	Good silencing?	Spreads far?
 Double-stranded siRNA	✓	✓	✗
 Single-stranded siRNA	✗	✗	✓
 Modified Single-stranded siRNA	✓	✓	✓

Single-stranded siRNA drugs could combine the silencing power of double-stranded RNA with the ability of single-stranded molecules to spread throughout the brain.

Finally, how are we going to measure and monitor the effectiveness of treatment? In animal models, we can do this by looking at brain tissue and measure how much huntingtin protein is made. This is a lot more tricky to do in humans, but top scientists are working on it and we think we're ready to start trials in Huntington's disease patients.

More potential uses for single-stranded siRNA

A final sneak preview of another possible use of single-stranded siRNA. Researchers are also looking at using it in combination with stem cell research. Basically, they are working on taking skin cells from individuals with Huntington's disease, and turning those skin cells into neurons. Those neurons could then be treated with single-stranded siRNAs to reduce the levels of harmful HD protein before transplanting them back into the brain.

It'll be many years before the combination of single-stranded siRNA with patient-derived stem cells might be used as a treatment, but it's a neat idea and good to know all avenues are being explored. Meanwhile we expect single-stranded siRNA to progress fairly rapidly towards clinical trials.

The authors have no conflicts of interest to declare. For more information about our disclosure policy see our FAQ...

Glossary

motor neuron disease A progressive neurological disease in which motor (movement) neurons die. Also known as ALS or Lou Gehrig's disease.

huntingtin protein The protein produced by the HD gene.

RNA interference A type of gene silencing treatment in which specially designed RNA molecules are used to switch off a gene

gene silencing An approach to treating HD that uses targeted molecules to tell cells not to produce the harmful huntingtin protein

stem cells Cells that can divide into cells of different types

anti-sense the half of the DNA double-helix that is mostly used as a backup, but sometimes produces message molecules

neuron Brain cells that store and transmit information

siRNA A way of silencing genes using specially designed molecules of RNA – like DNA but made of only a single strand – that target the message molecules in cells and tell them not to make a certain protein

messenger RNA A message molecule, based on DNA, used by cells as the final set of instructions for making a protein.

ASOs A type of gene silencing treatment in which specially designed DNA molecules are used to switch off a gene

RNA the chemical, similar to DNA, that makes up the 'message' molecules that cells use as working copies of genes, when manufacturing proteins.

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