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Genetic Technology Restores Normal BMP

Signaling and Suppresses Bone Cell

Differentiation in a Human Stem Cell Model of

FOP: Proof-of-Principle for the Treatment of

FOP

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In the most direct and stealth-like approach yet to the treatment of FOP, scientists at the Center for Research in FOP and Related Disorders at the University of Pennsylvania have developed a new genetic approach using small sequences of ribonucleic acid (RNA) to specifically block the damaged copy of the FOP gene in cells while leaving the normal copy untouched.

Every human being has two copies of the ACVR1/ALK2 gene in every cell in their body. Individuals with FOP have one normal copy and one damaged copy of the gene in each cell – a dangerous occurrence that causes over-activity of ACVR1 and that tips the scales to renegade bone formation, the dreaded consequence of FOP.

Using inhibitory RNA designed and engineered to specifically silence the damaged copy of the gene rather than the normal copy (a process known as RNA interference or RNAi), the scientists restored the cellular function that was deranged by the FOP mutation by virtually ridding the cells of the damaged, mutant, and dangerous ACVR1/ALK2 mRNA. The cells were essentially left with only normal copies of ACVR1/ALK2 mRNA, thus adjusting the cellular activity to normal. The tremendous advantage of this approach is that ACVR1/ALK2 activity is not abolished in the cell, but brought to more normal levels, similar to that of cells without the FOP mutation.

Imagine identical twin pilots of a commercial jet airplane. The twins are both in the cockpit. For all outward appearances, they look identical and behave identically.

However, one of the twins is a good pilot and the other is a terrorist. What distinguishes

them is one letter in a simple instruction on how to fly the airplane, tattooed on each of their wrists.

And, as in all of genetics, the instructions are written in code words of three letters each. The good twin's tattoo says: "Now fly the jet." The tattoo of the terrorist twin says: "Now fry the jet." Just one letter difference, but the fate of the plane hangs on that one letter. The new approach developed by members of the Penn FOP research team use allele-specific (twin specific) inhibitory RNA designed to recognize the one letter difference in the tattoos, and eliminate the evil twin so that the good twin can fly the airplane without danger or interference. After the preemptory stealth attack by the allele-specific inhibitory RNA (that identifies and knocks-out the evil twin terrorist pilot) the plane is left with only one pilot, but a good responsible pilot. All is well!

In a landmark paper published in the Thursday October 20, 2011 online edition of **Gene Therapy** (a **Nature** journal), Dr. Josef Kaplan (lead author), Dr. Frederick Kaplan, and Dr. Eileen Shore (senior author), all from the FOP Research Laboratory of the Perelman School of Medicine at The University of Pennsylvania, describe in detail this new stealth-like proof-of-principle approach for treating FOP.

FOP is a rare genetic disorder of progressive heterotopic ossification for which there is presently no cure. FOP is caused by a recurrent activating mutant of ACVR1/ALK2, a bone morphogenetic protein (BMP) type 1 receptor that occurs in all classically affected individuals. Individuals who have FOP harbor one normal copy and one damaged copy

of the ACVR1/ALK2 gene in each cell. The FOP mutation (in one of the two copies of ACVR1/ALK2) increases BMP signaling to greater than normal levels and initiates the formation of a disabling second skeleton of heterotopic bone.

In their study, the authors generated specific inhibitory RNA duplexes capable of suppressing the expression of the mutant copy of the gene in connective tissue progenitor cells from FOP patients (while leaving the normal or good copy untouched) and importantly show that this approach decreased the elevated BMP signaling in FOP cells to levels observed in control cells. The cells used in the experiments were adult stem (or progenitor) cells obtained directly from discarded baby teeth of FOP patients and thus contained the exact combination of damaged and normal ACVR1/ALK2 receptors found in all classically affected FOP patients worldwide. The discarded teeth were obtained from FOP pediatric patients and normal controls in the ongoing “FOP Good Tooth Fairy Program.”

While the approach outlined in this landmark study provides proof-of-principle for the use of allele-specific inhibition of ACVR1/ALK2 in the treatment of FOP, the *in vivo* utility of this approach must be confirmed in mouse models of classic FOP prior to its consideration for human use. Additionally, other hurdles stand in the way of human application at the present time, most notably safe delivery of the small RNA duplexes to cells in the human body. “We have a long way to go,” acknowledge the investigators, “but we have taken a big first step.”

Improvements in RNAi design are advancing at a rapid rate and will enhance the stability, potency, and specificity of the inhibitory RNA allowing for long-term experiments both *in vitro* and *in vivo*. The new RNA interference approach developed by the Penn scientists can be applied to emerging mouse models of FOP providing hope for a novel therapeutic strategy to decrease and perhaps eliminate the catastrophic heterotopic bone formation in FOP patients.

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