New hope for rare 'stone man' disease, where flesh turns to bone

September 2 2015, by Dennis Thompson, Healthday Reporter

Abnormal bone formation in soft tissue, a process known as heterotopic ossification, in the hindlimb region and in the back of a mouse model of fibrodysplasia ossificans progressiva. Yellow pseudo-coloring has been applied to highlight the heterotopic bone. Credit: LiQin Xie and Nanditha Das

Researchers working with mice believe there's hope for patients with a rare genetic disorder that turns their muscle into bone, in essence immobilizing them in an extra skeleton.
The disease, fibrodysplasia ossificans progressiva (FOP), gradually replaces muscle, tendons, ligaments and other soft tissue with bone.

Called "stone man syndrome," this lethal disorder appears to be caused by mutations in a gene called ACVR1 that controls bone and muscle development, researchers report Sept. 2 in the journal Science Translational Medicine.

The mutations are thought to push the gene into overdrive when it's exposed to activin A, a signaling protein that normally inhibits the gene's function, said senior author Aris Economides, executive director at Regeneron Pharmaceuticals Inc., a biopharmaceutical company based in Tarrytown, N.Y.

"Imagine you are driving your car down the road and you come to a red light. You press on the brakes, and the car stops," Economides said. With this disorder, he said, "not only does your car not stop, actually the brakes are hot-wired to the accelerator."

Armed with that knowledge, researchers at Regeneron developed an antibody that blocks activin A from triggering the gene. The antibody successfully arrested bone formation in lab mice that had been genetically engineered to carry the disease mutation, they report.

Betsy Bogard, director of global research development for the International FOP Association, called this an "extraordinary" finding.

"We are very fortunate and grateful that not only did Regeneron make this basic science discovery, but that, as a biotechnology company with expertise in developing antibodies, they are in a position to act on it and answer the next questions about whether this could lead to a meaningful therapy," she said.
Fibrodysplasia ossificans progressiva affects only one in every 2 million people worldwide, according to the U.S. National Institutes of Health (NIH).

The International FOP Association says there are 800 confirmed cases across the globe, 285 of them in the United States. However, it is suspected that more people have the disease but have not been located, Bogard said.

Children with fibrodysplasia ossificans progressiva typically are born with malformed big toes. Growth of excess bone usually starts in the neck and shoulders, proceeding down the body and into the limbs, according to the NIH.

These patients may be unable to fully open their mouths, causing difficulty speaking and eating. They also may have trouble breathing properly because of excess bone growth around the rib cage. And as their joints fuse, they become less and less mobile.

Up to now, the only treatment for fibrodysplasia ossificans progressiva has been the steroid drug prednisone, which is used to help manage pain, Bogard said.

"It's not proven to do anything. It just helps them feel better through the pain of a flare-up," she said. "There's nothing else that people can do. This disease is on a march through their bodies, and it's not going to stop."

The newly developed antibody arrested bone formation in lab mice for up to six weeks, and did not have any "untoward, obvious side effects" in adult mice, Economides said.

The ACVR1 gene is crucial for healthy bone and muscle development as
early as in the womb, Economides said. As children age, the ACVR1 gene controls the growth of bone and muscle, including the gradual replacement of cartilage by bone that occurs in normal skeletal development from birth to young adulthood.

Activin A is often secreted by the immune system in response to injury and inflammation, researchers said, which may help explain why bone growth in patients with fibrodysplasia ossificans progressiva is often triggered by tissue swelling or inflammation.

Surgery makes the disease worse, Economides said. Trying to cut away the excess bone growth intensifies the patient's excess bone development. Even routine surgeries that don't involve bone can create inflammation and prompt a reaction, he said.

Economides could not say when the antibody would be ready for testing in humans. His company is currently performing preclinical safety testing.

Also, while the research looks promising in mice, results of animal studies often fail to be replicated in humans.


For more on fibrodysplasia ossificans progressiva, visit the [U.S. National Institutes of Health](https://www.ninds.nih.gov/Disorders/Disease/Fibrodysplasia_Ossificans_Progressiva/Fibrodysplasia_Ossificans_Progressiva_Actor/...)

Copyright © 2015 HealthDay. All rights reserved.