

Anti-inflammatory drug can prevent neuron loss in Parkinson's model

July 25 2014

An experimental anti-inflammatory drug can protect vulnerable neurons and reduce motor deficits in a rat model of Parkinson's disease, researchers at Emory University School of Medicine have shown.

The findings demonstrate that the drug, called XPro1595, can reach the brain at sufficient levels and have beneficial effects when administered by subcutaneous injection, like an insulin shot. Previous studies of XPro1595 in animals tested more invasive modes of delivery, such as direct injection into the brain.

"This is an important step forward for anti-inflammatory therapies for Parkinson's disease," says Malu Tansey, PhD, associate professor of physiology at Emory University School of Medicine. "Our results provide a compelling rationale for moving toward a clinical trial in early Parkinson's disease patients."

The new research on subcutaneous administration of XPro1595 was funded by the Michael J. Fox Foundation for Parkinson's Research (MJFF). XPro1595 is licensed by FPRT Bio, and is seeking funding for a clinical trial to test its efficacy in the early stages of Parkinson's disease.

"We are proud to have supported this work and glad to see positive preclinical results," said Marco Baptista, PhD, MJFF associate director of research programs. "A therapy that could slow Parkinson's progression would be a game changer for the millions living with this disease, and



this study is a step in that direction."

In addition, Tansey and Yoland Smith, PhD, from Yerkes National Primate Research Center, were awarded a grant this week from the Parkinson's Disease Foundation to test XPro1595 in a non-human primate model of Parkinson's.

Evidence has been piling up that inflammation is an important mechanism driving the progression of Parkinson's disease. XPro1595 targets tumor necrosis factor (TNF), a critical inflammatory signaling molecule, and is specific to the soluble form of TNF. This specificity would avoid compromising immunity to infections, a known side effect of existing anti-TNF drugs used to treat disorders such as rheumatoid arthritis.

"Inflammation is probably not the initiating event in Parkinson's disease, but it is important for the neurodegeneration that follows," Tansey says. "That's why we believe that an anti-inflammatory agent, such as one that counteracts soluble TNF, could substantially slow the progression of the disease."

Postdoctoral fellow Christopher Barnum, PhD and colleagues used a model of Parkinson's disease in rats in which the neurotoxin 6-hydroxydopamine (6-OHDA) is injected into only one side of the brain. This reproduces some aspects of Parkinson's disease: neurons that produce dopamine in the injected side of the brain die, leading to impaired movement on the opposite side of the body.

When XPro1595 is given to the animals 3 days after 6-OHDA injection, just 15 percent of the dopamine-producing neurons were lost five weeks later. That compares to controls in which 55 percent of the same neurons were lost. By reducing dopamine neuron loss with XPro1595, the researchers were also able to reduce motor impairment. In fact, the



degree of dopamine cell loss was highly correlated both with the degree of motor impairment and immune cell activation.

When XPro1595 is given two weeks after injection, 44 percent of the vulnerable neurons are still lost, suggesting that there is a limited window of opportunity to intervene.

"Recent clinical studies indicates there is a four or five year window between diagnosis of Parkinson's disease and the time when the maximum number of vulnerable neurons are lost," Dr. Tansey says. "If this is true, and if inflammation is playing a key role during this window, then we might be able to slow or halt the progression of Parkinson's with a treatment like XPro1595."

The results were published in the Journal of Parkinson's Disease.

Provided by Emory University

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