Study shows new way to treat stroke using an already FDA-approved drug
29 January 2020, by Gisele Galoustian

Jang-Yen (John) Wu, Ph.D., holds a bottle containing granulocyte colony-stimulating factor (GCSF), currently used to treat neutropenia due to chemotherapy and used is for patients who require bone marrow transplants. Results from the study showed that GCSF improved neurological deficits that occur in the first few days following cerebral ischemia and improved long-term behavioral outcomes while also stimulating a neural progenitor recovery response. Credit: Florida Atlantic University

Stroke is the third leading cause of death and disability in the United States. More than 87 percent are ischemic strokes, caused by obstruction of one or more cerebral arteries. With limited progress in developing treatments, there is a critical need for neuroprotective agents to effectively treat stroke.

A study from Florida Atlantic University’s Schmidt College of Medicine holds promise for a new way to treat stroke using an already FDA-approved drug—granulocyte colony-stimulating factor (GCSF). GCSF enhances blood cellular development and is currently used to treat neutropenia (low white blood cells) caused by chemotherapy and has successfully been used with very few side effects for patients who require bone marrow transplants to stimulate blood cell formation.

The study, published in the Journal of Biomedical Science, is the first to report on the neuroprotective effect of GCSF against autophagy and mitochondrial stress in vivo. The data support the hypothesis that GCSF is one of the few growth factors that can reduce infarction by decreasing endoplasmic reticulum (ER) and mitochondrial stress while improving behavioral performance.

Results showed that GCSF improved neurological deficits that occur in the first few days following cerebral ischemia and improved long-term behavioral outcomes while also stimulating a neural progenitor recovery response. Researchers tested behavioral performance on corner and locomotor tests, used as an indicator of brain injury.

Using a mouse model, researchers investigated the efficacy of GCSF beyond the typical four-hour thrombolytic therapy (tPA) clot-busting drug—the gold standard to treat stroke for global ischemia. They examined the pro-survival mechanisms of GCSF against apoptosis resulting from autophagy, mitochondrial stress and ER stress.

"In recent years, many studies including ours have shown that as an endogenous growth factor and immune system modulator factor, GCSF is beneficial in models of neurological disorders such as stroke and traumatic brain injury," said Jang-Yen (John) Wu, Ph.D., corresponding author, distinguished professor of biomedical science in FAU's Schmidt College of Medicine, and a member of the FAU Brain Institute (I-BRAIN). "Although the anti-apoptotic activity of GCSF is reported in global cerebral ischemia, this mechanism has not been fully explored."

Researchers used a mechanism-based therapeutic approach for stroke first to examine the connection
of mitochondrial, autophagy and ER stress inhibition in the protective action of GCSF and then to analyze relevant ER stress pathways in the bilateral common carotid artery occlusion (BCAO) model of stroke. They confirmed the neuroprotection of GCSF gene therapy in the BCAO mouse stroke model by a decrease of dynamin-related protein (DRP1), a marker of mitochondrial stress, in the frontal and middle brain of the GCSF treated group.

The initial dose of GCSF was administered 24 hours post-BCAO and then followed by a single application of the same dose for another three days for a total of four days of administration. Researchers examined behavior and used immunoblotting to analyze key proteins in ER stress, autophagy and mitochondrial stress-induced apoptosis. BCAO mice receiving GCSF protein showed significantly less asymmetric turning in the corner test than BCAO mice without GCSF. In the behavioral assays, GCSF elicited increased locomotor sensitization verified by greater activity in the locomotor activity test, demonstrating the neuroprotective properties of the drug.

"More than 15 million people worldwide suffer from stroke and our study provides new and important insights into GCSF induced protection as it relates to ER stress and mitochondrial stress activated apoptosis," said Howard Prentice, Ph.D., corresponding author, a professor of biomedical sciences in FAU's Schmidt College of Medicine, and a member of FAU's I-BRAIN. "Future research will need to focus on uncovering the complete mechanisms by which GCSF retains the ER and mitochondrial homeostasis."

Wu and Prentice have been developing GCSF as a therapeutic method to replenish new brain cells because of its ability to preserve the central nervous system, suppress cell death and at the same time elicit neurogenesis as well as angiogenesis. GCSF works the same way for other neurological diseases such as Parkinson's disease due to its neuroprotective properties.

More information: Mode of action of granulocyte-colony stimulating factor (G-CSF) as a novel therapy for stroke in a mouse model, https://jbiomedsci.biomedcentral...