

## Bone transcription factor controls nervous system gene expression, finds study

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Splits in compact myelin at the paranode area were observed in Sp7fl/fl;Olig2-Cre mice, where paranodal loops were growing between compact myelin layers. Credit: Benayahu Elbaz-eilon, Ph.D.

An established transcription factor known for bone formation also supports specialized cells in the central nervous system to promote brain tissue stiffness, findings that could inform new therapeutics for neuronal regeneration, according to a Northwestern Medicine <u>study</u> published in *Neuron*.

"We have uncovered the <u>molecular mechanisms</u> that control <u>extracellular</u> <u>matrix</u> production by oligodendrocytes that, interestingly, includes the same regulatory molecule that controls <u>bone formation</u>. Whenever this regulation is disrupted, oligodendrocytes do not function properly, which leads to a disruption in <u>neuronal function</u>," said Brian Popko, Ph.D., the William Frederick Windle Professor of Neurology and co-corresponding author of the study.

Oligodendrocytes are a specialized type of <u>central nervous system</u> (CNS) cells called <u>glial cells</u>, which support communication between neurons. The main function of oligodendrocytes is to produce myelin, which surrounds axons (<u>nerve fibers</u>) in an insulating sheath to help nerve signals properly travel throughout the CNS.

Recent work suggests that oligodendrocytes also provide metabolic support to neurons and support a wide range of brain functions as well as being involved in different neurodegenerative diseases. These cells also produce key components of the CNS's extracellular matrix (ECM), which is composed of large molecules that surrounds cells of all tissues and organs, providing structural support, as well as support for myriad cellular functions.



In the current study, the investigators aimed to characterize which <u>transcription factors</u> control ECM-related gene expression in oligodendrocytes.

"The role of oligodendrocytes in generation of the extracellular matrix and in establishing the bio-mechanical properties of the brain is overlooked and was currently unknown," said Benayahu Elbaz-eilon, Ph.D., assistant professor in the Ken and Ruth Davee Department of Neurology Division of Multiple Sclerosis and Neuroimmunology and cocorresponding author of the study.

Using a combination of atomic force microscopy and genetic and transcriptional study techniques, the investigators explored how oligodendrocytes contribute to CNS stiffness, which plays a fundamental role in nearly all biological processes in the brain, including development, regeneration and aging, according to Elbaz-eilon.

"Establishing accurate CNS biomechanics is an important but largely overlooked function of oligodendrocytes," Elbaz-eilon said.

Using these techniques, they discovered that Osterix, an established transcription factor in bone formation, binds to genes in oligodendrocytes essential for ECM function and gene expression related to nodes of Ranvier, or small gaps in myelin sheaths. Osterix has been shown to control the maturation of osteoblasts, which are essential for promoting bone formation and growth.

The findings suggest Osterix is required in oligodendrocytes for the development, maintenance and regeneration of myelin in the CNS and supports brain and bone tissue stiffness.

"Many neurological disorders include the loss of oligodendrocytes. This, of course, disrupts CNS myelination, but it also changes the ECM



composition of the brain. We are just beginning to understand the implications of an altered ECM in the functioning, repair and aging of the nervous system," Popko said.

The investigators are now interested in characterizing ECM composition and brain stiffness inside and near focal lesions in the brain. They also aim to identify new therapeutic targets that will help transform the ECM around brain lesions into a microenvironment that promotes neural and myelin regeneration.

"The <u>scientific community</u> is constantly looking for new ways to promote brain regeneration and formatting of new myelin, instead of the myelin that was lost in disease. The ability of the brain to regenerate depends in many ways on brain stiffness. Full understanding of the molecular pathways that control <u>brain</u> stiffness in health and disease will enable us to harness these findings for finding new therapeutic options," Elbaz-eilon said.

**More information:** Benayahu Elbaz et al, The bone transcription factor Osterix controls extracellular matrix- and node of Ranvier-related gene expression in oligodendrocytes, *Neuron* (2023). DOI: 10.1016/j.neuron.2023.10.008

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