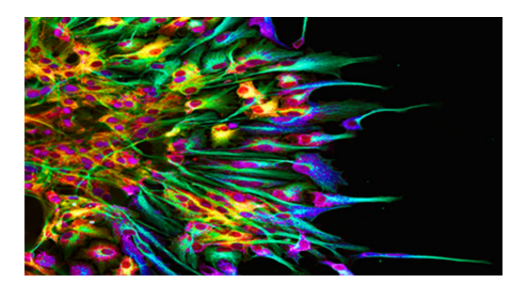


## Lack of protein Sp2 disrupts neuron creation in brain

January 14 2013



A rendered image of a primary neuronal stem cell culture in which cells were labeled with different fluorescently labeled proteins that differentiate between stem cells (orange/yellow) and their neuronal 'offspring' (blue/green/purple).

(Medical Xpress)—A protein known as Sp2 is key to the proper creation of neurons from stem cells, according to researchers at North Carolina State University. Understanding how this protein works could enable scientists to "program" stem cells for regeneration, which has implications for neural therapies.

Troy Ghashghaei and Jon Horowitz, both faculty in NC State's Department of Molecular Biomedical Sciences and researchers in the



Center for <u>Comparative Medicine</u> and Translational Research, wanted to know more about the function of Sp2, a cell cycle regulator that helps control how cells divide. Previous research from Horowitz had shown that too much Sp2 in skin-producing stem cells resulted in tumors in experimental mice. Excessive amounts of Sp2 prevented the stem cells from creating normal cell "offspring," or skin cells. Instead, the stem cells just kept producing more stem cells, which led to <u>tumor formation</u>.

"We believe that Sp2 must play a fundamental role in the lives of normal stem cells," Horowitz says. "Trouble ensues when the mechanisms that regulate its activity are overwhelmed due to its excess abundance."

Ghashghaei and his team – led by <u>doctoral candidate</u> Huixuan Liang – took the opposite approach. Using <u>genetic tools</u>, they got rid of Sp2 in certain <u>neural stem cells</u> in mice, specifically those that produce the major neurons of the brain's <u>cerebral cortex</u>. They found that a lack of Sp2 disrupted normal <u>cell formation</u> in these stem cells, and one important result was similar to Horowitz's: the abnormal stem cells were unable to produce normal cell "offspring," or neurons. Instead, the abnormal stem cells just created copies of themselves, which were also abnormal.

"It's interesting that both an overabundance of this protein and a total lack of it result in similar disruptions in how stem cells divide," Ghashghaei says. "So while this work confirms that Sp2 is absolutely necessary for stem cell function, a lot of questions still remain about what exactly it is regulating, and whether it is present in all stem cells or just a few. We also need to find out if Sp2 deletion or overabundance can produce brain tumors in our mice as in the skin.

"Finally, we are very interested in understanding how Sp2 regulates a very important decision a stem cell has to make: whether to produce more of itself or to produce offspring that can become neurons or <u>skin</u>



<u>cells</u>," Ghashghaei adds. "We hope to address those questions in our future research, because these cellular mechanisms have implications for cancer research, neurodevelopmental diseases and regenerative medicine."

The results appear online in Development.

NC State graduate students Guanxi Xiao, and Haifeng Yin, as well as Dr. Simon Hippenmeyer, a collaborator with the Ghashghaei lab from Austria's Institute of Science and Technology, contributed to the work. The work was funded by the National Institutes of Health and the American Federation for Aging Research.

**More information:** "Neural development is dependent on the function of specificity protein 2 in cell cycle progression" *Development*, 2013.

## Abstract

Faithful progression through the cell cycle is crucial to the maintenance and developmental potential of stem cells. Here, we demonstrate that neural stem cells (NSCs) and intermediate neural progenitor cells (NPCs) employ a zinc-finger transcription factor specificity protein 2 (Sp2) as a cell cycle regulator in two temporally and spatially distinct progenitor domains. Differential conditional deletion of Sp2 in early embryonic cerebral cortical progenitors, and perinatal olfactory bulb progenitors disrupted transitions through G1, G2 and M phases, whereas DNA synthesis appeared intact. Cell-autonomous function of Sp2 was identified by deletion of Sp2 using mosaic analysis with double markers, which clearly established that conditional Sp2-null NSCs and NPCs are M phase arrested in vivo. Importantly, conditional deletion of Sp2 led to a decline in the generation of NPCs and neurons in the developing and postnatal brains. Our findings implicate Sp2-dependent mechanisms as novel regulators of cell cycle progression, the absence of which disrupts neurogenesis in the embryonic and postnatal brain.



## Provided by North Carolina State University

Citation: Lack of protein Sp2 disrupts neuron creation in brain (2013, January 14) retrieved 1 May 2024 from <u>https://medicalxpress.com/news/2013-01-lack-protein-sp2-disrupts-neuron.html</u>

This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.