

Turning human stem cells into brain cells sheds light on neural development

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Medical researchers have manipulated human stem cells into producing types of brain cells known to play important roles in neurodevelopmental disorders such as epilepsy, schizophrenia and autism. The new model cell system allows neuroscientists to investigate normal brain development, as well as to identify specific disruptions in biological signals that may contribute to neuropsychiatric diseases.

Scientists from The Children's Hospital of Philadelphia and the Sloan-Kettering Institute for Cancer Research led a study team that described their research in the journal *Cell Stem Cell*, published online today.

The research harnesses human <u>embryonic stem cells</u> (hESCs), which differentiate into a broad range of different cell types. In the current study, the scientists directed the stem cells into becoming cortical interneurons—a class of <u>brain cells</u> that, by releasing the <u>neurotransmitter GABA</u>, controls electrical firing in <u>brain circuits</u>.

"Interneurons act like an orchestra conductor, directing other excitatory brain cells to fire in synchrony," said study co-leader Stewart A. Anderson, M.D., a research psychiatrist at The Children's Hospital of Philadelphia. "However, when interneurons malfunction, the synchrony is disrupted, and seizures or mental disorders can result."

Anderson and study co-leader Lorenz Studer, M.D., of the Center for Stem Cell Biology at Sloan-Kettering, derived interneurons in a laboratory model that simulates how neurons normally develop in the



human forebrain.

"Unlike, say, <u>liver diseases</u>, in which researchers can biopsy a section of a patient's liver, neuroscientists cannot biopsy a living patient's brain tissue," said Anderson. Hence it is important to produce a cell culture model of brain tissue for studying neurological diseases. Significantly, the human-derived cells in the current study also "wire up" in circuits with other types of brain cells taken from mice, when cultured together. Those interactions, Anderson added, allowed the study team to observe cell-to-cell signaling that occurs during forebrain development.

In ongoing studies, Anderson explained, he and colleagues are using their cell model to better define molecular events that occur during brain development. By selectively manipulating genes in the interneurons, the researchers seek to better understand how gene abnormalities may disrupt brain circuitry and give rise to particular diseases. Ultimately, those studies could help inform drug development by identifying molecules that could offer therapeutic targets for more effective treatments of neuropsychiatric diseases.

In addition, Anderson's laboratory is studying interneurons derived from stem cells made from skin samples of patients with chromosome 22q.11.2 deletion syndrome, a genetic disease which has long been studied at The Children's Hospital of Philadelphia. In this multisystem disorder, about one third of patients have autistic spectrum disorders, and a partially overlapping third of patients develop schizophrenia. Investigating the roles of genes and signaling pathways in their model cells may reveal specific genes that are crucial in those patients with this syndrome who have neurodevelopmental problems.

More information: Maroof et al, "Directed Differentiation and Functional Maturation of Cortical Interneurons from Human Embryonic Stem Cells," *Cell Stem Cell*, published online May 2, 2013.



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