Different origins discovered for medulloblastoma tumor subtypes

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Investigators have demonstrated for the first time that the most common malignant childhood brain tumor, medulloblastoma, is actually several different diseases, each arising from distinct cells destined to become different structures. The breakthrough is expected to dramatically alter the diagnosis and treatment of this major childhood cancer.

St. Jude Children's Research Hospital investigators led the international effort, which confirms that certain brain tumors and possibly other cancers regarded as the same disease; are in fact separate diseases with different origins. The finding supports the belief that the tumors will likely need different therapies. The study, which involved scientists at seven institutions in the United States, Japan and Great Britain, appears in the December 8 advance online edition of the journal Nature.

"Ten years ago medulloblastoma was regarded as a single disease, and all children with this brain tumor got the same treatment. This study shows clearly that there are distinct subtypes of this cancer that come from uniquely susceptible cell types in the brain that acquire specific mutations," said Richard Gilbertson, M.D., Ph.D., the study's senior author and a member of the St. Jude departments of Developmental Neurobiology and Oncology. "These findings will allow us to better model the heterogeneity we see in the clinic and to move toward designing directed therapies for each tumor subtype."

The research builds on work published in Nature earlier this year from Gilbertson and his colleagues. That research used the same method to show similar mechanisms at work in generating subtypes of ependymoma, the third most common brain tumor in children and the most common adult spinal tumor.

The approach in both investigations mapped gene expression to compare cells in the normal nervous system with cells in different subtypes of brain tumors. The goal was to identify the specific origins of different brain tumor subtypes.

The latest study focused on the wingless (WNT) and sonic hedgehog (SHH) subtypes of medulloblastoma, which together account for about 40 percent of the estimated 400 medulloblastoma tumors diagnosed annually in U.S. children and adolescents. The subtypes are named for the biochemical pathways abnormally activated in particular tumors.

Investigators used gene expression mapping to unmask a set of cells in the brainstem as the possible source of WNT-subtype medulloblastoma. The cells had not previously been linked to cancer. The cells also are located beneath and apart from the cerebellum. In the past, this brain structure was thought to be the source of all medulloblastomas.

Since mutations in the CTNNB1 genes occur specifically in patients with WNT-subtype medulloblastoma, researchers mutated the CTNNB1 gene in these same cells in the developing mouse brainstem. By the age of 6 months, about 20 percent of those mice developed medulloblastomas that mimicked the anatomy, histology and genetics of human WNT-subtype medulloblastoma.

Amar Gajjar, M.D., co-chair of the St. Jude Department of Oncology, said the mouse model will facilitate development of novel, less toxic therapies for children. He is co-author of the study.

Earlier research traced the origin of SHH-subtype medulloblastoma to a subset of cells known as granule neuron precursor cells destined to become part of the cerebellum. In this study, Gilbertson and colleagues show that over-expression of CTNNB1 in those precursor granule cells has no impact on the developing mouse cerebellum and does not lead to medulloblastoma.
The latest findings suggest WNT-subtype medulloblastoma begins in a subset of cells that become mossy fiber cells in the adult brainstem, but Gilbertson said additional research is needed to confirm the observation. The study also points to the loss of the p53 tumor suppressor gene and possibly the TULP4 tumor suppressor gene as fueling WNT-subtype tumor development.

Gilbertson led the team that in 2006 showed medulloblastoma could be categorized based on whether the WNT, the SHH or another biochemical pathway was abnormally activated. Researchers went on to demonstrate that the WNT and SHH subtypes also look different under the microscope, target patients of different ages and have different outcomes. The SHH subtype tends to arise in very young children. About 80 percent become long-term survivors. The WNT subtype usually strikes older adolescents and is considered curable in all cases.

Gilbertson said these latest findings indicate patients with WNT-subtype tumors might be candidates for less intensive therapy and thus less likely to suffer long-term treatment side effects. Medulloblastoma treatment includes surgery, radiation and chemotherapy.