Participants Needed for a Feasibility Safety Study

Participants needed for a feasibility safety study using the experimental surgical device ExAblate 4000 system in the management of benign, centrally located intracranial tumors that require clinical intervention in pediatric and young adult subjects.

Candidates include:

- Males and females ages 8 to 22 years with benign centrally located intracranial tumors that require clinical intervention who are scheduled for neurosurgery
- Subjects should be on a stable dose of all condition-related medications for 30 days prior to study entry
- Subjects and/or parent(s)/legal representative are able and willing to give informed consent and able to attend all study visits

To learn more or determine whether your child may be eligible to participate, please contact Tami Quintero via email at Tami.Quintero@mch.com or at 305-496-4188.