

Development of Pre-Clinical Assays for Quality Control of Cancer Imaging Agents

Chu, Ashley (School: Hathaway Brown School)

Glioblastoma is the most aggressive primary brain tumor, with an average survival of 14.6 months even with treatment. Prior research has shown that a protein, PTP μ , is cleaved in tumor tissue, and the abundant fragments allow them to serve as a tumor biomarker. Glioblastoma rapidly proliferates, making it difficult for surgeons to fully resect the tumor. Peptides derived from a short portion of PTP μ 's amino acid sequence are thought to bind to the fragments. By conjugating these peptides to fluorophores, surgeons can better identify what they need to resect. Using two in vitro assays, a bead aggregation assay and a fluorescent bead binding assay, we evaluated the effects of buffer conditions on peptide-protein interactions. Additionally, we sought to identify optimal conditions for peptide binding to PTP μ in hopes of improving clinical imaging agents for glioblastoma resection. With the bead aggregation assay, we found that particular types of media support the best aggregation. There is also a possible effect of pH on aggregation. When moving onto the fluorescent bead binding assay, preliminary research suggests that certain peptides have a strong affinity to PTP μ . Many more buffer conditions and peptides will be tested in future research to determine the conditions under which the peptides have the strongest binding to PTP μ , so they may be used as imaging agents.