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Immunostaining infiltrating spheroids as preparation for quantitative light-sheet imaging V.2

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Protocol status: Working

We use this protocol and it's working

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Abstract

Although various in vivo and in vitro models for studying glioblastoma cell invasion has progressed the field, there is still a need for optimized procedures. In particular to reveal key features of glioblastoma biology and infiltrating growth. In this protocol, we present an approach using indirect immunofluorescence in a 3D human xenograft glioblastoma spheroid model embedded in a naturally derived extracellular matrix

Attachments



[protocol.pdf](#)

415KB

