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PELSA protocol V.2

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Protocol status: Working

We use this protocol and it's working

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Abstract

This protocol details a step-by-step procedure of PELSA method, a modification-free approach that achieves unprecedented sensitivity in proteome-wide target identification and binding-region determination. Following this procedure, one can determine the binding proteins, binding regions, and binding affinities of the analyte ligand.

Attachments



[PELSA_protocol.pdf](#)

255KB

