

May 14, 2024

Quantification of IHC Using an Inverted Confocal Microscope and NIS-Elements Program-Killinger Lab 2024

DOI

dx.doi.org/10.17504/protocols.io.5qpvokrbzl4o/v1

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OPEN ACCESS



DOI: dx.doi.org/10.17504/protocols.io.5qpvokrbzl4o/v1

Protocol Citation: Solji Choi 2024. Quantification of IHC Using an Inverted Confocal Microscope and NIS-Elements Program-Killinger Lab 2024. protocols.io https://dx.doi.org/10.17504/protocols.io.5qpvokrbzl4o/v1

Manuscript citation:

Choi SG, Tittle T, Garcia-Prada D, Kordower JH, Melki R, Killinger BA. Alpha-synuclein aggregates are phosphatase resistant. bioRxiv [Preprint]. 2024 Apr 9:2023.11.20.567854. doi: 10.1101/2023.11.20.567854. PMID: 38645137; PMCID: PMC11030248.

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Protocol status: Working
We use this protocol and it's

working

Created: May 14, 2024

Last Modified: May 14, 2024

Protocol Integer ID: 99764



Funders Acknowledgement:

Michael J. Fox Foundation Grant ID: ASAP-021030 MIchael J. Fox Foundation Grant ID: ASAP-024442

Abstract

Protocol to quantify DAB stained sections using NIS-Elements.



4

1 Capture brightfield images with an inverted confocal microscope with a 20X objective (Nikon A1R).

- 2 Conduct annotation of each tissue section within a bounding box of 2000×2000 pixels for mouse tissues and 2863×2454 pixels for human tissues.
- 3 Use a manual RGB-based color thresholding algorithm for mouse tissues to exclude pure methyl green nuclei staining from the measurement. For human tissues, use NIS-elements (version 5.10.01, https://www.microscope.healthcare.nikon.com/products/software/nis-elements, RRID:SCR_014329) auto-thresholding algorithm.

Export the percentage area of the thresholded signal (object area fraction provided in the program) and normalize to the average value of non-CIAP treated tissues.