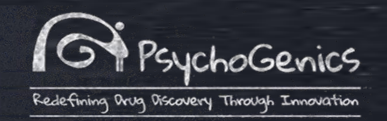
TIMECOURSE ASSESSMENT FOLLOWING BILATERAL AND UNILATERAL-DUAL INOCULATION OF ALPHA-SYNUCLEIN PREFORMED FIBRILS IN C57BL/6 MICE



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INTRODUCTION

Alpha-synuclein (α Syn) is a 140 amino acid protein implicated both genetically and neuropathologically in Parkinson's disease (PD). Increased levels of α Syn lead to neurodegeneration and aggregated α Syn is the primary component of Lewy bodies, the histopathological hallmark of PD. Synthetic α Syn (murine and human) are capable of 'seeding' and propagating α Syn pathology in both α Syn transgenic and non-transgenic (WT) neuronal cultures and mice (Luk, et al., 2012a; Luk, et al., 2012b; Volpicelli-Daley, et al., 2014). Unilateral striatal administration of synthetic murine alpha-synuclein preformed fibrils [PFFs] in WT mice has been shown to induce reliable pS129 α Syn-positive pathology at 30, 90, and 180 days post inoculation [DPI]. However, motor disruption at any timepoint hasn't been a reliable outcome measure.

We sought to build on previously published α -synuclein Parkinson's disease models by examining the pathology development over time post-inoculation of mouse α -synuclein PFF into the striatum of both hemispheres, or into the striatum and substantia nigra of one hemisphere, of C57BI6/J WT mice. PFF were inoculated via stereotaxic surgery into bilateral mouse striatum, or unilateral dual mouse striatum-substantia nigra. Behavior and immunohistochemistry were assessed 30, 60, and 90 days DPI.

METHODS

Animals: A total of 35 (n=6-16) per treatment group and dpi, six-week-old gender mixed C57Bl6/J mice were purchased from The Jackson Laboratories (Bar Harbor, ME). All animals were examined, manipulated, and weighed prior to study initiation to ensure adequate health and suitability for the study.

Alpha-synuclein pre-formed fibrils: Mouse alpha-synuclein preformed fibrils (mPFFs) (2.5 μ g/ μ l) were prepared as per Luk's preparation protocol (Luk et al, Science, 2012a).

Surgical method: Mice were anesthetized with isoflurane and stereotactically injected into the striatum of both hemispheres (for bilateral injections) or into the striatum and substantia nigra of one hemisphere (for unilateral-dual injections) - coordinates (striatum: anteroposterior: +0.2, mediolateral: ± 2.0 , dorsoventral: -2.6; substantia nigra: anteroposterior: -3.3, mediolateral: ± 1.3 , dorsoventral: -4.4), with recombinant mPFF (5.0 µg; 2.0 µl total volume). Control animals received inoculations with PBS. Injections were performed with a 10 µl syringe (Hamilton, NV) at a rate of 0.1 µl/min with the needle left in place for 5 minutes following administration at each target. All mice received analgesia both pre- and post-operatively. Wet feed was provided for post-surgical animals for a period of 3 consecutive days following surgery.

Behavior assessment: At each timepoint, all the animals were tested in wire hang and tapered balance beam tasks. The wire hang test, Santa-Maria et al. (2012)'s modified protocol in which mice were placed on top of a standard wire cage lid and the lid was then turned upside down. The latency of mice to fall off was measured, and average values were computed from three trials (30 seconds apart). Trials were stopped if the mouse remained on the lid after 5 minutes. Tapered balance beam consisted of a 100 cm in length black acrylic strip tapered from a width of 1.5 cm to 0.5 cm. The beam was angled of 17° from horizontal running from low to high [58 cm from the floor] topped with a goal box. After training, animals are tested 24 hours later. During testing, mice received 3 trials with an intertrial interval (ITI) of 30 seconds. The following measures are captured: the latency to turn [animal turn to face the goal box]; latency to traverse; total number of steps and footslips for each paw and expressed as a ratio of the total number of footslips to the total number of steps.

Brain collection and preparation for histology: Brain hemispheres were collected according to PGI' standard procedures. In short, mice were flush perfused with saline to remove blood cells before harvesting brains. Hemibrains were cryoprotected in 15% sucrose after overnight post fixation in 4% phosphate-buffered PFA. Brains were frozen within OCT in molds in dry ice cooled liquid isopentane. Tissue blocks were cut coronally in a uniform systematic random protocol on a Leica CM3050 cryotome at 10-microns section thickness.

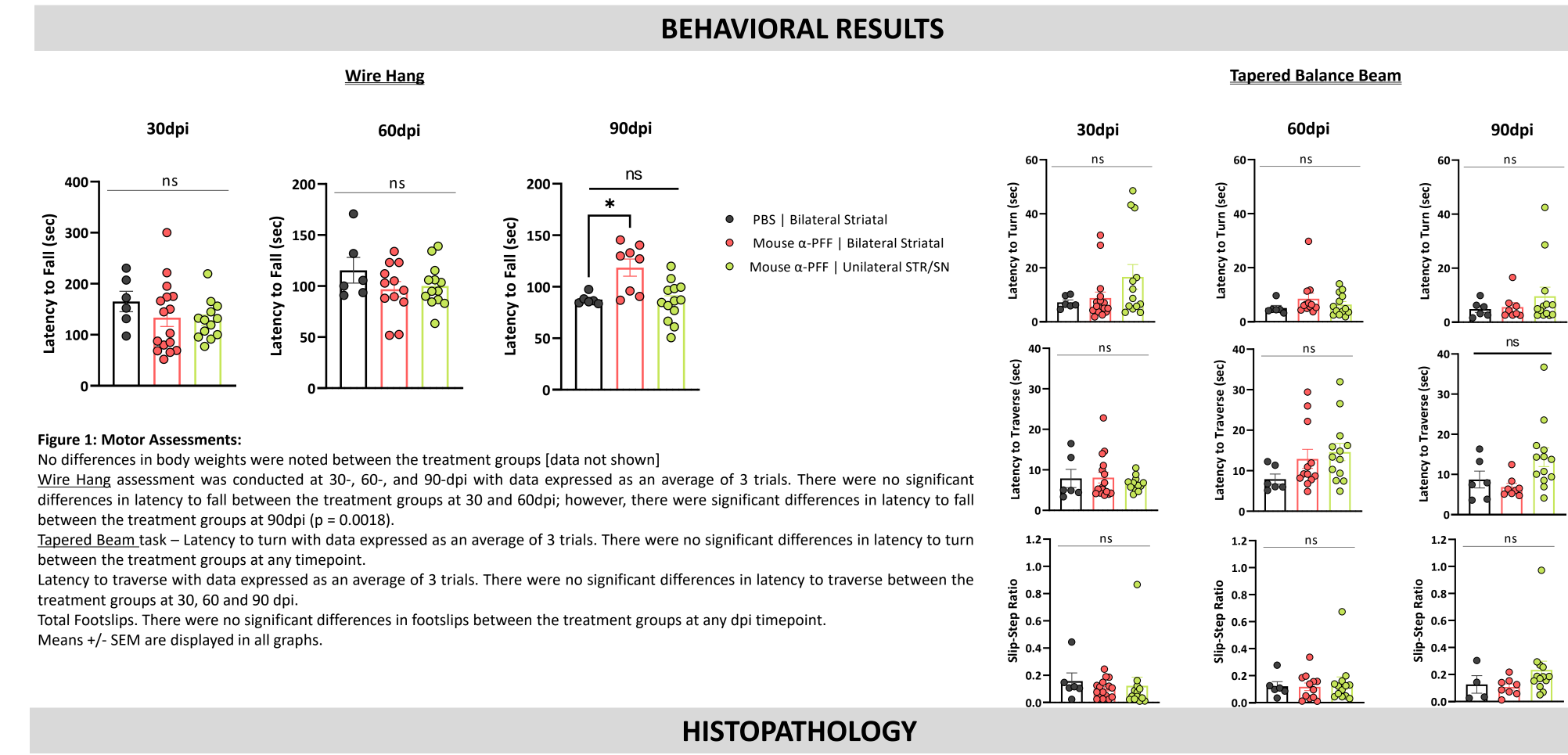
Labeling, imaging and quantification: 8 sections per sample were immunohistochemically labeled with pSer129 aSyn [EP1536Y] and TH [1B7] primary antibodies. AlexaFluor-conjugated secondary antibodies were used to visualize primary binding, all sections were counterstained with DAPI to visualize nuclei. Whole slices images were acquired on an Axio.Scan Z1 slide scanner. Dorsal striatum (CPu) and substantia nigra (SN) were manually delineated. The saved ROIs were used to count immunoreactive objects above threshold and size restrictions fully automatedly using Image Pro Premier image analysis software (v9.1 or higher).

Brain tissue preparation and rat neuronal culture experiments: Total homogenates were prepared from fresh frozen, brain tissues of PFF or PBS inoculated mice. Hemibrains were regionally dissected to isolate the cortex and midbrain including the substantia nigra. Tissues were lysed in 9x volume of homogenization buffer complemented with cOmpleteTM protease inhibitor cocktail (Sigma) and sonicated on ice prior to storage in single-use aliquots.

Rat cortical neurons at DIV5 (days *in vitro*) were exposed to brain homogenates from the cortex and midbrain regions. The formation of pS129 aSyn positive aggregates was observed upon fixation of cells on DIV22 or DIV25. An antibody for MAP2 [ab5392] was used to visualize neurons and pS129 aSyn [EP1536Y] positive-area was normalized to MAP2-positive area.

References:

Kelvin C. Luk,1 Victoria Kehm,1 Jenna Carroll,1 Bin Zhang,1 Patrick O'Brien,1 John Q. Trojanowski,1 and Virginia M.-Y. Lee1, **Pathological α-Synuclein Transmission Initiates Parkinson-like Neurodegeneration in Non-transgenic Mice**; *Science*. 2012 Nov 16; 338(6109): 949–953. PMCID: PMC3552321



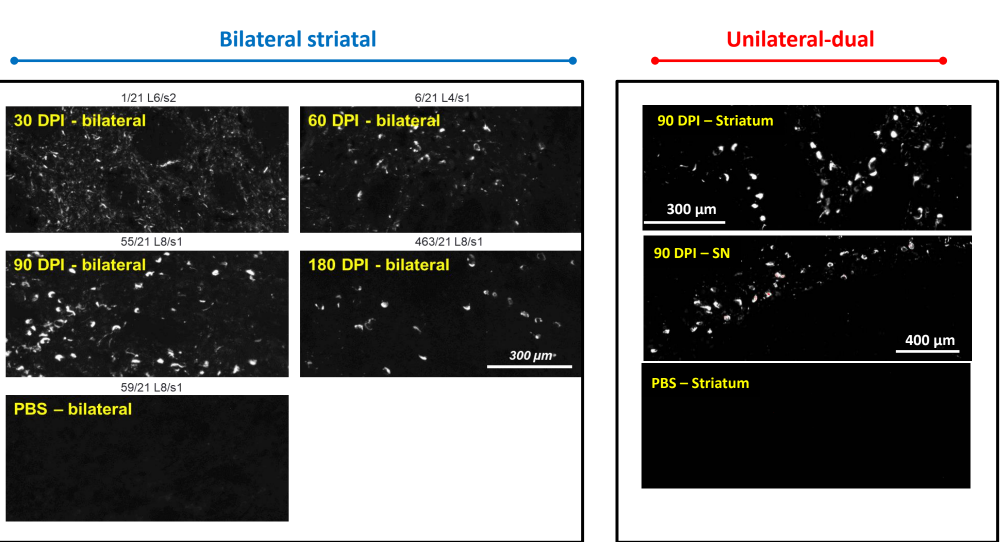


Figure 2: Representative images of pSer129 αSyn in the **dorsal striatum** of PFF-injected mice bilaterally striatal at 30, 60, 90 and 180DPI (left panel) and of unilateral-dual injected striatal and substantial nigra mice at 90 DPI (ipsi, right panel): Density of positive labeling is highest 30DPI consisting of many small sized fibrils that accumulate over time to larger size Lewy body-like aggregates. Those decrease with time due to the loss of positive neurons. No positive fibrils are present in PBS injected controls and only very few in contralateral side at 90DPI in the unilateral model (left). The pattern of pSer129+ fibrils was similar with dual-unilateral and bilateral PFF inoculation at 90 DPI.

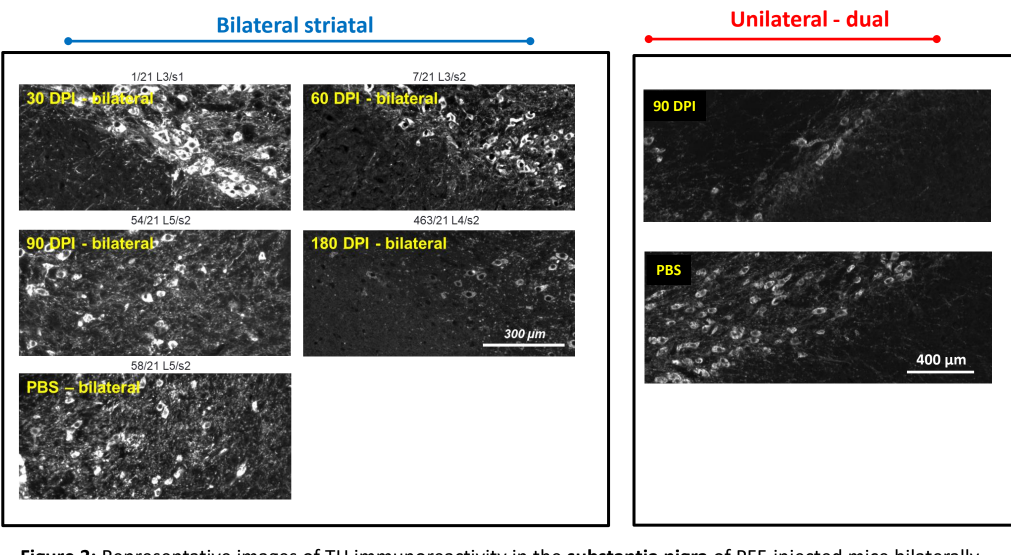


Figure 3: Representative images of TH immunoreactivity in the **substantia nigra** of PFF-injected mice bilaterally striatal at 30, 60, 90 and 180DPI (left panel) and unilaterally-dual injected striatal and substantia nigra mice at at 90 DP (ipsi, right panel): From 60DPI onwards, the number of the TH+ neurons starts to drop and this leads to significantly lower positive cells in the uni- and bilateral model at 90DPI. At 180DPI, only few cells were found positive for TH in the SN. Unilateral-dual injections led to more pronounced decrease in TH+ neurons ipsilaterally compared to unilateral or bilateral PFF injections.

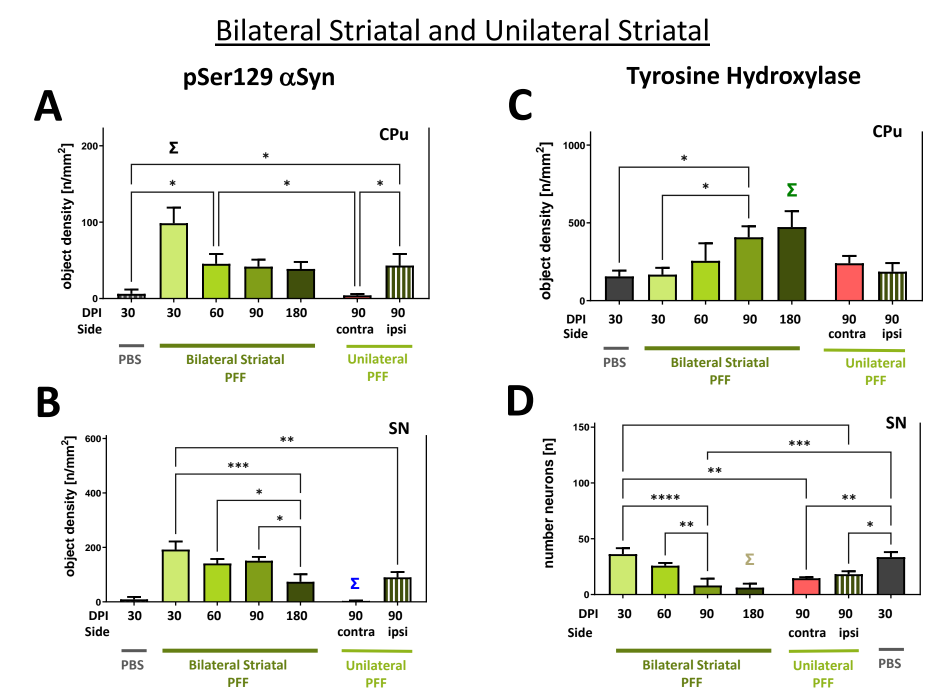


Figure 4: (A,B) pSer129 α Syn-positive object density in the CPu and SN. In bilaterally striatal-injected animals, the highest load of pSer129 α Syn is seen at 30DPI and then drops significantly. In the SN, the significant drop is seen between 90 and 180 DPI in the bilaterally striatal-injected groups. (C) TH object density in the CPu increases as a sign of fragmentation of TH labeling in bilaterally striatal-injected groups (D) TH cell number in the SN of bilaterally striatal-injected animals drops significantly from 90DPI onwards.

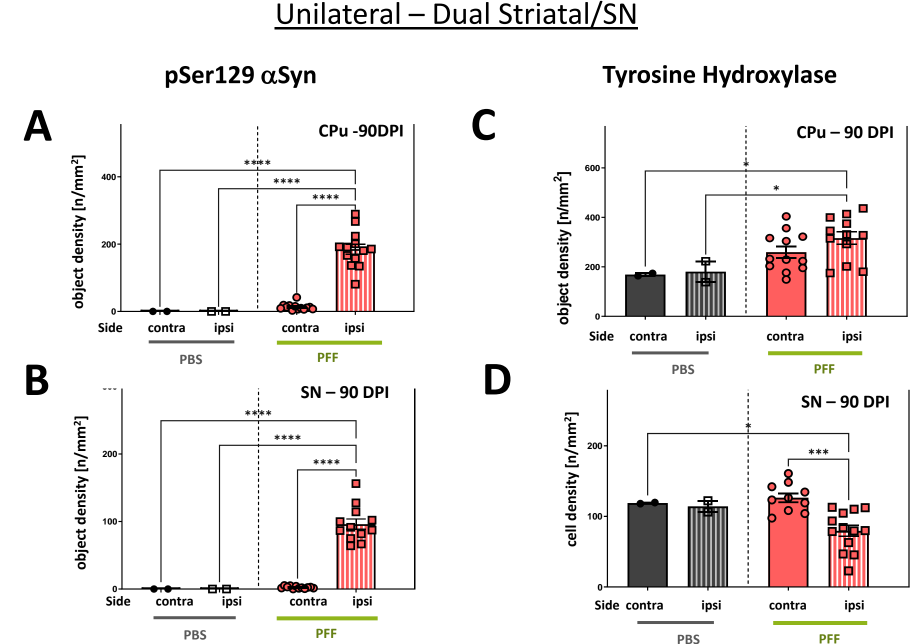
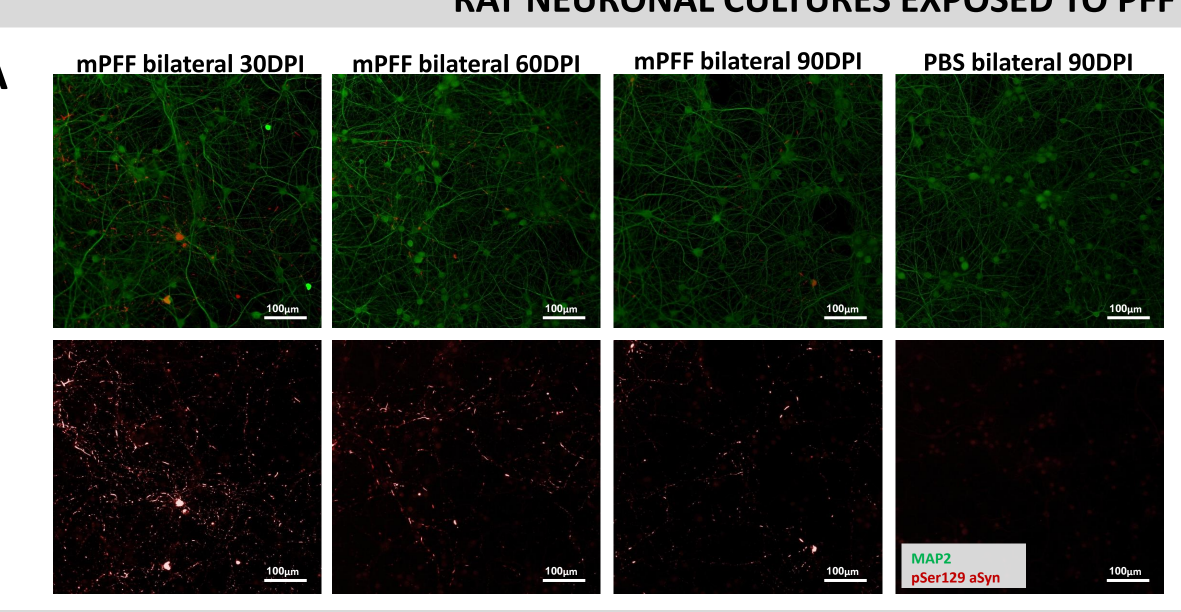


Figure 5: (A,B) pSer129 α Syn-positive object density in the CPu and SN of unilateral-dual striatal/SN inoculated animals. Increased load of pSer129 α Syn is seen in the ipsilateral CPu and SN compared to the contralateral side and to the PBS-treated groups. (C,D) TH object density in the Cpu and TH cell density in the SN. Increase in object density in the CPu and a small reduction in dopaminergic cell density in ipsilateral SN versus contralateral hemisphere.

RAT NEURONAL CULTURES EXPOSED TO PFF MOUSE TISSUES



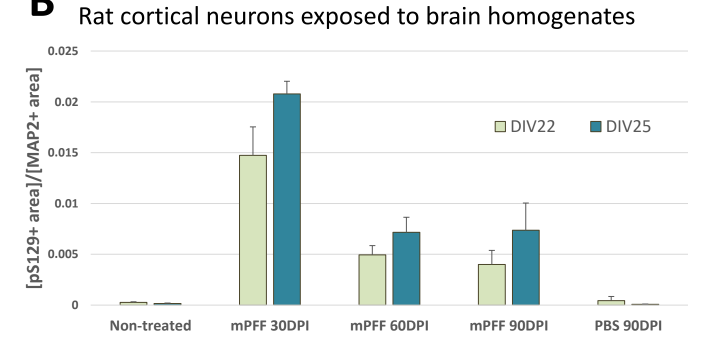


Figure 6: (A) Images of phosphorylated α Syn (red) associated with MAP2 (green) labeling in primary rat neurons fixed on DIV25 following exposure to brain tissues from PFF or PBS bilaterally-injected mice (B) When cells were exposed for longer and fixed on DIV25 vs DIV22, the pS129 α Syn signal was increased. The highest burden of pS129 α Syn positive puncta was observed in lysates from the 30DPI group. No aggregates were formed in untreated neurons or in neurons exposed to homogenates from the PBS-injected group.

CONCLUSIONS

- Bilateral striatal or dual-unilateral striatal-substantia nigral (90-day only) α-synuclein mPFF inoculation, does not significantly alter the behavioral phenotype when examined at 30-, 60-, or 90-days post inoculation at a dose of 10 μg per mouse.
- Bilateral Striatal: Highest load of pSer129 αSyn at 30DPI while size of aggregates increase with time; Reduction in pSer129 αSyn over time likely linked to cell loss as it's accompanied by drop of TH-positive cells after 90DPI.
- Unilateral-dual: increased pSer129 αSyn at 90DPI in the CPu and SN which is accompanied by an increased load of pSer129 αSyn in the CPu and SN and drop of dopaminergic cell density in the SN.
- Brain homogenates from animals inoculated with α-synuclein PFF can induce the accumulation of pSer129 αSyn-positive inclusions in a neuronal culture system of rat cortical neurons demonstrating the seeding capacity of the *in vivo* formed aggregates. Importantly, the *in vitro* pathology load correlated with *in vivo* data, as the largest load of pSer129 αSyn aggregates was observed in neurons exposed to brain homogenates collected at 30 DPI.
- Newly developed models of PFF inoculations results in overall higher load of α Syn aggregates (bilateral and dual-unilateral) and allow parallel biochemical and histological investigations in drug efficacy studies.

