

## **Characterization of pathology-inducing $\alpha$ -synuclein species from human diseased brain tissue**

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Short Title:  $\alpha$ Synuclein species from human diseased brain

## Abstract

Synucleinopathies are a group of neurodegenerative diseases characterized by the presence of pathological accumulations of misfolded, phosphorylated  $\alpha$ -synuclein ( $\alpha$ Syn) protein. Multiple lines of evidence indicate that synucleinopathy disease progression is driven by a prion-like process of transmission of a pathologic form of  $\alpha$ Syn. One potential therapeutic approach to prevent cell-to-cell propagation is to target this transmissible species with selective antibodies. In this study, a rodent primary neuronal culture reporter system was developed to monitor induction of detergent-insoluble, phosphorylated (pS129) aggregates of  $\alpha$ Syn. Induction of pS129  $\alpha$ Syn pathology was observed with both synthetic  $\alpha$ Syn fibrils (PFFs) and brain lysates from multiple system atrophy (MSA) patients but not  $\alpha$ Syn monomers or human brain lysate controls. The induction-competent species in MSA lysates could be enriched by high-speed centrifugation suggesting that it is present as a high molecular weight aggregate. Furthermore, samples derived from brain lysates from Parkinson's disease (PD) and Dementia with Lewy Bodies (DLB) patients also induced pS129  $\alpha$ Syn pathology, but required longer incubation times. Lastly, the potential of  $\alpha$ Syn selective antibodies to immunodeplete induction-competent forms of  $\alpha$ Syn from both PFF and synucleinopathy brain samples is described. The results demonstrate that antibodies targeting the C-terminal of  $\alpha$ Syn are most effective for immunodepletion of pathology-inducing forms of  $\alpha$ Syn from samples derived from human synucleinopathy brains. Furthermore, the data support the hypothesis that antibodies that recognize a C-terminal epitope and exhibit selectivity for oligomeric forms over monomeric forms of  $\alpha$ Syn represent a desirable target for immunotherapy for synucleinopathy patients.

## Introduction

$\alpha$ -synuclein ( $\alpha$ Syn) is a 140 amino acid protein preferentially expressed in neurons at pre-synaptic terminals where it is thought to play a role in regulating synaptic transmission (1). It has been proposed to exist natively as both an unfolded monomer (2) and a stable tetramer of  $\alpha$ -helices (3, 4) and has been shown to undergo several posttranslational modifications (5). One modification that has been extensively studied is phosphorylation of  $\alpha$ Syn at amino acid serine 129 (pS129). Normally, only a small percentage of  $\alpha$ Syn is constitutively phosphorylated at S129, whereas the amount of pS129-positive  $\alpha$ Syn present in pathologic intracellular inclusions is vastly increased (6). These pathological inclusions consist of aggregated, insoluble accumulations of misfolded  $\alpha$ Syn proteins and are a characteristic feature of a group of neurodegenerative diseases collectively known as synucleinopathies (7).

The pathological aggregates of  $\alpha$ Syn found in neurons are called Lewy bodies and are the characteristic hallmarks of both Parkinson's Disease (PD) and dementia with Lewy bodies (DLB). Additionally, abnormal  $\alpha$ Syn-rich lesions called glial cytoplasmic inclusions (GCIs) are found in oligodendrocytes, and represent the hallmark of a rapidly progressing, fatal synucleinopathy known as multiple systems atrophy (MSA). The idea that the progression of synucleinopathies is in part due to cell to cell transmission of pathologic forms of  $\alpha$ Syn species has gained support in recent years (8-10). The initial evidence for this hypothesis derives from the observed stereotypical progression of brain pathology described in PD (11) and from evidence of host-to-graft spreading of  $\alpha$ Syn aggregates in PD patients (12). In addition, reports of either undetectable (13-15) or low levels (16) of  $\alpha$ Syn mRNA expression in oligodendrocytes suggests that a pathological form of  $\alpha$ Syn is propagated from neurons to oligodendrocytes in MSA. Recent work supports this idea of  $\alpha$ Syn propagation, demonstrating that  $\alpha$ Syn is taken up by oligodendrocytes both *in vitro* and *in vivo* (17). Moreover, inoculation of human brain homogenates from MSA patients into  $\alpha$ Syn over-expressing transgenic mice results in neurological dysfunction and extensive pS129-positive neuronal deposits, supporting the hypothesis that MSA may be transmitted by the prion-like propagation of pathologic forms of  $\alpha$ Syn (10, 18).

In this study, an *in vitro* reporter assay was developed using rat hippocampal neuronal cultures overexpressing the human A53T  $\alpha$ Syn mutant protein. The A53T mutation is associated with familial forms of PD (19) and has been shown to accelerate  $\alpha$ Syn fibrillization *in vitro* (20)

leading to greater neurotoxicity in transgenic mice (21). We used both synthetic, pre-formed  $\alpha$ Syn fibrils (PFFs) and brain lysates from MSA patients to induce detergent-insoluble, pS129-positive immunofluorescence in neurons as a marker of  $\alpha$ Syn pathology. Furthermore, we demonstrate that aggregates isolated from brain lysates from both PD and DLB patients also induce pS129  $\alpha$ Syn pathology, but only after prolonged incubation. We then used  $\alpha$ Syn selective antibodies to immunodeplete the aggregate-inducing form from both PFF and synucleinopathy brain samples in order to understand which  $\alpha$ Syn epitopes and species hold potential as therapeutic targets to prevent the pathological transmission of  $\alpha$ Syn aggregates.

## Results

### *Assay characterization using PFF*

To determine if application of human  $\alpha$ Syn PFF could induce a detergent-insoluble pS129  $\alpha$ Syn signal in primary rat neuronal cultures, hippocampal cells at 7 days in vitro (DIV) were treated with 150 ng/ml  $\alpha$ Syn PFF generated from full-length recombinant hA53T- $\alpha$ Syn. This treatment led to a modest induction of pS129  $\alpha$ Syn immunofluorescence after 11 days in the absence of cellular toxicity (supplemental Figure S1). These results confirm previous reports that exogenous recombinant human  $\alpha$ Syn PFF can induce the recruitment of endogenous rodent  $\alpha$ Syn into insoluble pS129  $\alpha$ Syn aggregates (22, 23). In order to create a more sensitive assay system, primary rat hippocampal neurons were transduced with AAV-hA53T- $\alpha$ Syn at DIV 4 to overexpress hA53T- $\alpha$ Syn. Transduction with AAV-hA53T- $\alpha$ Syn alone did not induce accumulation of insoluble pS129  $\alpha$ Syn signal; however, application of 150 ng/ml PFF led to a robust induction of insoluble pS129  $\alpha$ Syn staining 11 days following treatment (Figure 1A-B). PFF-mediated induction in the AAV-hA53T- $\alpha$ Syn-transduced neurons was ~20-fold higher than induction in the neurons exposed to an empty AAV vector (Figure 1B). In addition, increasing the MOI of AAV-hA53T- $\alpha$ Syn led to an increase in the PFF-induced pS129 signal (supplemental Figure S2A) suggesting that the concentration of intra-cellular  $\alpha$ Syn is a key parameter regulating induction by PFFs. Due to the report of cross-reactivity of the 81A anti- $\alpha$ Syn pS129 antibody with phosphorylated neurofilament light chain (NfL) (24), specificity of the 81A signal was confirmed by demonstrating a comparable staining pattern of insoluble pS129 with a second anti-pS129 antibody, MJFR13, which does not cross-react with phosphorylated NfL (supplemental Figure S3) (25).

PFF-mediated induction of insoluble pS129 in AAV-hA53T- $\alpha$ Syn-transduced hippocampal neurons was concentration-dependent and did not cause significant toxicity (Figure 1C). In contrast, a pS129  $\alpha$ Syn signal could not be elicited with 4500 ng/ml of hA53T- $\alpha$ Syn monomer (Figure 1D) or with PFF of tau protein (data not shown). Furthermore, PFF-dependent increases in pS129  $\alpha$ Syn were inversely correlated with the level of branch points in MAP2-positive neurites (supplemental Figure S2B). These data are consistent with a previous publication reporting that overexpression of hA53T- $\alpha$ Syn in rat primary midbrain neurons results in altered neurite morphology, including decreased neurite branching (26). Our results demonstrate that detergent

insoluble pS129  $\alpha$ Syn pathology can be induced with hA53T- $\alpha$ Syn PFF in primary rat hippocampal neurons overexpressing hA53T- $\alpha$ Syn and is correlated with impaired neurite morphology.

#### *Assay characterization using human brain lysates*

Having developed a highly sensitive reporter assay, we next sought to determine if lysates generated from brain samples of patients diagnosed with various synucleinopathies could induce pS129  $\alpha$ Syn pathology. PBS-soluble lysates were generated using frontal cortex tissue samples from MSA, PD, DLB+AD patients and normal controls (demographic and neuropathological information are included in supplemental Table S1 and S2, respectively). AAV-hA53T- $\alpha$ Syn-transduced hippocampal neurons were treated with lysates and the induction of insoluble pS129  $\alpha$ Syn measured. MSA lysates induced a significant increase in pS129  $\alpha$ Syn signal 11 days post treatment in contrast to brain lysates from PD, DLB+AD and normal controls (Figure 2A-B, supplemental Table S3). Induction was robust (~100-fold of untreated control) and significant toxicity was not observed. A total of 9 different MSA brain lysates were then tested at different concentrations (Figure 2C, supplemental Table S4). Statistically significant induction was observed with 5 of the 9 MSA lysates tested at a 1:300 dilution, with the highest measured intensity (MSA #1, MSA #7) resulting in ~50% of the pS129 signal induced by a 150 ng/ml PFF treatment (Figure 2C, supplemental Table S4). Variability in the absolute degree of induction for MSA #1 and MSA #3 (compare Figure 2B and 2C) was observed and likely reflects inter-batch differences in the primary neuron cultures. Nevertheless, the rank order of the ability and degree to which individual MSA brain samples induced was consistent across several primary neuron preparations. MSA lysate induction was concentration dependent and lysates did not produce toxicity with the exception of the highest concentration tested (1:100; data not shown). These results are in agreement with a previous reports demonstrating that only lysates from MSA patients and not PD patients induce  $\alpha$ Syn aggregation in HEK cells expressing  $\alpha$ Syn A53T-YFP, and that different MSA brain lysates produce different magnitudes of induction (10, 27).

The magnitude of induction by the MSA lysates did not correlate with  $\alpha$ Syn levels (supplemental Figure S4A) suggesting that the inducing species may be a minor component of the overall  $\alpha$ Syn in the extract. In addition, induction did not correlate with the extent of glial cytoplasmic

inclusions (GCIs) observed in tissue sections, suggesting that there may be multiple variables affecting the level of the PBS-soluble inducing species (supplemental Figure S4B).

High-speed centrifugation has been used to partially-purify the pathology-inducing species of tau from brain lysates (28) and a similar approach was used to further characterize the inducing species from the MSA samples. PBS-soluble MSA brain lysates were subjected to high-speed centrifugation (100,000 x g, 30 minutes) and the resulting supernatant and pellet fractions isolated and tested for induction in transduced neurons. As shown in Figure 2D, inducing activity was completely recovered in the re-suspended pellets but not the supernatant fractions from the MSA lysates. This finding indicates that the inducing species can be enriched by high speed centrifugation and is likely composed of high molecular weight aggregates.

#### *Longer incubation times with PD and DLB+AD brain lysates lead to pS129 $\alpha$ Syn induction*

Both the data presented here, as well as recently published data, suggest that MSA and PD may contain different strains of toxic  $\alpha$ Syn species due to their differential ability to induce pathology in various biological systems (10, 27). Because of these potential strain differences, we hypothesized that DLB+AD or PD brain lysates may require longer incubation times for induction of pS129  $\alpha$ Syn pathology in our in vitro system. As with the MSA samples, high-speed centrifugation was used to enrich the presumed  $\alpha$ Syn aggregates from control, PD and DLB+AD lysates. Levels of  $\alpha$ Syn in starting brain lysates and isolated supernatant and pellet fractions were measured by ELISA (Figure 3, supplemental Table S5). As shown in Figure 3, similar levels of  $\alpha$ Syn were detected across the starting lysates and the pellet fractions regardless of clinical diagnosis. On average ~3% of the starting  $\alpha$ Syn ELISA signal was recovered in the pellet fraction while ~32% was recovered in the supernatant fraction (supplemental Table S5). The reasons for the modest recovery of total  $\alpha$ Syn starting material (~35%) following centrifugation are not clear. These results indicate that high molecular weight species of  $\alpha$ Syn, presumably aggregates, are present in all of the brain lysates, including control lysates. Furthermore these high molecular aggregates represent a minor fraction (~3%) of the total  $\alpha$ Syn present in the starting lysate. Re-suspended pellets were then tested for inducing activity in AAV-hA53T- $\alpha$ Syn-transduced hippocampal neurons. Cells were treated for up to 32 days, 3 weeks longer than previous experiments. As expected, significant induction of pS129 was observed with MSA pellet fractions

following 11 days of treatment (Figure 4, supplemental Table S6) and the induction was significant at all time points. The ~2-fold higher induction at 11 days compared to the later time points likely reflects inter-plate differences in background normalization. Interestingly, significant induction was also observed with the DLB+AD and PD pellet fractions but only after 25 days of incubation (Figure 4, supplemental Table S6). In contrast, pellet samples from control brain lysates failed to induce pS129  $\alpha$ Syn, regardless of duration of exposure (Figure 4, supplemental Table S6). As the total  $\alpha$ Syn levels were similar across all of the pellet fractions (Figure 3), the observed differences in induction are likely related to differences in the molecular nature and/or conformation of the high molecular weight species enriched by high speed centrifugation.

#### *Binding of $\alpha$ Syn antibodies to $\alpha$ Syn monomers and PFFs*

To further investigate the molecular nature of the induction-competent species found in  $\alpha$ Syn PFF and MSA brain lysates, a set of ten commercially available  $\alpha$ Syn antibodies were selected for immunodepletion studies. Antibodies were selected to target different epitopes across the  $\alpha$ Syn sequence (Figure 5, supplemental Table S7). A competitive binding assay was used to characterize antibody affinity to  $\alpha$ Syn monomers and PFF and example binding curves for MJFR1 and MJFR14642 are shown in Figure 5. A summary of relative affinity  $IC_{50}$  values is included in Table 1. As shown in Figure 5B, MJFR1 exhibited comparable binding to both  $\alpha$ Syn monomer and  $\alpha$ Syn PFF with  $IC_{50}$  values of 8.0 ng/ml and 29.2 ng/ml, respectively, and a monomer to PFF binding ratio of 0.27 (Table 1). In contrast, MJFR14642 exhibited 50-fold higher affinity for PFF and over 400-fold weaker affinity for monomer compared to MJFR1 (Figure 5C, Table 1) demonstrating that MJFR14642 is more selective for the oligomeric form. Indeed, MJFR14642 exhibited the greatest degree of selectivity for PFF (monomer to PFF binding ratio of 7173) of all the antibodies tested (Table 1). Only 4D6 and LB509, two other C-terminal antibodies, produced monomer to PFF binding ratios greater than 10 (26 and 39, respectively; Table 1).

#### *Immunodepletion of PFF and MSA-induced pS129 $\alpha$ Syn pathology with $\alpha$ Syn antibodies*

The ability of each antibody to immunodeplete the  $\alpha$ Syn inducing species from either  $\alpha$ Syn PFF or a MSA brain lysate sample was then evaluated. Antibodies were tested at a concentration of 50 nM. As we were unable to demonstrate binding of antibody 5C2 to either monomer or PFF up to 15000 ng/ml (Table 1), this antibody, along rabbit IgG, were included as negative controls. PFF

$\alpha$ Syn or MSA brain lysate sample #7 were incubated overnight with each antibody, the antibody complexes removed and the immunodepleted samples tested for induction of pS129  $\alpha$ Syn in AAV-hA53T- $\alpha$ Syn-transduced hippocampal neurons after 11 days of treatment (Figure 6). The greatest depletion of the inducing species from PFF was observed with the C-terminal antibodies LB509, 14H2, MJFR1 and MJFR14642 with reductions of >90%. In contrast, the N-terminal antibodies Syn303 and EP1646Y exhibited limited ability to immunodeplete the PFF inducing species, producing reductions of ~30% and ~50%, respectively. This result is consistent with the lower activity of Syn303 compared to the C-terminal antibody Syn211 in inhibiting PFF-mediated induction of pathology in mouse neurons as previously reported (23). The pS129-selective antibody MJFR13 was unable to immunodeplete the inducing species from PFF, consistent with the lack of pS129  $\alpha$ Syn in the PFF samples.

MJFR1 and MJFR14642 were the most active antibodies for immunodepleting the induction-competent species from the MSA brain lysate with reductions of >80% observed (Figure 6). Interestingly, LB509, another C-terminal antibody, exhibited limited ability to deplete the inducing species from the MSA lysate (~50%) even though this antibody depleted >90% of the inducing species from PFF. The lower than expected activity of LB509 for depleting the inducing species from MSA could reflect potential differences in the conformation and/or modifications of the MSA species compared to PFF. Similar to the results with PFF, both N-terminal antibodies were less effective for depleting the inducing species from MSA. In contrast to the results with PFF, the pS129 antibody MJFR13 immunodepleted the MSA  $\alpha$ Syn inducing species by ~70%, indicating the presence of pS129  $\alpha$ Syn. This observation is intriguing considering that phosphorylated oligomeric  $\alpha$ Syn has been shown to be significantly elevated in the CSF of MSA patients as compared to other synucleinopathies (29).

Since both MJFR1 and MJFR14642 were highly effective for depleting the inducing species from PFF and the MSA lysates, these antibodies were selected for concentration response curve (CRC) analysis (Figure 7). Both antibodies exhibited similar maximal efficacy and potency in depleting the PFF inducing species (Figure 7A;  $IC_{50}$  values derived from several experiments are included in Table 2);  $IC_{50}$  values of 0.036 nM and 0.079 nM were derived for MJFR1 and MJFR14642, respectively. To determine if differences in oligomer selectivity influenced the ability of each antibody to deplete the inducing species in the presence of monomer, PFF immunodepletion reactions with either 1 nM MJFR1 or 1 nM MJFR14642 were spiked with increasing

concentrations of  $\alpha$ Syn monomer and the depleted samples tested for inducing activity (Figure 7C). In the absence of spiked monomer, both antibodies completely depleted the inducing species from PFF as expected. Addition of  $\alpha$ Syn monomer effectively blocked the ability of MJFR1 to deplete the inducing species in PFF in a dose-dependent manner (Figure 7C). In contrast, MJFR14642 was ~100-fold less sensitive to competition with  $\alpha$ Syn monomer (Figure 7C). Taken together, these results confirm the superior selectivity of MJFR14642 and suggest that this property may translate into more potent activity against the induction-competent species in biological fluids that contain competing levels of  $\alpha$ Syn monomer (e.g. CSF and ISF).

MJFR1 and MJFR14642 also depleted the inducing species from MSA lysate #7 in a concentration dependent manner (Figure 7B). Both antibodies were weaker for immunodepleting the inducing  $\alpha$ Syn species from the MSA lysate compared to PFF (Table 2). Interestingly, the oligomer-selective antibody MJFR14642 was ~10-fold more potent in depleting the MSA inducing species compared to the non-selective MJFR1 antibody (Table 2). This ~10-fold difference in potency was also maintained when the MSA brain pellet was used to induce  $\alpha$ Syn aggregates (13.7 fold, Table 2, N=1). This finding suggests that the MJFR14642 vs. MJFR1 potency difference is likely due to differences in conformation and/or epitope exposure of the induction-competent species, rather than competitive binding to other non-inducing  $\alpha$ Syn found in the crude brain lysates (e.g.,  $\alpha$ Syn monomers).

Both  $\alpha$ Syn antibodies MJFR1 and MJFR14642 were also able to immunodeplete the inducing species from pelleted PD and DLB brain lysate samples in a concentration dependent manner (Table 2). Interestingly, the  $\alpha$ Syn oligomer-selective MJFR14642 exhibited 7.1-fold and 13.8-fold greater potency than MJFR1 for immunodepleting the inducing  $\alpha$ Syn species from a DLB brain pellet and a PD brain pellet, respectively (Table 2). These data confirm that induction of pS129  $\alpha$ Syn by PD and DLB extract pellets is mediated by  $\alpha$ Syn aggregates and further support the hypothesis that C-terminal, oligomer-selective  $\alpha$ Syn antibodies are more specific for the physiologically-relevant transmissible forms of  $\alpha$ Syn from synucleinopathy patients.

## Discussion

The hypothesis that the pathological initiation and progression of synucleinopathies is in part due to cell to cell transmission of toxic  $\alpha$ Syn species in a prion-like fashion has gained support in recent years (8, 9). As such, the concept of an immunotherapy in which  $\alpha$ Syn-selective antibodies bind to and prevent the spread of the pathological  $\alpha$ Syn species responsible for templating toxic  $\alpha$ Syn aggregates has become an attractive disease-modifying hypothesis and approach (30). Our data demonstrate that  $\alpha$ Syn antibodies can bind to induction-competent  $\alpha$ Syn species in synthetic  $\alpha$ Syn PFFs and pathologically-relevant  $\alpha$ Syn aggregates found in brain lysates from synucleinopathy patients, thereby preventing the induction of insoluble pS129  $\alpha$ Syn-positive aggregates in primary rat hippocampal cultures.

There are three main findings from the experiments described in this report. First, we demonstrate that frontal cortex lysates from synucleinopathy patients induce pathological pS129  $\alpha$ Syn in a rodent primary neuron system. While pS129  $\alpha$ Syn induction has been shown previously in rodent primary cultures using synthetic  $\alpha$ Syn PFFs (22, 23), and brain tissue samples from MSA patients have been reported to induce  $\alpha$ Syn aggregates in  $\alpha$ Syn-expressing HEK cells (10, 27), this is the first report to demonstrate the ability of MSA brain tissue samples to induce insoluble pS129-positive aggregates of  $\alpha$ Syn in primary neuronal cultures. Moreover, while our initial data demonstrating a lack of induction of pS129  $\alpha$ Syn following 11 days of treatment with either PD or DLB brain lysates agree with previous data showing the inability of PD brain tissue to induce  $\alpha$ Syn aggregates in  $\alpha$ Syn-expressing HEK cells (10, 27), we demonstrate for the first time that PD and DLB brain tissue samples can induce a pathological signal in neurons after prolonged incubation times. This result is especially intriguing given the clinical observation that MSA typically progresses much more rapidly than PD and DLB (31). This finding is consistent with the idea that different toxic  $\alpha$ Syn strains are present contingent on the differing synucleinopathy clinical and pathologic states (32, 33). Furthermore, the current data support the hypothesis that different conformations and/or posttranslational modifications can confer varying degrees of  $\alpha$ Syn strain toxicity or infectivity, which may in turn drive the rate of disease progression (10, 32, 34, 35). The potential differences between the discrete  $\alpha$ Syn strains have yet to be clearly elucidated. There are reports of different amounts of SDS-soluble and insoluble  $\alpha$ Syn species in PD, DLB and MSA (36, 37) but the significance of these findings is unclear and additional work is required to

identify the specific conformational and/or posttranslational differences across toxic  $\alpha$ Syn strains. Finally, while the strain hypothesis is an attractive explanation for the observed temporal differences in induction between MSA, PD and DLB, we cannot rule out the possibility that there is simply a greater concentration of the same strain of induction-competent  $\alpha$ Syn in MSA relative to PD and DLB brain lysates.

A second finding is the superior potency and maximal efficacy of C-terminal antibodies in immunodepleting toxic  $\alpha$ Syn species from diseased brain tissue samples. The potential for  $\alpha$ Syn immunotherapy in synucleinopathies with C-terminal antibodies has been demonstrated in transgenic mouse models overexpressing  $\alpha$ Syn (36-38), and immunotherapy with C-terminal antibodies has been proposed as a preferred strategy for targeting exposed regions of aggregated  $\alpha$ Syn in order to inhibit fibrillization and membrane permeabilization (39). Here we report that C-terminal antibodies exhibit higher potency to immunodeplete induction-competent  $\alpha$ Syn species present in MSA patient brain tissue compared with antibodies targeting the N-terminal region of  $\alpha$ Syn. The C-terminal is the site of S129 phosphorylation and also contains a protease cleavage site at D119 that allows for the generation of a C-terminal truncated form of  $\alpha$ Syn (40). The C-terminal truncated form of  $\alpha$ Syn has been shown to be both a component of  $\alpha$ Syn lesions (41, 42) and to enhance oligomerization and toxicity (41, 43-45). Furthermore, it has been shown that preventing C-terminal truncation with antibodies directed to the cleavage site is protective both in vitro and in vivo (46, 47). Of the 5 C-terminal antibodies tested here, 3 have epitopes that overlap this cleavage site (MJFR1, 14H2L1, LB509), while the other 2 are carboxy to this site (MJFR14642, 4D6). It is interesting to note that the most potent antibody for immunodepletion of the inducing  $\alpha$ Syn species in MSA brain tissue, MJFR14642, binds to an epitope at the most distal region of  $\alpha$ Syn and carboxy to the cleavage site. This observation suggests that while C-terminal truncated forms of  $\alpha$ Syn may initiate the formation of and/or comprise a part of the inducing species in MSA brain tissue, an  $\alpha$ Syn species with an intact C-terminal must be a major component of the induction-competent  $\alpha$ Syn species found in the brains of MSA patients.

A third main finding is the increased potency of the highly oligomer-selective antibody MJFR14642, compared with the non-selective antibody MJFR1, for immunodepletion of the induction-competent  $\alpha$ Syn species present in MSA, PD and DLB brain tissue samples. While both of these antibodies were equipotent and equally efficacious for immunodepleting the induction-

competent  $\alpha$ Syn species in PFF samples, MJFR14642 exhibited a ~10-fold greater potency for immunodepleting the induction-competent  $\alpha$ Syn species in synucleinopathy brain samples. The reasons for the increased potency are unclear but may be related to a mixture of inducing species in the MSA samples for which the antibodies have varying affinity, or improved avidity of MJFR14642 for the inducing species. One limitation of this current study is that estimates of potency with the clinical samples are based on a limited number of samples and replicates. Nevertheless, these data support the notion of differences between inducing  $\alpha$ Syn species in PFF and MSA brain tissue samples, as well as suggesting that  $\alpha$ Syn oligomer-selective antibodies may be a preferred immunotherapy for synucleinopathies compared to  $\alpha$ Syn antibodies that do not distinguish between conformations. Several oligomer-selective antibodies have been generated (48, 49), indicating the feasibility of generating antibodies targeting specific  $\alpha$ Syn conformations. Moreover, due to the relatively high plasma and CSF concentrations of soluble  $\alpha$ Syn monomer compared with the oligomeric / aggregated form of  $\alpha$ Syn (50-52), the binding of oligomer-selective antibodies to the toxic  $\alpha$ Syn species would be substantially less hindered due to low affinity for the more prevalent monomeric species. Reduced binding to monomeric  $\alpha$ Syn represents at least two advantages for oligomer-selective antibodies. First, the probability of interference by endogenous monomeric  $\alpha$ Syn function is reduced and second, clinical doses of oligomer-selective antibodies would be predicted to be lower than non-selective antibodies. This point is critical in terms of low brain to plasma exposures seen with antibody treatments, with approximately 0.1-0.3% brain penetration typically reported (53). High concentrations of free antibody will be required to achieve relevant concentrations in brain parenchyma. Therefore,  $\alpha$ Syn antibodies that are selective for  $\alpha$ Syn oligomers and bind less  $\alpha$ Syn monomer in plasma and the interstitial space will result in a greater fraction of free antibody available to access the CNS, reach therapeutic levels in the brain and neutralize oligomeric species in the interstitial fluid.

Finally, the in vitro system used in this study represents a robust assay for detecting pathological induction with relatively low amounts of the toxic  $\alpha$ Syn species. However, the limitations of the system include that it is a rat rather than human preparation, that overexpression of mutant human  $\alpha$ Syn is required to produce a pS129  $\alpha$ Syn signal with sufficient signal-to-noise to enable antibody potency estimates and that incubation with diseased brain parenchyma lysates rather than interstitial fluid are used as the source of inducer. Nevertheless, the assay system also includes unique strengths. It is a relatively sensitive assay conducive for identifying potent  $\alpha$ Syn antibodies

that recognize pathologically-relevant  $\alpha$ Syn aggregates, it could serve as a potential biomarker assay if optimized for CSF samples in terms of both patient selection and target engagement studies for clinical trials, and it holds promise to further explore the process of  $\alpha$ Syn aggregate uptake and seeding mechanisms (54). Finally, it would be of interest to investigate potential functional consequences of this robust pS129  $\alpha$ Syn signal in future studies to determine if changes in neuronal activity either precede or follow these phosphorylation events. Indeed, alterations in network connectivity have been previously reported in PFF-treated wildtype mouse hippocampal neurons as early as 4 days post treatment (22).

In conclusion, we have demonstrated discrete pS129  $\alpha$ Syn pathology in rodent primary neurons following exposure to MSA, PD and DLB brain tissue, and that antibodies targeting the C-terminal of  $\alpha$ Syn are most effective at immunodepleting these potentially distinct transmissible forms of  $\alpha$ Syn. We propose that C-terminal antibodies selective for oligomeric forms over monomeric forms of  $\alpha$ Syn represent a desirable immunotherapy for synucleinopathies due to immunodepletion of pathologically-relevant forms of transmissible  $\alpha$ Syn species found in the diseased brain of synucleinopathy patients.

## Materials and Methods

Chemicals, reagents and kits were purchased from ThermoFisher (Waltham, MA) unless indicated otherwise.

### *Preparation of $\alpha$ -synuclein pre-formed fibrils (PFFs)*

Recombinant human  $\alpha$ Syn (rPeptide; Bogart, GA) and human  $\alpha$ Syn containing the A53T mutation (rPeptide; Bogart, GA) were reconstituted with H<sub>2</sub>O to a final concentration of 1 mg/ml in 20 mM Tris/HCl, 100 mM NaCl, pH 7.4. Samples were incubated in 2 ml eppendorf tubes (~1 ml/vial) at 37°C with constant shaking for 4 days and then centrifuged at 100,000 x g at room temperature (RT) for 20 minutes (min). The pellets were re-suspended with PBS for a final PFF concentration of 1 mg/ml (PFF concentrations expressed as monomer equivalent). PFF aliquots were frozen in liquid nitrogen and stored at -20°C. Fibrillization of  $\alpha$ Syn was monitored by a Thioflavin T binding assay (supplemental Figure S5). Briefly, samples were diluted to 0.5 mg/ml in PBS and added to an equal volume of 25  $\mu$ M Thioflavin-T. Samples were then measured using an Envision multilabel plate reader (PerkinElmer; Waltham, MA) with the excitation and emission wavelengths set at 485 nm and 535 nm, respectively. Total protein levels were assessed with a Micro BCA Protein Assay Kit.

### *Preparation of human brain lysates*

Frontal cortex brain tissue samples from normal controls and Multiple System Atrophy (MSA) and Dementia with Lewy Bodies plus Alzheimer's disease (DLB+AD) cases were obtained from the Banner Sun Health Research Institute (Sun City, AZ). Frontal cortex brain tissue samples from Parkinson's disease (PD) cases and normal controls were obtained from John Q. Trojanowski (University of Pennsylvania, Philadelphia, PA). Demographic information and neuropathology for each case are included in supplemental Tables S1 and S2, respectively. Brain samples were sonicated in filtered PBS (1 ml PBS/100 mg weight tissue) with a KONTES Micro Ultrasonic Cell Disrupter for 2 x 10 seconds (sec). Samples were placed in 2 ml eppendorf tubes and the tubes were kept on wet ice during sonication. Brain lysates were centrifuged at 3,000 x g, 4°C for 5 min to remove particulate material. The supernatant (PBS-soluble) was isolated, divided into aliquots and frozen in liquid N<sub>2</sub> and stored at -80°C.

### *High-speed centrifugation isolation of human brain lysates*

Sedimentation of high molecular weight  $\alpha$ Syn aggregates was adapted from (28). PBS-soluble brain homogenates previously prepared at 100 mg/ml in PBS were diluted 3-fold to 33.3 mg/ml in ice cold PBS followed by centrifugation at 100,000 x g for 30 min at 4°C. The supernatant (Supe) was removed and saved for analysis. The pellet was re-suspended in ice cold PBS in the same volume as the starting sample.

### *Total and oligomer $\alpha$ -synuclein binding assay*

Oligomer and monomer binding assays were performed essentially as described (49). Briefly, 96 well plates (Corning CoStar) were coated with 100  $\mu$ l of 1  $\mu$ g/ml  $\alpha$ Syn WT PFF in PBS at RT for 2 hours (hr) or 4°C overnight (similar results were obtained with both). The plates were then washed 3 times with ~300  $\mu$ l of wash buffer (0.05% Tween in PBS) and blocked with 150  $\mu$ l of 3% BSA (Fraction V, protease-free, Roche Diagnostics Corporation; Indianapolis, IN) in PBS at room temperature for 2 hr (or 4°C overnight). 3-fold serial dilutions of PFF were prepared (final concentration range 1  $\mu$ g/ml to 6 pg/ml, monomer equivalent) and  $\alpha$ Syn WT monomer (final

concentration range 10 µg/ml to 60 pg/ml) for 12 concentrations in sample buffer (0.1% BSA/0.05% Tween/PBS, 2 tablets of cOmplete Protease Inhibitor Cocktail (Roche Diagnostics Corporation; Indianapolis, IN) in 50ml buffer). An equal volume of 2-fold concentrated αSyn PFF or monomer was mixed with 2-fold concentrated antibodies in low binding plates (Becton Dickinson; Franklin Lakes, NJ) and incubated at room temperature for 2 hr. 100 µl of mixtures of antibody and PFF/monomer were loaded into the PFF coated plate and incubated at room temperature for 10 min, then washed 3 times. 100 µl of donkey anti-human IgG (Jackson ImmunoResearch; West Grove, PA), with 50% glycerol 1:1000 diluted in PBSTB (1% BSA/0.2% Tween/PBS) were then added and incubated at RT for 1 hr. Plates were washed 3 times for 5~10 min per wash. 100 µl of AP substrate (Tropix CDP Star Ready-to-Use with Sapphire II, Applied Biosystems; Waltham, MA) were then added and the reaction was developed at RT for 30 min. Luminescence counts were read with EnVision. PFF was sonicated 15-times at 1 sec/pulse before coating or mixing with antibodies. The plates were maintained with constant shaking during the assay.

#### *Total αSyn ELISA*

Corning High Binding 96-well ELISA plates were coated with MJFR1 (Abcam, Cambridge, United Kingdom) (0.25 µg/ml) in Tris-buffered saline (TBS) (50 µl/well) for 1 hr at 37°C. Plates were then washed 4-times with 300 µl/well of TBS with 0.05% tween-20 (TBST) and then blocked for at least 2 hrs with 3% BSA/TBS at RT with shaking. The plates were washed as above followed by addition of the standard curve of synuclein peptide prepared in sample diluent (1% BSA/TBST). The highest standard concentration was 1000 pg/ml. Test samples were diluted in sample diluent, added at 50 µl/well to the ELISA plates and the plates incubated overnight at 4°C. After the overnight incubation, the secondary antibody (Clone-42, BD Biosciences, San Jose, CA), previously conjugated to alkaline phosphatase (AP) using EZ-link AP conjugation kit (Novus Biologicals, LLC, Littleton, CO), was added to the ELISA plates at 50 µl/well without washing and incubated for 1 hr at RT. Plates were then washed 4-times with 300 µl/well TBST and prior to addition of 100µl/well Tropix CDP star ready substrate. Signal was measured using a TopCount (PerkinElmer, Waltham, MA). Data analysis was performed using Graphpad Prism (Graphpad Software, Inc. La Jolla, CA).

#### *Primary cell culture isolation*

Primary rat hippocampal neuronal cultures were prepared weekly from ~7 litters at embryonic day 19 (E19) using the Papain Dissociation System, according to manufacturer's instructions (Worthington Biochemical; Lakewood, NJ). Rat hippocampal cells were plated on poly-D lysine coated 96 well imaging plates (BD Biosciences, San Jose, CA) (~16 plates per week) at 30,000/well in neuronal culture medium (neural basal medium (NBM) containing 0.5 mM GlutaMax and B-27 supplement (Invitrogen; Carlsbad, CA).

#### *Immunoprecipitation experiments*

Previously prepared brain homogenate (100 mg/ml) was diluted 150-fold in neuronal culture medium. Protein A/G agarose beads were prewashed once with PBS + 0.05% tween-20, then 3-times with PBS and then blocked with PBS + 1% BSA for 2 hr at 4°C prior to use. Test αSyn antibodies were added to the diluted brain homogenate and samples incubated for 2 hr at 4°C with end-over-end rotation. Prewashed and preblocked Protein A/G agarose bead slurry was then added at a 1:10 dilution and samples incubated overnight at 4°C with end-over-end rotation. Samples

were then centrifuged at 1500 x g for 2 min to pellet the beads. The depleted supernatant was then removed and used for treatment in the primary neuron induction assay.

### *Immunofluorescence assay*

On day *in vitro* (DIV) 4, rat hippocampal neurons were transduced with an adeno-associated viral vector, AAV1 containing the cDNA for human  $\alpha$ Syn harboring the A53T mutation (GeneDetect; Sarasota, FL) at a multiplicity of infection (MOI) of 3,000 unless indicated otherwise. On DIV 7 cells were treated with test samples (6 wells per treatment). All treatments were added by half medium exchange with 2-fold concentrated samples. Each plate contained a negative control (no treatment condition) and positive controls including 150 ng/ml PFF or a non-depleted brain extract. On DIV 18 (11 days post treatment), cells were fixed and stained for either total  $\alpha$ Syn (soluble and insoluble) or insoluble  $\alpha$ Syn or pS129  $\alpha$ Syn. Neurons were fixed with a solution containing 4% paraformaldehyde, 4% sucrose and either 0.1% Triton X-100 (Sigma-Aldrich, Saint Louis, MO) to fix all  $\alpha$ Syn species, or 1% Triton X-100 to fix only the insoluble forms. Fixative was added to the cells for 15 min. Following fixation, cells were washed 3-times with wash buffer containing PBS plus 0.05% tween. Cells were then blocked with 3% BSA and 0.3% triton in PBS for 1-2 hr. Following the blocking step, cells were treated with primary antibody overnight in blocking buffer. Primary antibodies used include anti- $\alpha$  and  $\beta$ -Synuclein (EP1646Y, N-terminal rabbit monoclonal, 1:100 dilution; Millipore, Billerica, MA), anti-pS129  $\alpha$ Syn (81A, mouse monoclonal, 1:1000 dilution, Biolegend; San Diego, CA), anti-pS129  $\alpha$ Syn (MBJR13, ab168381, Abcam, Cambridge, UK) and anti-MAP2 (ab5392, 1:10,000 dilution; Abcam, Cambridge, UK). Following incubation with antibodies, plates were washed 3-times with PBS containing 0.05% tween and then incubated 1 hr with fluorescent-conjugated secondary antibodies. The secondary antibodies used include Alexa Fluor 647 goat anti-mouse IgG (1:500 dilution); Alexa Fluor 488 goat anti-rabbit IgG (1:500 dilution), Alexa Fluor 568 goat anti-chicken IgG (1:500 dilution) and Hoechst, 1:800 dilution. Plates were then washed 3-times for 15 min each with PBS plus 0.05% tween, followed by a final wash in PBS alone.

### *Immunofluorescent analysis*

Images were acquired on ArrayScan<sup>TM</sup> VTi automated microscopy and image analysis system (Cellomics Inc., Pittsburgh, PA, USA) with a  $\times 10$  objective. Imaged plates were analyzed with the High Content Studio 3.0 software package using the neuronal profiling application. Cells were identified with Hoechst fluorescence which defines the nuclear area, and neurites were identified by MAP2 staining. The cell soma was identified by overlapping nuclear and MAP2 staining. The total insoluble  $\alpha$ Syn and total insoluble pS129  $\alpha$ Syn were identified by the fluorescence intensities in two additional channels. Pathological induction is quantified by the total detergent-insoluble pS129 spot intensity co-localized to identified neurites. Fold induction is determined by normalizing to the mean of the negative control wells (untreated cells) within each plate.  $Z'$  values were calculated for each plate based on negative control wells and 150 ng/ml PFF as the positive control. The average  $Z'$  over the course of these experiments was 0.68. A toxicity index was calculated by multiplying the normalized nuclei count, normalized neuronal count, and normalized neurite length for each well. Values were normalized to the respective mean values of the negative control wells. Wells with toxicity scores of  $< 0.6$  were excluded from analysis. Concentration response curves were generated and analyzed using Graphpad Prism (Graphpad Software, Inc. La Jolla, CA).

### *Statistical analysis*

Data expressed as average  $\pm$  SD. Multiple comparisons were performed using one-way ANOVA or two-way ANOVA with Dunnett's post hoc correction as appropriate.

### **Acknowledgements**

We are grateful to the Banner Sun Health Research Institute Brain and Body Donation Program of Sun City, Arizona and the Alzheimer's Disease Core Center, the Morris K. Udall Parkinson's Research Center of Excellence, and the Center for Neurodegenerative Disease Research, Perelman School of Medicine, the University of Pennsylvania (Philadelphia, PA) for providing human brain tissues. The Brain and Body Donation Program is supported by the National Institute of Neurological Disorders and Stroke (U24 NS072026 National Brain and Tissue Resource for Parkinson's Disease and Related Disorders), the National Institute on Aging (P30 AG19610 Arizona Alzheimer's Disease Core Center), the Arizona Department of Health Services (contract 211002, Arizona Alzheimer's Research Center), the Arizona Biomedical Research Commission (contracts 4001, 0011, 05-901 and 1001 to the Arizona Parkinson's disease Consortium) and the Michael J. Fox Foundation for Parkinson's Research. The Alzheimer's Disease Core Center, the Morris K. Udall Parkinson's Research Center of Excellence, and the Center for Neurodegenerative Disease Research are supported by P30 AG10124 and P50 NS053488.

### **Conflict of Interest**

All authors were or are employees of Bristol-Myers Squibb during the data generation and writing of this manuscript.

### **Author Contributions**

JDG, JEM, ND and MKA wrote the manuscript, designed the experiments and oversaw data generation and analysis. NH, CP, LY, LI and YC generated and analyzed the data and aided in the experimental design.

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**Table 1. Antibody binding IC<sub>50</sub> values**

Antibody	Monomer Binding (ng/ml)			PFF Binding (ng/ml)			Mono/PFF IC <sub>50</sub> Ratio
	AVE	SD	n	AVE	SD	n	
MJFR 14642	3658	1981	5	0.51	0.10	5	7173
MJFR1	8.0	1.3	2	29.2	7.0	2	0.27
4D6	54	18	2	2.1	0.2	2	26
4B12	5.6	1.2	2	23.3	3.2	2	0.24
Syn303	51	9	2	109.3	6.6	2	0.47
LB509	62	24	2	1.6	0.1	2	39
EP1646Y	17	0.4	2	52.8	9.9	2	0.32
5C2	>10000		2	>1000		2	NA
14H2	21	5	2	10	3	2	2.1

Average IC<sub>50</sub> for inhibition of binding of antibodies to PFF-coated plates by  $\alpha$ Syn monomers and  $\alpha$ Syn PFF. Antibodies were preincubated in solution with a titration of either  $\alpha$ Syn monomer or PFF and then the amount of free antibody determined and IC<sub>50</sub> calculated. Data represent the average  $\pm$  SD from 2-5 experiments. The monomer to PFF IC<sub>50</sub> ratios are also shown.

**Table 2. Antibody immunodepletion IC<sub>50</sub> values**

Sample	MJFR1 IC <sub>50</sub> (nM)			MJFR14642 IC <sub>50</sub> (nM)			R1/R14642 IC <sub>50</sub> ratio
	AVE	SD	n	AVE	SD	n	
PFF	0.036	0.017	4	0.079	0.054	4	0.5
MSA #7 lysate	7.2	2.2	3	0.72	0.38	5	10.0
MSA #7 pellet	0.52		1	0.038		1	13.7
PD #3 pellet	0.29		1	0.021		1	13.8
DLB+AD #2 pellet	0.44		1	0.062		1	7.1

Average IC<sub>50</sub> for immunodepletion of the induction-competent species from PFF, MSA, PD and DLB+AD brain extracts. Samples were preincubated with a titration of either MJFR1 or MJFR14642, the antibody-complexes removed and the immunodepleted samples evaluated for induction of pS129  $\alpha$ Syn pathology in primary rat hippocampal neurons overexpressing hA53T  $\alpha$ Syn following 11 days of treatment. In some cases, the inducing species was enriched by high-speed centrifugation (pellets) prior to immunodepletion. Data represent the average  $\pm$  SD from 3-5 experiments or the IC<sub>50</sub> from a single experiment. The MJFR1 to MJFR14642 IC<sub>50</sub> ratios are also shown.

## Legends to Figures

**Figure 1. Characterization of pS129  $\alpha$ Syn induction with PFF.** (A) Primary rat hippocampal neurons overexpressing hA53T  $\alpha$ Syn were fixed in the presence of 1% Triton X-100 to extract soluble proteins 11 days (DIV 18) after treatment with 150 ng/ml PFF. Nuclei were stained with Hoechst (blue), dendrites with an anti-MAP2 antibody (green) and pS129  $\alpha$ Syn stained with antibody 81A (white/purple). Neurons treated with PFF exhibited robust pS129  $\alpha$ Syn staining compared to untreated controls. (B) The relative level of pS129  $\alpha$ Syn intensity normalized to untreated controls. Neurons were transduced with either an empty AAV (Empty) or A53T  $\alpha$ Syn AAV (A53T) and then left untreated or treated with 150 ng/ml PFF for 11 days. Treatment with PFF resulted in a significant increase in pS129 signal in both empty AAV and hA53T  $\alpha$ Syn AAV transduced neurons compared to control. Induction was ~20-fold higher in the hA53T  $\alpha$ Syn AAV transduced neurons as compared to empty AAV transduced neurons. Data represents average  $\pm$  SD from n=6 replicates. Statistical analysis based on one-way ANOVA with Dunnett's post hoc test. (\*\*\*\* p < 0.0001). (C)  $\alpha$ Syn PFF concentration response curve generated 11d post treatment in neurons transduced with hA53T  $\alpha$ Syn AAV. A concentration-dependent increase in pS129 signal observed in the absence of cell toxicity. Data are normalized to untreated controls and represents average  $\pm$  SD from n=6 replicate wells. (D) Neurons transduced with hA53T  $\alpha$ Syn AAV were treated for 11 days with either 450 ng/ml PFF or 4500 ng/ml monomer and the induction of pS129 or the effect on cell health measured. Significant induction of pS129 signal observed with PFF but no induction with  $\alpha$ Syn monomer. Data are normalized to untreated controls and represents average  $\pm$  SD from n=6 replicates. Statistical analysis based on one-way ANOVA with Dunnett's post hoc test. (\*\*\*\* p < 0.0001).

**Figure 2. Characterization of pS129  $\alpha$ Syn induction with brain lysates.** (A) Primary rat hippocampal neurons overexpressing hA53T  $\alpha$ Syn were fixed in the presence of 1% Triton X-100 to extract soluble proteins and stained with Hoechst (blue), an anti-MAP2 antibody (green) and the anti-pS129  $\alpha$ Syn antibody 81A (purple) 11 days (DIV 18) after treatment with a 1:300 dilution of lysates from control brain and MSA brain tissue. (B) Quantification of pS129  $\alpha$ Syn intensity normalized to untreated control wells shows robust and statistically significant elevated levels of pS129  $\alpha$ Syn induction for neurons treated with MSA brain tissue lysates as compared to either control, PD or DLB+AD brain lysates. Data represent average  $\pm$  SD from n= 6-12 replicates for untreated controls and 4-6 replicates for brain lysates. <5% of treated wells were excluded from analysis due to toxicity (toxicity index <0.6). Statistical analysis based on one-way ANOVA with Dunnett's post hoc test. (\*\*\*\* p < 0.0001). (C) Brain tissue lysates from 9 different MSA patients showing varying amounts of pS129  $\alpha$ Syn induction 11days post treatment normalized to the untreated control. Data represent average  $\pm$  SD from n=6 replicates. All wells exhibited acceptable viability scores (toxicity index > 0.6). Statistical analysis based on comparison to untreated controls using one-way ANOVA with Dunnett's post hoc test. (\*p<0.5; \*\*\*\* p < 0.0001). (D) MSA brain tissue lysates were fractionated by high-speed centrifugation and the resulting pellet and supernatant (Sup) fractions measured for pS129 induction. Induction is expressed relative to induction with the starting lysate. Data represent average  $\pm$  SD from n=6 replicates.

**Figure 3.  $\alpha$ Syn levels in brain lysates.**  $\alpha$ Syn levels were measured by ELISA in (A) total brain lysates and (B) aggregates isolated from brain lysates following high-speed centrifugation (Pellet).  $\alpha$ Syn levels are expressed relative to a  $\alpha$ Syn monomer standard. Each data point represents individual control, DLB, MSA and PD samples. Group averages are indicated. No significant differences between groups were observed based on one-way ANOVA.

**Figure 4. Induction of pS129  $\alpha$ Syn signal with pellet fractions from brain lysates.** (A) Primary rat hippocampal neurons overexpressing hA53T  $\alpha$ Syn were fixed in the presence of 1% Triton X-100 to extract soluble proteins and stained with Hoechst (blue), an anti-MAP2 antibody (green) and the anti-pS129  $\alpha$ Syn antibody 81A (white) 25-days (DIV 32) after treatment with re-suspended brain lysate pellet fractions from control, MSA, PD, DLB cases. (B) Quantification of pS129  $\alpha$ Syn intensity normalized to untreated controls following 11, 18, 25 or 32 days of treatment. Significant induction of pS129  $\alpha$ Syn can be seen for all three synucleinopathy brain tissue samples. No induction of pS129  $\alpha$ Syn signal was detected with control brain tissue pellets at any timepoint. Data represent average  $\pm$  SD from 3 different brain lysate pellets for each group. Individual brain pellet data are included in supplemental Table S6. Statistical analysis based on comparison to the control group using two-way ANOVA with Dunnett's post hoc test. (\*\*\*)  $p < 0.001$ ; (\*\*\*\*)  $p < 0.0001$ .

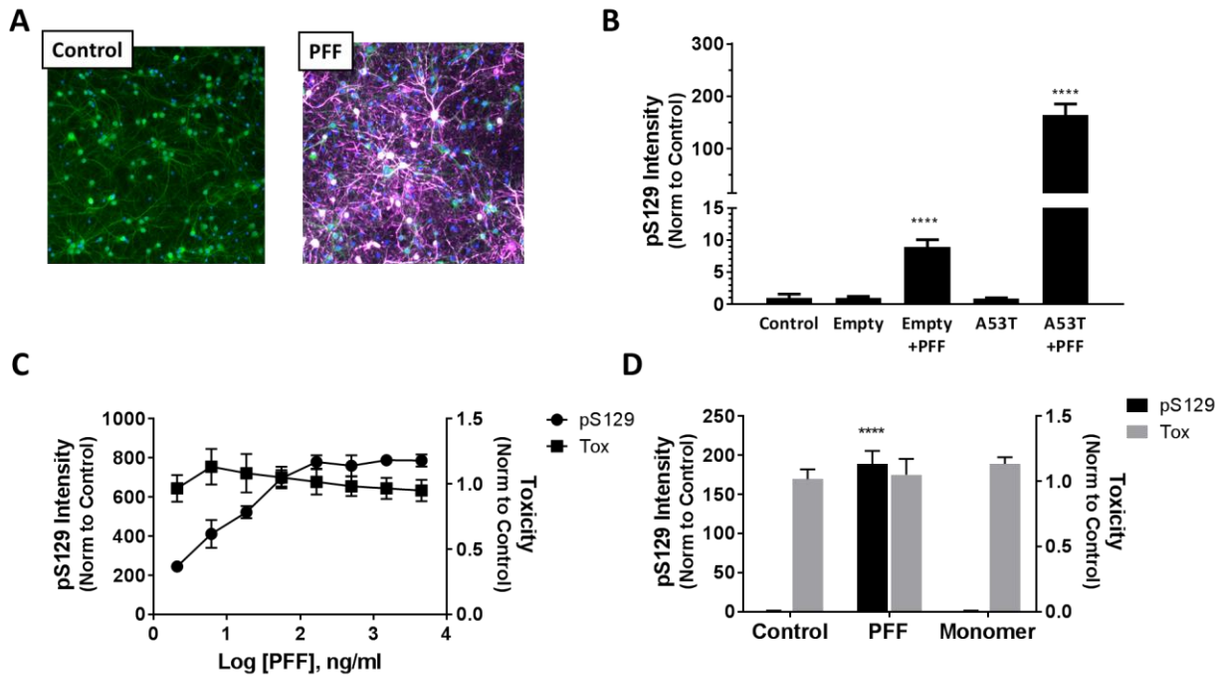
**Figure 5. Binding characterization of  $\alpha$ Syn antibodies.** (A) Amino acid sequence for  $\alpha$ Syn showing the putative binding regions for the commercial  $\alpha$ Syn antibodies tested. \*S129 residue. (B) Example curves for MJFR1 binding to increasing concentrations of either  $\alpha$ Syn monomer (black) or  $\alpha$ Syn PFF (gray). Data represent average  $\pm$  SD from duplicate wells. Summary  $IC_{50}$  values included in Table 1. (C) Example curves for MJFR14642 binding to increasing concentrations of either  $\alpha$ Syn monomer (black) or  $\alpha$ Syn PFF (gray). Data represents average  $\pm$  SD from duplicate wells. Summary  $IC_{50}$  values included in Table 1.

**Figure 6. Immunodepletion of inducing activity from PFF and a MSA brain lysate with  $\alpha$ Syn antibodies.** PFF (15 ng/ml) or MSA brain lysate #7 were immunodepleted with  $\alpha$ Syn antibodies at a concentration of 50 nM. Depleted PFF or brain lysate samples were then tested for induction of pS129  $\alpha$ Syn in primary rat hippocampal neurons overexpressing hA53T  $\alpha$ Syn following 11 days of treatment. Data are normalized to the undepleted PFF or undepleted MSA lysate positive controls and represents the average  $\pm$  SD from 2-6 experiments. Statistical analysis based on comparison to the IgG control group using two-way ANOVA with Dunnett's post hoc test. (\*  $p < 0.05$ ; \*\*  $p < 0.01$ ; \*\*\*\*  $p < 0.0001$ ; ns = not significant).

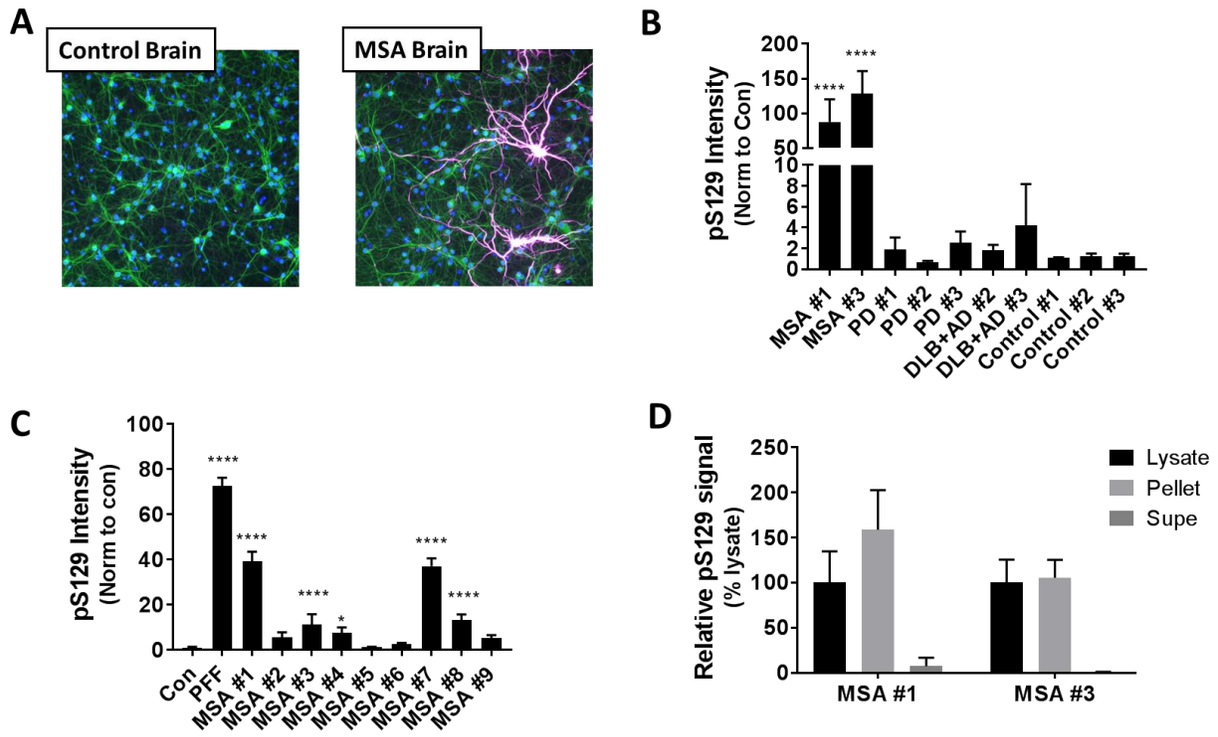
**Figure 7. Immunodepletion of PFF and a MSA brain tissue sample with antibodies MJFR1 and MJFR14642.** Samples were preincubated with a titration of either MJFR1 or MJFR14642, the antibody-complexes removed and the immunodepleted samples evaluated for induction of pS129  $\alpha$ Syn pathology in primary rat hippocampal neurons overexpressing hA53T  $\alpha$ Syn following 11 days of treatment. (A) Example immunodepletion curves for MJFR1 (black) and MJFR14642 (gray) for removing the inducing  $\alpha$ Syn species from 15 ng/ml PFF. Data was normalized to an undepleted PFF positive control. Cumulative curves were generated by combining data from multiple experiments. Data represent average  $\pm$  SD from n=4 experiments. Summary  $IC_{50}$  values are included in Table 2. (B) Example immunodepletion curves for MJFR1 (black) and

MJFR14642 (gray) for removing the inducing  $\alpha$ Syn species from MSA brain tissue lysates #7. Data were normalized to an undepleted MSA brain lysate as positive control. Cumulative curves were generated by combining data from multiple experiments. Data represent average  $\pm$  SD from n=3-6 experiments. Summary IC<sub>50</sub> values are included in Table 2. (C) Example immunodepletion curves for 1 nM MJFR1 (black) and 1 nM MJFR14642 (gray) for removing the inducing  $\alpha$ Syn species from 15 ng/ml PFF samples in the presence of increasing concentrations of  $\alpha$ Syn monomer. Data were normalized to an undepleted PFF positive control. Data represent average  $\pm$  SD from n=3 replicates.

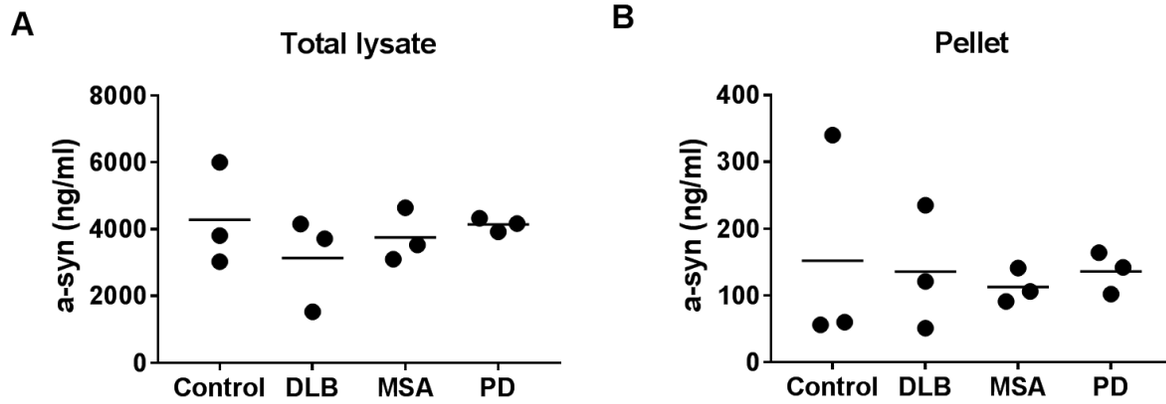
Figure 1.



**Figure 2.**



**Figure 3.**



**Figure 4.**

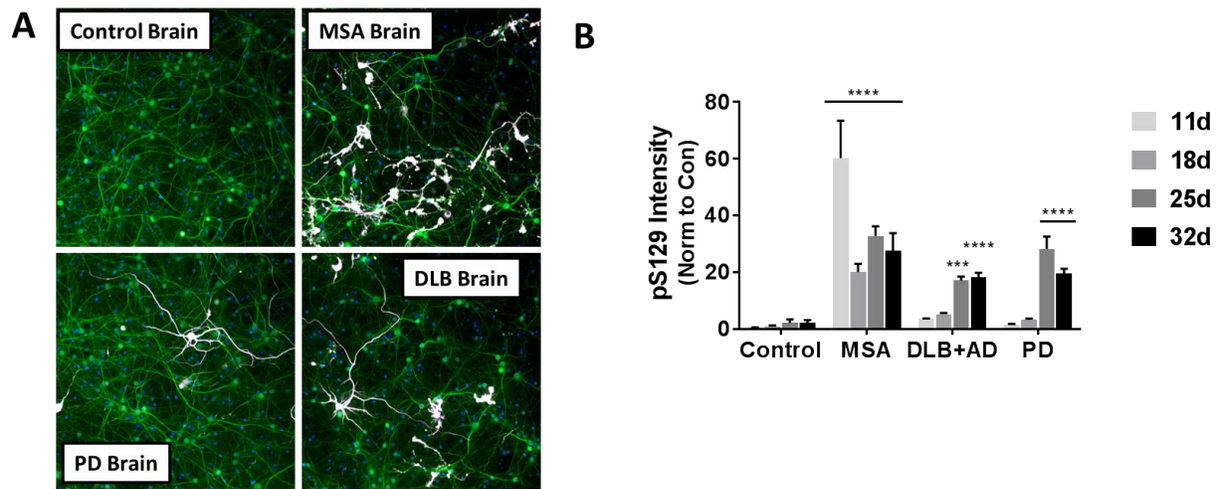




Figure 6.

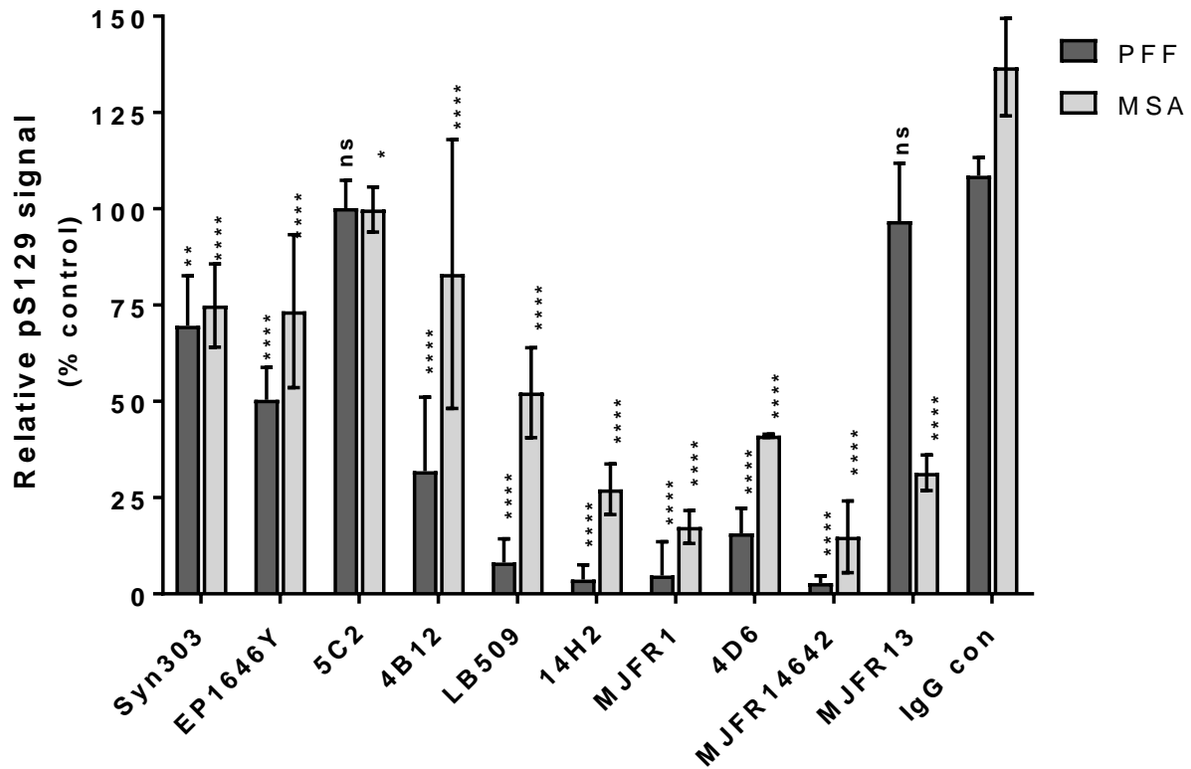


Figure 7.

