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Rho Kinase Inhibition as a Therapeutic for Progressive Supranuclear Palsy and Corticobasal Degeneration

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Introduction

Progressive supranuclear palsy (PSP) and corticobasal degeneration (CBD) are neurodegenerative four-repeat tauopathies with no cure. Mitigating pathogenic tau levels is a rational strategy for tauopathy treatment, but therapeutic targets with clinically available drugs are lacking. Here, we report that protein levels of the Rho-associated protein kinases (ROCK1 and ROCK2), p70 S6 kinase (S6K), and mammalian target of rapamycin (mTOR) were increased in PSP and CBD brains. RNAi depletion of ROCK1 or ROCK2 reduced tau mRNA and protein level in human neuroblastoma cells. However, additional phenotypes were observed under ROCK2 knockdown, including decreased S6K and phosphorylated mTOR levels. Pharmacologic inhibition of Rho kinases in neurons diminished detergent-soluble and -insoluble tau through a combination of autophagy enhancement and tau mRNA reduction. Fasudil, a clinically approved ROCK inhibitor, suppressed rough eye phenotype and mitigated pathogenic tau levels by inducing autophagic pathways in a Drosophila model of tauopathy. Collectively, these findings highlight the Rho kinases as rational therapeutic targets to combat tau accumulation in PSP and CBD.

Key words: autophagy; Drosophila; Rho kinase; ROCK; tau; tauopathy

Significance Statement

Studies of progressive supranuclear palsy (PSP) and corticobasal degeneration (CBD) suggest that mitigating pathogenic tau levels is a rational strategy for tauopathy treatment. In this report, the Rho-associated protein kinases (ROCK1 and ROCK2) are identified as novel drug targets for PSP and CBD. We show that elevated insoluble tau levels are associated with increased ROCK1 and ROCK2 in PSP and CBD brains, whereas experiments in cellular and animal models identify pharmacologic inhibition of ROCKs as a mechanism-based approach to reduce tau levels. Our study correlates bona fide changes in PSP and CBD brains with cellular models, identifies drug targets, and tests the therapeutic in vivo.
Bakhoum et al., 2014). Other studies revealed that mTOR and the mTOR effector p70 S6 kinase (S6K) likely have direct effects on tau synthesis, phosphorylation, and aggregation (An et al., 2003; Tang et al., 2013).

Rho-associated coiled-coil containing protein kinases 1 (ROCK1) and ROCK2 are ubiquitous serine/threonine kinases that share 65% similarity in their amino acid sequences and 92% identity in their kinase domains (Nakagawa et al., 1996). Pharmacologic inhibition of ROCKs can induce protein degradation pathways, including autophagy, in mammalian cells (Bauer et al., 2009; Koch et al., 2014). However, whether ROCKs influence tau synthesis or protein homeostasis is unclear. In this study, observations from PSP and CBD brains are linked to in vitro and in vivo models that provide mechanistic insight, identifying ROCKs as therapeutic targets for PSP and CBD.

Materials and Methods

Antibodies. The following antibodies were used: Tau Dako A0024; phospho-Tau (S202) CP13 Cell Signaling Technology 11834; phospho-Tau (S396) PHF13 Cell Signaling Technology 9632; ROCK1 Abcam ab45171; ROCK2 Abcam ab56661; mTOR Cell Signaling Technology 2972s; phospho-mTOR (S2448) Cell Signaling Technology 2971s; phospho-S6K (T389) Cell Signaling Technology 108D2; S6K Cell Signaling Technology 2708; p62 Abcam ab109012; LC3 Novus NB100-2220; Actin Abcam ab6276.

Human brain tissue preparation and immunoblotting. Postmortem frontal cortex tissues from control, PSP, and CBD cases were selected for comparison from the Emory University Alzheimer’s Disease Research Center brain bank. The PSP and CBD cases in this study underwent extensive neuropathological characterization required for diagnosis based on established criteria (Hauw et al., 1994; Cairns et al., 2007). Cases were matched as closely as possible for age at death, gender, and postmortem interval (Table 1). Sarkosyl-soluble and -insoluble fractions were prepared exactly as described previously (Diner et al., 2014). Protein concentration was determined by bichinchoninic acid method (Pierce). Immunoblotting was performed using standard procedures as described previously (Herskowitz et al., 2011). Fifty micrograms of protein per sample were used for immunoblots. Actin was used as loading control. Images were captured using an Odyssey Image Station (Li-Cor), and band intensities were quantified using Odyssey Application Software version 3.0 (Li-Cor).

For densitometry analysis, control human brain samples were divided randomly into two groups (1–5 and 6–10). Control samples 1–5 were run on an immunoblot with all PSP cases and a separate immunoblot with all CBD cases. In parallel, control samples 6–10 were run on an immunoblot with all PSP cases and a separate immunoblot with all CBD cases. For each immunoblot, densitometry measurements for the control samples were averaged and normalized to an arbitrary unit of 1. Because each control case was used on two immunoblots, two densitometry measurements for each control case were acquired for each control case. The two densitometry measurements for each control sample were averaged, and that average represents the single data point that is represented in the densitometry graphs in Figure 1.

Cell culture, transduction, and transfection. SH-SY5Y human neuroblastoma cells were maintained in MEM Eagle (Lonza) with 10% fetal bovine serum and 1% penicillin/streptomycin. For transductions or transfections, equivalent amounts of cells were plated. DNA plasmid or siRNA transfections were performed using Lipofectamine 2000 (Invitrogen) or DharmaFECT 1 siRNA Transfection Reagent (Dharmacon), respectively, according to the instructions of the manufacturer. Primary cortical neuronal cultures were prepared from embryonic day 17 C57BL/6 mouse embryos and maintained in Neurobasal medium supplemented with 0.8 mM l-glutamine and B27 as described previously (Herskowitz et al., 2013).

DNA constructs, lentivirus, and siRNA. Lentivirus vectors for shRNA expression were constructed and generated as described previously (Herskowitz et al., 2012): ROCK1 shRNA1, 5′-GCAATGACTTACTTACAGA and shRNA2, 5′-CTACAAGTGGTGTAGTTT; ROCK2 shRNA1, 5′-ATCGACAGCATCCTTCTT and shRNA2, 5′-GCAAATCTGTAA TACTCG; and scramble, 5′-GGACATCTCGATCGATA. ROCK2 mutants resistant to shRNA1 sequences were produced as described previously (Herskowitz et al., 2013). To generate ROCK2 K121G, cDNA encoding human ROCK2 with shRNA-resistant mutations was used as a template, and the QuikChange XL Site-Directed Mutagenesis kit (Stratagene) was used: sense primer, 5′-GGTCAAGGAAGTGGTTTTGCTGATCGTTCT TACTAAGTGGTGA, and antisense primer, 5′-TCAACTTACTAAG ACGCCATACGATAACCCTCTGCGATGCC. Constructs were verified by sequencing. For S6K, the following were used: siRNA1, 5′-CAUGUGAC AUUGUGAGAAA and siRNA2, 5′-GGAUGGCAGAUUGUGUUAG.

Chemicals. To inhibit ROCKs, Fasudil (HA-1077; Sigma-Aldrich) or SR4677 (N-[2-[[dimethylamino]ethoxy]-4-(1H-pyrazol-4-yl)phenyl-2,3-dihydro-1,4-benzodioxin-2-carboxamide dihydrochloride; Feng et al., 2008) was dissolved in H2O, and mock was H2O. MG132 (carbobenzoxy-L-leucyl-L-leucyl-L-leucinal), bafilomycin, and rapamycin (Sigma catalog #C2211, #B1793, and #R0395, respectively) were dissolved in 100% dimethylsulfoxide (DMSO) and used at 10 μM, 100 nM, or 5 μM, respectively. Mock was DMSO.

Cell lysate preparation and viability assay. Cells were lysed in PBS plus protease inhibitor mixture (Roche Diagnostics), Halt phosphatase inhibitor mixture (Pierce), and lysis buffer containing 0.5% Nonidet P-40, 0.5% deoxycholate, 150 mM sodium chloride, and 50 mM Tris, pH 7.4. Cell lysate was subjected to a 13,000 rpm spin, and the cleared lysate was designated as the soluble fraction. Sarkosyl-insoluble fractions were prepared as described above for human brain tissue analysis; however, a Dounce homogenizer was not used (Diner et al., 2014). Protein concentration was determined for all fractions as described above. Cell viability was measured using the CellTiter 96 Non-Radioactive Cell Proliferation Assay (Promega) following the instructions of the manufacturer. Absor-
insoluble PSP and CBD fractions, tau immunoreactivity was captured using a DFC295 camera (Leica) with Leica software. Eye phenotypes were examined using a S8APO Scientific). For drug treatments, flies were grown on Formula 4-24 In-Bloomington Drosophila Stock Center (stock 51360). They were main-
tination protocol was used to generate Sarkosyl-soluble and
soluble fraction cell lysate.

Statistical analysis. Statistical analysis was performed using Student’s t test for independent samples. Data are expressed as the percentage of the mean ± SEM with respect to the control. Error bars represent SEM. Prism software (GraphPad Software) was used for correlation analysis and the generation of all graphs, correlation, and p value matrices.

Results

To measure tau levels in the cases under study, a one-step frac-
tionation protocol was used to generate Sarkosyl-soluble and
-insoluble fractions from 10 control, nine PSP, and nine CBD postmortem cortical tissue homogenates (Table 1). All fractions were subjected to SDS-PAGE and subsequent immunoblot using polycyclonal tau antibody generated against amino acids 243–441, containing microtubule-binding 4R (Fig. 1A,B). In Sarkosyl-insoluble PSP and CBD fractions, tau immunoreactivity was observed prominently at 68 and 64 kDa, representing hyperphorylated full-length 4R tau isoforms (Liu et al., 2001). Densi-
tometry analysis indicated that Sarkosyl-insoluble tau level was increased significantly in PSP and CBD cases compared with control brains (Fig. 1E). To quantify Sarkosyl-soluble tau species, immunoreactive bands from 50 to 68 kDa were measured, but changes in soluble tau level were not statistically significant among control, PSP, and CBD samples (Fig. 1F).

Tau degradation can be mediated by autophagy, and mTOR signaling through S6K can initiate autophagic cascades in neu-
rons (Boland et al., 2008; Krüger et al., 2012). Moreover, ROCK1 and/or ROCK2 are proposed to influence autophagy in mamma-
lian cells (Bauer et al., 2009). To assess whether protein levels of these kinases change in PSP or CBD, Sarkosyl-soluble samples were subjected to SDS-PAGE and subsequent immunoblot (Fig. 1G,D). Densitometry analysis indicated that ROCK1, ROCK2, mTOR, and S6K are increased significantly in PSP and CBD cases compared with controls (Fig. 1G–J). Statistical correlation analysis was performed, and correlation coefficients and p values indi-
cate strong positive correlation among ROCK1, ROCK2, mTOR, and S6K in PSP and CBD cases (Fig. 1K–L).

Next, we sought to determine how these kinases may affect tau levels in cellular models. First, SH-SYSY human neuroblastoma cells were transduced with lentivirus expressing ROCK1-targeted, ROCK2-targeted, or scramble shRNA (two shRNA sequences per target), and 96 h later, levels of endogenous tau protein and tau mRNA were measured by immunoblot or quantitative PCR, respec-
tively. Tau protein was reduced 34 and 65%, whereas tau mRNA was reduced ~51 and ~61% in ROCK1 or ROCK2 knockdown cells, respectively, suggesting that lower tau mRNA levels may contribute to tau protein loss (Fig. 2A,B).

Phosphorylation of mTOR at serine 2448 (pmtOR) sup-
presses autophagy induction, and pmtOR level can be regulated by phosphatidylinositol 3-kinase signaling through AKT or via
direct phosphorylation by S6K (Chiang and Abraham, 2005; Holz and Blenis, 2005). Densitometry analysis indicated that S6K was decreased 17 and 45% after RNAi depletion of ROCK1 or ROCK2, respectively, and that ROCK2, but not ROCK1, knockdown decreased pmTOR, whereas total mTOR levels were unchanged (Fig. 2A). Based on this, we hypothesized that depletion of ROCK2 reduced S6K levels, which in turn suppressed pmTOR, stimulating autophagy and tau degradation. To test the first half of the hypothesis, SH-SY5Y cells were transfected with S6K-targeted or scramble siRNA, and 96 h later, tau levels and pmTOR

Figure 2. ROCK2 kinase activity is required for the effects of ROCK2 on tau. Representative immunoblots of kinase knockdown compared with mock or scramble (SCR) controls. Unless otherwise noted, n = 3 biological replicates per condition. A, ROCK1 shRNA (R1, and R11) reduced tau (***p = 0.0003) and S6K (***p = 0.0005). ROCK2 shRNA (R2, and R22) reduced tau, S6K (***p < 0.0001), and pmTOR (normalized to total mTOR; **p = 0.0026) reduced relative tau mRNA level. n = 6 biological replicates per condition. C, S6K knockdown reduced pmTOR and tau (***p < 0.0001). B, R11 (**p = 0.005) and R22 (**p = 0.0026) reduced relative tau mRNA level. *p < 0.05. D, Tau accumulates in bafilomycin-treated samples. SCR, *p = 0.0228; R11, **p = 0.0013; R21, **p = 0.0057. E, ROCK2 expression reversed S6K and tau levels compared with R2, plus vector control (***p = 0.0002, ***p = 0.0004 or L121G (**p = 0.0039, ***p = 0.0001). All data are expressed as the percentage of the mean ± SEM.
were measured by immunoblot (Fig. 2C). Densitometry analysis indicated that tau and pmTOR were reduced 57 and 60%, respectively, in S6K-depleted cells. Notably, ROCK2 levels were unchanged after S6K knockdown (Fig. 2C). These results suggest that ROCK2 knockdown reduces S6K levels and in turn decreases pmTOR and tau. Next, ROCK1-targeted, ROCK2-targeted, or scramble shRNA-expressing cells were treated for 24 h with MG132, a proteasome inhibitor, or 6 h with bafilomycin, an endosomal acidification inhibitor that blocks autophagy, and tau levels were measured by immunoblot. MG132 treatment did not alter tau levels, whereas bafilomycin increased tau significantly compared to scramble shRNA-expressing cells (Fig. 2D). These observations indicate that inhibiting autophagy, but not the proteasome, accumulates tau protein in ROCK1 or ROCK2 knockdown cells.

To determine whether ROCK2 kinase activity mediates the effects of ROCK2 on S6K and tau, site-directed mutagenesis was used to substitute glycine for lysine 121, generating the ROCK2 mutant K121G. Lysine 121 is the ATP-binding site in the kinase domain and is absolutely required for ROCK2 activity (Amano et al., 1997). SH-SY5Y cells were transduced with lentivirus expressing ROCK2-targeted or scramble shRNA, and 72 h later, cells were transiently transfected with the indicated ROCK2 plasmids (resistant to shRNA) or empty vector control. In ROCK2-depleted cells, expression of ROCK2 reversed effects on S6K and tau, whereas K121G did not (Fig. 2E). These results demonstrate that ROCK2 kinase activity is required for the effects of ROCK2 on S6K and tau.

To test whether pharmacologic inhibition of ROCK2 stimulates autophagy and reduces tau levels, primary murine cortical neurons were exposed to SR3677, a characterized ROCK2 small-molecule inhibitor (Feng et al., 2008; Herskovitz et al., 2013). After 16 h, endogenous proteins were measured by immunoblot (Fig. 3A). Densitometry analysis indicated that SR3677 treatment showed a linear dose-dependent decrease in soluble tau, S6K, and pmTOR protein levels, which were consistent with ROCK2 knockdown studies (Fig. 3A). Suppressing mTOR phosphorylation of S6K threonine 389 (pS6K) induces autophagy in neurons; therefore, pS6K was measured, as well as LC3-II, a specific marker for autophagosomes, and p62, a known substrate of autophagy (Kabeya et al., 2000; Björkoy et al., 2005; Boland et al., 2008). SR3677 treatment increased LC3-II, whereas p62 and p56K levels were decreased in a dose-dependent manner (Fig. 3A). Sarkosyl-insoluble fractions were also prepared, and tau was measured by immunoblot. Densitometry analysis indicated that SR3677 treatment showed a linear dose-dependent decrease in Sar-
kosyl-insoluble tau levels that mimicked SR3677-mediated effects on soluble tau (Fig. 3A). Notably, tau immunoreactivity was observed prominently at two bands in Sarkosyl-insoluble fractions, which may represent phosphorylated tau isoforms.

Past studies demonstrated that pharmacologic inhibition of ROCKs can suppress phosphorylation of tau at serine 202 (pTau S202; Past studies demonstrated that pharmacologic inhibition of ROCKs can suppress phosphorylation of tau at serine 202 (pTau S202; Gentry et al., 2012). To determine whether SR3677 reduces endogenous pTau S202 or pTau S396 in primary neurons, immunoblots were performed. Levels of pTau S202 and pTau S396 were normalized to soluble tau, and densitometry analysis indicated that exposure to SR3677 led to a linear dose-dependent decrease in pTau S202 and pTau S396 (Fig. 3A). Quantitative PCR analysis revealed that tau mRNA levels were lower but not significantly reduced after SR3677 treatments compared with mock (Fig. 3B). Autophagy can mediate protein aggregate degradation (Verhoef et al., 2002). Based on this and the findings presented here, we hypothesize that ROCK2 inhibition stimulates autophagy and that this mechanism in combination with lower tau mRNA levels contributes to soluble and insoluble tau protein reduction in neurons.

To assess the clinical relevance of our hypothesis, cortical neurons were exposed for 16 h to increasing doses of Fasudil, a clinically approved ROCK inhibitor (Shibuya et al., 2005). Densitometry analysis indicated that soluble tau protein level was significantly reduced at doses ≥20 μM, whereas insoluble tau and pTau S202 was decreased significantly at 40 μM (Fig. 3C). Notably, LC3-II was increased significantly at 40 μM compared with mock (Fig. 3C). Quantitative PCR analysis showed lower, but not significantly reduced, tau mRNA levels in neurons treated with Fasudil ≥20 μM (Fig. 3D). Notably, cell viability of primary neuronal cultures was not affected by Fasudil or SR3677 treatment (Fig. 3E). Detailed statistical information for densitometry analyses of SR3677 and Fasudil treatments is presented in Table 2.

To determine whether Fasudil can suppress pathogenic tau in vivo, a Drosophila model of tauopathy was used. In this model, the glass multiple reporter element drives expression of a single-copy cDNA encoding the 4R human wild-type tau in the eye (referred to as gl-tau). gl-tau eyes display a rough phenotype in anterior regions, and large autophagic vacuoles accumulate in the retina, suggesting that autophagy is impaired (Jackson et al., 2002; Bakhoun et al., 2014). Drosophila express a single isoform of ROCK (DROCK), and amino acid sequence alignments of DROCK with rodent ROCKs indicate a mismatch that is adjacent to the ATP binding site in the DROCK kinase domain (Mizuno et al., 1999). Based on this, Fasudil was selected to pharmacologically suppress DROCK. Rapamycin induces autophagic pathways and reduces the rough eye phenotype in gl-tau flies (Bakhoun et al., 2014). Therefore, rapamycin was used as a positive control in our experiments. gl-tau flies were grown in the presence or absence of 30 μM Fasudil or 5 μM rapamycin, and immediately after eclosion, adult fly heads were harvested for microscopy and biochemical analyses. The rough eye phenotype was similarly alleviated in Fasudil- and rapamycin-treated cohorts compared with mock (Fig. 4A). In parallel, immunoblot and subsequent densitometry analysis indicated that 4R human tau protein level was reduced 62 and 64% in flies exposed to Fasudil and rapamycin, respectively (Fig. 4B). Additional bio-

Table 2. Immunoblot measurements and statistics for SR3677 and Fasudil treatments

<table>
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<tr>
<th>Compound</th>
<th>Soluble tau</th>
<th>Insoluble tau</th>
<th>p202 tau/tau</th>
<th>p396 tau/tau</th>
<th>LC3-II</th>
<th>p70/mTOR</th>
<th>p62</th>
<th>LC3-II</th>
<th>LC3-II</th>
<th>LC3-II</th>
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<td>5 μM</td>
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<td>78.22</td>
<td>84.94</td>
<td>57.48</td>
<td>127.6</td>
<td>74.47</td>
<td>88.12</td>
<td>81.82</td>
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<td>10 μM</td>
<td>56.48</td>
<td>65.13</td>
<td>56.88</td>
<td>28.39</td>
<td>170.7</td>
<td>50.78</td>
<td>58.2</td>
<td>56.46</td>
<td>87.01</td>
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<td>20 μM</td>
<td>48.82</td>
<td>31.21</td>
<td>29.24</td>
<td>17.03</td>
<td>235.5</td>
<td>40.12</td>
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<td>77.48</td>
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<td>40 μM</td>
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<td>5 vs 10 μM</td>
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<td>20 vs 40 μM</td>
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<td>0.8888</td>
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<td>20 vs 40 μM</td>
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<td>0.5014</td>
<td>0.1629</td>
<td>0.8698</td>
<td>0.3743</td>
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<td>5 vs 20 μM</td>
<td>0.2758</td>
<td>0.0224</td>
<td>0.0104</td>
<td>0.1406</td>
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Measurements and statistics data for Figure 3A, B, and C. Percentage values represent the percentage of the mean with respect to mock. *p values are provided for the indicated comparison. Mock versus 40 μM.
chemical evaluation indicated that rapamycin treatment decreased levels of Ref(2)P, the Drosophila homolog of mammalian p62, and suppressed phosphorylation of Drosophila target of rapamycin (pTOR) in gl-tau flies (Miron et al., 2003; Nezis et al., 2008). Ref(2)P and pTOR were similarly reduced in cohorts exposed to Fasudil (Fig. 4B). Based on these findings, we propose that Fasudil induces autophagic pathways that mitigate pathogenic tau levels in gl-tau flies.

Discussion
Mitigating pathogenic tau levels is a rational strategy for tauopathy treatment, but therapeutic targets with clinically available drugs are lacking. Currently, 29 small-molecule protein kinase inhibitors are used for the treatment of human diseases. Of these, Fasudil and Ripasudil are ROCK inhibitors. Our findings indicate that ROCK1 and ROCK2 are elevated in PSP and CBD brains and that shRNA-mediated depletion or pharmacologic inhibition of ROCK2 stimulates tau protein degradation in ROCK2-coupled with decreased pmTOR, we propose that autophagic mechanisms may also play a role in tau protein loss. Blocking this further, ROCK2 knockdown decreased tau mRNA and protein effects are appreciable by immunoblot. To extrapolate theseuhl RNA-mediated depletion or pharmacologic inhibition of ROCKs reduces tau through combinatorial mechanisms at the mRNA and protein levels.

Whether increased ROCK1, ROCK2, S6K, and/or mTOR contribute to underlying pathogenic mechanisms or manifest during PSP and CBD progression is a critical question. Analysis of PSP and CBD brains at earlier disease stages or complimentary experiments in tauopathy mouse models may help address these questions. It is possible that increased kinase levels may be attributable in part to reactive gliosis in PSP and/or CBD brains (Togo and Dickson, 2002). Hence, it will be important to determine whether changes to ROCKs, S6K, or mTOR are exclusive to specific cell populations and whether enhanced gene transcription, inefficient protein degradation, or a combination of these effects could elevate kinases at the cellular level.

In human neuroblastoma cells, ROCK1 knockdown lowered tau mRNA ~51% but reduced tau protein only 34% (Fig. 2A,B). This disconnect may reflect lengthy tau protein half-life, which is ~20 h in SH-SY5Y cells (Kim et al., 2014). Therefore, the effects of lower tau mRNA may require another 20 h or more before tau protein effects are appreciable by immunoblot. To extrapolate this further, ROCK2 knockdown decreased tau mRNA ~61% and reduced tau protein 65%, suggesting that posttranslational mechanisms may also play a role in tau protein loss. Blocking autophagy, but not the proteasome, rescued tau protein level, and coupled with decreased pmTOR, we propose that autophagy stimulation facilitates tau protein degradation in ROCK2-depleted cells. Similarly, pharmacologic inhibition of ROCK2 using SR3677 in neurons lowered tau mRNA levels equivalently at all doses, whereas tau protein level was depleted in a dose-dependent manner (Fig. 3A,B). Diminished tau protein levels in SR3677-treated neurons were accompanied by protein changes that hallmark autophagy induction, including decreases in pmTOR, pS6K, and p62, as well as increases in LC3-II. These findings support our hypothesis that ROCK2 inhibition promotes autophagic degradation of tau, but how shRNA-mediated depletion or pharmacologic inhibition of ROCKs lowers tau mRNA level is an open question and will require additional investigation. Notably, tau inclusions are found in neurons, astrocytes, and oligodendroglia in PSP and CBD brains, and although strong evidence supports the hypothesis that tau degradation is mediated by autophagy in neurons, it is possible that other mechanisms, including proteasome-mediated degradation, are critical for tau disposal in non-neuronal cells (Wang and Mandelkow, 2012).

Compared with Fasudil, which inhibits ROCK1 and ROCK2 with similar potency, SR3677 offers approximately eightfold higher selectivity of ROCK2 over ROCK1 (Feng et al., 2008). This may account for the more robust SR3677-induced phenotypes and similarity to ROCK2 knockdown studies, yet it is likely that SR3677 also inhibits ROCK1 at the experimental doses used in this study. Moreover, at high doses (40 μM) of SR3677, it is possible that off-target effects, such as direct inhibition of mTOR, could contribute to the observed phenotypes. The off-target hit rate of SR3677 was 1.4%, and the cell-based IC50 for A7r5 smooth muscle cells was calculated to be ~3 nM. However, SR3677 potency in cell culture experiments is likely affected by cell accumulation, cell permeability, and/or ROCK1 and ROCK2 enzyme kinetics (Feng et al., 2008). We propose that suppressing ROCK1 activity likely has therapeutic benefit, but tau reduction is more robustly induced by ROCK2 inhibition through combinatorial effects on tau mRNA levels and autophagy stimulation. Coupled with the clinical availability of Fasudil, our findings raise enthusiasm for ROCK inhibitors as potential therapeutics to combat tau accumulation in PSP and CBD.