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Neurobiology of Disease

Pharmacologic Inhibition of ROCK2 Suppresses Amyloid-β Production in an Alzheimer’s Disease Mouse Model

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Alzheimer’s disease (AD) is the leading cause of dementia and has no cure. Genetic, cell biological, and biochemical studies suggest that reducing amyloid-β (Aβ) production may serve as a rational therapeutic avenue to delay or prevent AD progression. Inhibition of RhoA, a Rho GTPase family member, is proposed to curb Aβ production. However, a barrier to this hypothesis has been the limited understanding of how the principal downstream effectors of RhoA, Rho-associated, coiled-coil containing protein kinase (ROCK) 1 and ROCK2, modulate Aβ generation. Here, we report that ROCK1 knockdown increased endogenous human Aβ production, whereas ROCK2 knockdown decreased Aβ levels. Inhibition of ROCK2 kinase activity, using an isomeric-selective small molecule (SR3677), suppressed β-site APP cleaving enzyme 1 (BACE1) enzymatic action and diminished production of Aβ in AD mouse brain. Immunofluorescence and confocal microscopy analyses revealed that SR3677 alters BACE1 endocytic distribution and promotes amyloid precursor protein (APP) traffic to lysosomes. Moreover, SR3677 blocked ROCK2 phosphorylation of APP at threonine 654 (T654); in neurons, T654 was critical for APP processing to Aβ. These observations suggest that ROCK2 inhibition reduces Aβ levels through independent mechanisms. Finally, ROCK2 protein levels were increased in asymptomatic AD, mild cognitive impairment, and AD brains, demonstrating that ROCK2 levels change in the earliest stages of AD and remain elevated throughout disease progression. Collectively, these findings highlight ROCK2 as a mechanism-based therapeutic target to combat Aβ production in AD.

Introduction

Alzheimer’s disease (AD) is the leading cause of dementia with no effective treatment. There is abundant evidence that the amyloid precursor protein (APP) and its derivative, amyloid-β (Aβ) peptide, play central roles in AD. Aβ accumulates in senile plaques, a pathological hallmark of AD (Masters et al., 1985), and APP gene mutations cause chromosome 21-linked familial AD (FAD; Goate et al., 1991; Murrell et al., 1991). FAD cases display a neuropathological phenotype similar to sporadic AD, and the observation that all known FAD mutations enhance or modify the production of Aβ provided the mechanistic foundation for the amyloid cascade hypothesis (Hardy, 1997). While Aβ alone cannot account for all features of AD, reducing Aβ production or accumulation is central to therapeutic strategies aimed at disease modification. Aβ is generated by sequential proteolytic cleavage of APP by β-site APP cleaving enzyme (BACE) 1 and the subsequent action of γ-secretase. Characterization of the APP Swedish mutation at the BACE1 cleavage site highlighted the potency of modulating this pathway in regulating Aβ generation (Mullan et al., 1992; Citron et al., 1995). Moreover, genetic studies in a population of Icelanders indicated that an APP amino acid substitution that abrogates cleavage by BACE1 protects against AD (Jonsson et al., 2012). Generating viable BACE1 small-molecule inhibitors has met substantial roadblocks due to the size of the BACE1 enzymatic site and the lack of pharmacokinetic efficacy in vivo (Stachel, 2009). However, alternative small-molecule approaches to reduce Aβ production have been explored, including γ-secretase inhibitors/modulators and nonsteroidal anti-inflammatory drugs (NSAIDs; De Strooper et al., 2010).

One way that NSAIDs are proposed to reduce Aβ is through the inhibition of Rho-GTPases and their principal downstream effectors, Rho-associated, coiled-coil containing protein kinase (ROCK) 1 and ROCK2 (Zhou et al., 2003). 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). Exposure to Y-27632, a drug that inhibits ROCK1 and ROCK2 with similar potency (Uehata et al., 1997), reduced brain levels of Aβ42 in an AD mouse model but had an appreciable effect on total soluble Aβ (Zhou et al., 2003). These studies suggested that ROCK inhibition might serve as a rational avenue to curb Aβ production. However, this promising hypothesis has languished somewhat in recent years due to the limited understanding of how the Rho/ROCK pathway modulates Aβ generation and which ROCK isoform is responsible for these effects. In this report, the effects of ROCK1 or ROCK2 knockdown on Aβ generation are defined. We show that, following treatment with a small-molecule inhibitor of ROCK2, APP processing to Aβ was dramatically reduced in cellular and animal models of AD, and mechanisms contributing to the observed effects were identified.

Materials and Methods

Cell culture, transduction, and transfection. SH-SY5Y human neuroblastoma and HEK293 cells were maintained in Eagle’s minimal essential media or DMEM (Lonza), respectively, with 10% fetal bovine serum, and 1% penicillin/streptomycin. Primary cortical neuronal cells were prepared from embryonic day 17 mouse embryos and maintained in neauronal medium supplemented with 0.8 mm 1-glutamine and B27. Cortical tissue was dissected from mouse embryos and trypsin digested. Cells were plated at a density of 100,000 cells/cm² in 12-well dishes that were coated with 100 μg/ml poly-l-lysine. On day 3 postplating, neurons were transduced with indicated lentivirus with a multiplicity of infection of 1. For Aβ studies, at 72 h post-transduction cells were treated with drugs in conditioned media for 16 h. For R1/2A knockdown of ROCK1 or ROCK2, cells were harvested 96 h post-transduction. For transfections or transfections, equivalent amounts of cells were plated, and transfections were performed using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions.

DNA constructs and lentivirus. The plasmid template for lentivirus generation is pLenti, and lentivirus was generated by the Emory University Viral Vector Core. To express human wild-type (WT) APP695, cDNA encoding APP695 was cloned into pLenti using BamH1 and EcoR1 sites. Lentivirus vectors for shRNA expression were constructed as previously described (Herskowitz et al., 2012). The following shRNA sequences were used: human ROCK1 shRNA 1, 5'-GCAATGCTTTCATAGA; human ROCK1 shRNA 2, 5'-CTCAATGTGCTGAGT; human ROCK2 shRNA 1, 5'-ATCCACAAGCTCTTCT; human ROCK2 shRNA 2, 5'-GCAATCTGTATAATGCG; scramble, 5'-GGACATCTCTGATGTA; mouse ROCK2 shRNA 5'-CAATGAGCCTCTTGGAA. To generate ROCK1 and ROCK2 mutants resistant to human shRNA 1 sequences, cDNA encoding ROCK1 (a gift from Dr. Lei Wei, Indiana University School of Medicine, Indianapolis, IN) or ROCK2 (a gift from Dr. Anming Meng, Tsinghua University, Beijing, China) was used as a template, and QuikChange XL Site-directed Mutagenesis kit (Stratagene) was used (Lee et al., 2009; Zhang et al., 2009). For ROCK1, the sense primer was 5'-CTG AAG AAG AAA CAG TAC GCC TCA ATT CAT CAT GTT GTT GTG GAG, and the antisense primer was 5'-AGGGGTGTCGAGTCTTCT CCT AAG CAA ATC ATT GCC TTC TTC TAG CAG. For ROCK2, the sense primer was 5'-GGGGTG TAA AAG AAA CAC ACC ACC CTT TCT TTA AGA ATC AGT, and the antisense primer was 5'-CAG TAC TGA TCA TTA TTA AAG AAA GGG TGT TGT CTT ATT TCC ACC. To generate APP shRNA 6541 (T654A), cDNA encoding APP695 was used as a template, and the QuikChange XL Site-directed Mutagenesis kit was used. For T654A, the sense primer was 5'-CTG AAG AAG AAA CAG TAC GCC TCA ATT CAT CAT GTT GTT GTG GAG, and the antisense primer was 5'-CTC CAC CAC ACC ATG ATG AAT GGA TGC GTA GTG TTT CCT CTT CAG. Constructs expressing human wild-type BACE1 cDNA or BACE1 at serine 498 (S498A) were generated as previously described (Herskowitz et al., 2012). For ROCK inhibition, Rho Inhibitor 1 (RhoI; Cytoskeleton Inc) was used at 1 μg/ml. To inhibit ROCKs, Y-27632 (Calbiochem) was dissolved in H₂O at 50 μM, and Fasudil (HA-1077, Sigma-Aldrich) was dissolved in H₂O and used at the indicated doses. To inhibit ROCK2, SR3677 was dissolved in H₂O and used at indicated doses. To inhibit BACE1, β-secretase Inhibitor IV (BI IV; Calbiochem) was used at 10 μM. The vehicle control (labeled “mock”) was H₂O for all experiments.

Aβ measurements. For cells, media were conditioned for 16 h, and then collected with cells for biochemical analyses. For brain tissue, soluble Aβ was prepared as described below. Aβ was detected using a sandwich ELISA for human Aβ40 or Aβ42 (Millipore) following the manufacturer’s instructions. Plates were read at 450 nm on a Spectra Max Plus plate reader (Molecular Devices).

Cell lysate and mouse tissue preparation and immunoblotting. Cells were lysed in PBS plus protease inhibitor mixture (PIC; 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 μM Tris, pH 7.4 (PBS+PIC+Halt-lysis buffer). Tissue was homogenized (dounce homogenizer) in the PIC+Halt-lysis buffer described above. Homogenized tissue or cell lysate was subjected to a 13,000 rpm spin to remove nuclei and debris. Cleared lysate was used for the indicated biochemical assay. Immunoblotting was performed using standard procedures as described previously (Herskowitz et al., 2011). To load equivalent amounts of lysate per sample, protein concentration was determined by bicinchoninic acid method (Pierce). Actin was used as loading control. Images were captured using an Odyssey Image Station (LI-COR), and band intensities were quantified using Scion Image.

Antibodies. The following antibodies were used: ROCK1, ROCK2, and actin (Abcam); APP and α-secretase cleaved APP (sAPPα; 6E10, Covance); sAPPβ (192wt or 192swe for 5XFAD, Elan Pharmaceuticals); APP for immunocytochemistry (C8; a gift from Dr. Dennis Selkoe, Harvard Medical School, Boston, MA); BACE1 (Abcam); pMLC-Ser19 (Cell Signaling Technology); EEA1 (BD Biosciences); CD63 (BD Biosciences); LAMP1 (Developmental Studies Hybridoma Bank, University of Iowa, Iowa City, IA); ionized calcium-binding adapter molecule 1 (Iba1; Wako); and glial fibrillary acidic protein (GFAP; BIOCARE Medical).

Cell viability assay. Primary neuronal culture cell viability was measured using the CellTiter 96 Nonradioactive Cell Proliferation Assay (Promega) following the manufacturer’s instructions. Absorbance was measured at 570 nm with a reference wavelength at 650 nm. Cell death values were calculated as the percentage of control absorbance.

Stereotoxic injection. Twelve-week-old 5XFAD transgenic mice of either sex were anesthetized with 80 mg/kg ketamine and 8 mg/kg xylazine, and placed in a stereotaxic frame. Eyes were covered in ophthalmic ointment. Scap was sterilized before skull exposure and incision. Unilateral injections of H₂O or SR3677 were given at a rate of 1 μl/3 min into the left and right hemispheres (anteroposterior, −2.1 mm from bregma; mediolateral, ±1.8 mm from midline; and dorsoventral, −1.8 mm from the dura). Following surgery, mice were returned to their home cages, and stereotypic movements were scored. Twenty-four hours later, mice were anesthetized with isoflurane and decapitated. Hemispheres were rapidly harvested and stored at −80°C until further processing for biochemical analyses.

BACE1 activity assay. The BACE1 Activity Detection Kit (Sigma-Aldrich) was used to measure BACE1 activity from cells, brain, and in vitro assays following the manufacturer’s instructions. For cells and brain homogenate, 100 or 500 μg was used, respectively, per sample. All fluorescence readings were taken using a Synergy 2 Alpha microplate reader (BioTek). In vitro kinase assay and liquid chromatography coupled with tandem mass spectrometry. Synthetic BACE1 peptides (ab14766, Abcam) or APP peptides (AnaSpec) were incubated with recombiant ROCK2 protein (Abcam), 200 μM ATP, and 1X NEK PB buffer for 3 h at 30°C. Samples were trypsin digested, and the resulting peptides were analyzed independently by reverse-phase liquid chromatography coupled with tandem mass spectrometry (LC-MS/MS), as previously described (Herskowitz et al., 2010).

Immunocytochemistry and confocal microscopy. Immunocytochemistry and confocal microscopy experiments were performed as previously described (Herskowitz et al., 2012). Briefly, HEK293 cells were plated onto Matrigel extracellular matrix (BD)-coated coverslips in 12-well culture dishes 24 h after transfection. Cells were exposed to SR3677 for 6 h and subsequently fixed for 20 min in 4% paraformaldehyde and rinsed several times with PBS containing 0.5% normal horse serum, 0.5% normal goat
serum, and 0.05% saponin. The cells were blocked and permeabilized for 30 min with PBS containing 5% normal horse serum, 5% normal goat serum, 1% bovine serum albumin, and 0.05% saponin (blocking buffer) plus 0.05% Triton X-100. Cells were rinsed and incubated in primary antibody diluted in blocking buffer overnight at 4°C. The next day, cells were rinsed and incubated for 1 h with both fluorophore-conjugated anti-rabbit and mouse secondary antibodies (Jackson ImmunoResearch) in blocking buffer. Cell nuclei were stained with Hoechst 333258 (Invitrogen) and mounted onto slides using Vectashield (Vector Laboratories). Images were captured using an Olympus IX81 FV1000 inverted confocal microscope. Colocalization analysis was conducted with ImageJ using JACoP (Just Another Colocalization Plugin) version 2.0 according to published methods (Ibolye and Cordelieres, 2006). Briefly, rolling ball background subtraction was applied (radius = 20 pixels) to all images, and a threshold was selected (consistent across treatment and staining). The thresholded Mander’s coefficient was calculated to determine the fraction of red overlapping green (M2). Experiments were performed in triplicate, and the data are presented as the mean ± 1 SD.

Human brain tissue preparation and immunohistochemistry. Postmortem human brain tissue samples were provided by the University of Washington Alzheimer’s Disease Research Center (ADRC) and its Adult Changes in Thought Study, as well as the Johns Hopkins ADRC and the Baltimore Longitudinal Study of Aging (Table 1; Shock et al., 1984). The soluble (S2) fraction was prepared from each case as previously described (Donovan et al., 2012). A sample of 50 μg was used for immunoblots. Immunohistochemical staining procedures were performed as previously described (Herskowitz et al., 2010). Briefly, fixed 50 μm cryosectioned samples of postmortem human frontal cortex were incubated with primary antibodies for 48 h at 4°C. Tissue samples were incubated for 1 h with both fluorophore-conjugated anti-rabbit and mouse secondary antibodies (Jackson ImmunoResearch), and 5 min incubation with bisbenzamidine (Hoechst) was used to label nuclei. For ROCK2, tyramide signal amplification (PerkinElmer) was used.

Statistical analysis. All experiments were performed on at least biological triplicate samples, and statistical analysis was performed using Student’s t test for independent samples. Unless otherwise noted, all 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 all graphs.

Results

Opposing outcomes of ROCK1 or ROCK2 knockdown on endogenous human Aβ levels

To test whether Rho or pan-ROCK inhibition alters production of endogenous human Aβ, SH-SY5Y neuroblastoma cells were exposed to Rho1 (1 μg/ml), pan-ROCK inhibitor Y-27632 (50 μM), or vehicle (H2O, labeled mock), and secreted endogenous Aβ40 was measured by ELISA. Treatment with Rho1 or Y-27632 reduced Aβ40 by 12%, which is consistent with previous findings (Fig. 1A; Zhou et al., 2003). To determine how selective depletion of ROCK1 or ROCK2 influences APP processing to Aβ, SH-SY5Y cells were transduced with lentivirus expressing ROCK1-targeted, ROCK2-targeted, or scrambled shRNA. Ninety-six hours later, levels of endogenous full-length cell-associated APP, secreted sAPPα, and Aβ40 levels were measured by immunoblot or ELISA (sAPPβ as well as Aβ42 were below the limit of detection). Densitometry analysis indicated thatAPP and sAPPα were marginally reduced following RNAi depletion of ROCK2 (Fig. 1B). Secreted Aβ40 levels were increased 65% from ROCK1 knock-down cells but decreased 50% from ROCK2-depleted cells compared with scramble or mock controls (Fig. 1C). These results were confirmed following knockdown of ROCK1 or ROCK2 using a second set of shRNA sequences in human cells (Fig. 1D). To rescue effects on Aβ40 levels due to shRNA-mediated knockdown of ROCK1 or ROCK2, site-directed mutagenesis was used to generate ROCK1 and ROCK2 shRNA-resistant mutants (R1 rescue and R2 rescue, respectively). SH-SY5Y cells were transduced with lentivirus expressing ROCK1-targeted, ROCK2-targeted, or scramble shRNA, and 72 h later cells were transiently transfected with the indicated plasmids. Expression of R1 rescue or

Table 1. Postmortem human brain tissue samples

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Eleven cognitively normal, age-matched, pathology-free controls were compared to 5 asymptomatic AD (aAD) cases, 9 MCI cases, and 16 sporadic AD cases. Case numbers correspond to numbers indicating immunoblot samples. Patient information was not available. F, Female; M, male.
R2 rescue reversed effects on Aβ40 production that were observed following RNAi depletion of ROCK1 or ROCK2, respectively (Fig. 1E). Notably, the opposing outcomes of ROCK1 or ROCK2 knockdown on Aβ levels in SH-SY5Y cells suggest that the net result of inhibiting both kinases is a limited effect, which was observed following treatment with RhoI or Y-27632. These findings suggest that selective inhibition of ROCK2 activity may reduce amyloidogenic processing of APP.

SR3677 reduces sAPPα and Aβ in a dose-dependent manner

To address whether pharmacologic inhibition of ROCK2 activity influences Aβ production, primary murine cortical neurons were transduced with lentivirus expressing wild-type human APP<sub>swp</sub> and secreted Aβ40 and Aβ42 was measured by ELISA following exposure to RhoI, Y-27632, or SR3677, a highly selective ROCK2 small-molecule inhibitor (Feng et al., 2008). Consistent with the findings above, treatment with RhoI or Y-27632 had little to no

Figure 1. Aβ levels were increased following ROCK1 knockdown, but were decreased after ROCK2 knockdown. A, Human SH-SYSY neuroblastoma cells were exposed to RhoI, Y-27632, or vehicle (Mock), and secreted endogenous Aβ40 was measured by ELISA. RhoI or Y-27632 reduced Aβ40 12% compared with mock. ***p < 0.0001. B, SH-SYSY cells were transduced with lentivirus expressing ROCK1-, ROCK2-, or scramble-shRNA, and APP, sAPPα, and Aβ were measured (representative blot shown). ROCK2 knockdown reduced endogenous APP and sAPPα (20% and 32%, respectively). *p = 0.032, **p = 0.0037. C, Aβ40 was increased 65% and decreased 50% from ROCK1 or ROCK2 knockdown cells, respectively. ***p < 0.0001. D, Effects of ROCK1 or ROCK2 knockdown on human endogenous Aβ levels using a second set of shRNA sequences. HeLa293 cells were transfected with plasmids expressing an alternate ROCK1-, ROCK2-, or scramble shRNA sequence, and 96 h post-transfection, secreted endogenous Aβ40 was measured by ELISA. Representative immunoblot confirms shRNA-mediated knockdown of ROCK1 or ROCK2. Consistent with the shRNA experiments in B and C, ROCK1 knockdown increased Aβ40 levels 16%, whereas ROCK2 knockdown decreased Aβ40 levels 22% compared with scramble and mock-transfected controls. ***p < 0.0001. E, ROCK1 and ROCK2 shRNA-resistant mutants rescue effects on endogenous Aβ levels following RNAi depletion of ROCK1 or ROCK2. SH-SYSY cells were transduced with lentivirus expressing ROCK1-, ROCK2-, or scramble-shRNA and subsequently were transfected with plasmids expressing ROCK1 shRNA-resistant mutant (R1 rescue), ROCK2 shRNA-resistant mutant (R2 rescue), or empty vector, respectively. Forty-eight hours post-transfection, secreted endogenous Aβ40 was measured by ELISA. Representative immunoblot confirms shRNA-mediated knockdown of ROCK1 or ROCK2, and the expression of R1 and R2 rescue. Aβ40 levels were comparable among scramble, R1 rescue, and R2 rescue samples (SCR: 99%; R1 shRNA: 120%; R1 rescue: 94%; R1 shRNA vs R1 rescue, *p = 0.02; R2 shRNA: 67%; R2 rescue: 103%; R2 shRNA vs R2 rescue, *p = 0.0204). M, Mock; SCR, scramble. N = 3–6 biological replicates per condition. All data are expressed as the percentage of the mean ± SEM with respect to mock or scramble control.
effect on Aβ, whereas SR3677 dramatically reduced Aβ40 and Aβ42 levels >75% (Fig. 2A). To determine whether SR3677 has a similar effect on endogenous Aβ in human cells, SH-SY5Y neuroblastoma cells were also tested. SR3677 treatment diminished endogenous secreted Aβ40 levels by 68% (Fig. 2B). To further characterize the effects of SR3677 on APP processing to Aβ, primary neurons expressing APP<sub>695</sub> were exposed to increasing doses of SR3677, and levels of full-length cell-associated APP, sAPP<sub>α</sub>, sAPP<sub>β</sub>, and Aβ40 were measured by immunoblot or ELISA. Densitometry analysis indicated that the reduction of APP and sAPP<sub>α</sub> levels was significant at doses ≥20 μM, whereas sAPP<sub>β</sub> and Aβ40 both showed a more striking and linear dose-dependent decrease in levels (Fig. 2 C,E). Fusudil, the only clinically approved ROCK inhibitor, lacks selectivity to ROCK1 or ROCK2 but has been used successfully to treat stroke patients without incidence of severe side effects (Shibuya et al., 2005). In contrast to SR3677, Fusudil treatment reduced Aβ40 levels only 17–22% at concentrations ≥20 μM in primary neurons expressing APP<sub>695</sub> (Fig. 2E). Cell viability of primary neuronal cultures was not affected by SR3677 treatment (Fig. 2F). Furthermore, phosphorylation of myosin light chain 2 serine 19, a ROCK2 substrate (Totsukawa et al., 2000), was reduced in neurons exposed to SR3677 or RNAi-depleted of ROCK2, verifying that SR3677 inhibits ROCK2 kinase activity in neurons (Fig. 2G). These results indicate that ROCK2 inhibition strongly suppresses amyloidogenic processing of APP in a dose-dependent manner. Moreover, the observation that SR3677 mirrored the effects of RNAi depletions of ROCK2 on APP processing to Aβ strongly supports the conclusion that SR3677-induced phenotypes are the consequence of selective ROCK2 inhibition.

**SR3677 reduces BACE1 activity and Aβ in 5XFAD mouse brain**

Without comprehensive knowledge of pharmacokinetic and CNS penetration properties, we sought to examine how acute exposure to SR3677 impacts APP processing to Aβ in the 5XFAD AD mouse model (Oakley et al., 2006). Twelve-week-old 5XFAD mice were treated with 2 mg/kg SR3677 or the same volume of vehicle (H<sub>2</sub>O, labeled mock) by stereotoxic injection into the hippocampus. Twenty-four hours later, the primary injection site (molecular layer of dentate gyrus) and the overlying parietal cortex intimately surrounding the needle track were extracted for biochemical assays. Each hemisphere was harvested separately,site (molecular layer of dentate gyrus) and the overlying parietal cortex (tissue samples; however, quantitative analysis indicated that these changes were not significant compared with mock. These observations suggest that ROCK2 inhibition reduces BACE1 residence in early endosomes and promotes BACE1 traffic to lysosomes. Recent advances show that BACE1 cleavage of APP occurs in early endosomes; there-
Figure 2. SR3677 reduces Aβ40 and Aβ42 in a dose-dependent manner. A, Neurons were transduced with lentivirus expressing APP695 and exposed to Rho1, Y27632 (Y27, 50 μM), SR3677 (50 μM), or H2O (Mock). SR3677 reduced Aβ40 and Aβ42 levels 76% and 78%, respectively. ***p < 0.0001. B, Human SH-SYSY neuroblastoma cells were exposed to SR3677 (50 μM) or vehicle (H2O, labeled mock), and secreted endogenous Aβ40 was measured by ELISA. Treatment with SR3677 reduced Aβ40 68% compared with mock. ***p < 0.0001. C, Neurons expressing APP695 were exposed to SR3677 at indicated doses, and APP, sAPPα, and sAPPβ were measured (representative blot shown). Densitometry analysis indicated changes in APP and sAPPα reached significance at concentrations ≥ 20 μM (APP: 20 μM, 58%; 40 μM, 48%; 60 μM, 46%; sAPPα: 20 μM, 67%; 40 μM, 65%; 60 μM, 57%; APP: mock vs 20, 40, or 60 μM, ***p < 0.0001; sAPPα: mock vs 20, 40, or 60 μM, *p = 0.01). Reduction of sAPPβ was dose dependent (sAPPβ: Mock, 100%; 5 μM, 74%; 10 μM, 40%; 20 μM, 26%; 40 μM, 21%; 60 μM, 8%; Mock vs 10 μM, *p = 0.0341; Mock vs 20 μM, *p = 0.0138; Mock vs 40 μM, *p = 0.013; Mock vs 60 μM, ***p = 0.0089). D, Densitometry analysis in C was used to calculate the ratio of sAPPα to APP. At higher concentrations of SR3677, the ratio of sAPPα to APP increases. E, Neurons expressing APP695 were exposed to SR3677 or Fasudil at indicated doses, and Aβ40 was measured by ELISA. SR3677 reduces Aβ40 in a dose-dependent manner (mock, 100%; 5 μM, 88%; 10 μM, 70%; 20 μM, 56%; 40 μM, 45%; 60 μM, 26%; Mock vs 5 μM, ***p = 0.0088; 5 vs 10 μM, ***p = 0.0001; 10 vs 20 μM, ***p < 0.0001; 20 vs 40 μM, ***p = 0.0016; 40 vs 60 μM, *p = 0.0301). Fasudil reduces Aβ40 marginally at doses ≥ 20 μM (mock, 100%; 5 μM, 107%; 10 μM, 99%; 20 μM, 83%; 40 μM, 81%; 60 μM, 78%; Mock vs 20 μM, *p = 0.0151; Mock vs 40 μM, ***p = 0.0092; Mock vs 60 μM, ***p = 0.0022). F, Primary murine cortical neurons were exposed to vehicle (H2O, labeled mock) or the indicated doses of SR3677 for 16 h. No gross changes in cell viability were observed under the indicated SR3677 concentrations. G, To verify that SR3677 can reduce ROCK2 kinase activity in neurons, primary murine cortical neurons were treated with SR3677 (20 μM) or vehicle (mock) and harvested 16 h later to examine phosphorylation of myosin light chain 2 serine 19 (pMLC-Ser19), a ROCK2 substrate (Totsukawa et al., 2003), by immunoblot. In parallel, neurons were transduced with lentivirus driving expression of ROCK2-targeted or scramble shRNA, and 96 h later neurons were harvested. Immunoblot analyses using a phospho-specific antibody against Ser19 revealed that exposure to SR3677 or RNA depletion of ROCK2 similarly reduced levels of pMLC-Ser19, supporting that SR3677 inhibits ROCK2 kinase activity in neurons. Actin was used as a loading control. M, Mock; SCR, scramble. N = 3–6 biological replicates per condition. All data are expressed as the percentage of the mean ± SEM with respect to mock.
before, it is possible that the observed reduction in sAPPβ levels after SR3677 treatment is due in part to redistribution of BACE1 to LAMP1-positive compartments (Sannerud et al., 2011). Notably, BACE1 protein levels are not altered in the presence of SR3677, suggesting that BACE1 may be rescued from lysosomal degradation (Figs. 3A, 4C; Koh et al., 2005).

In addition, endogenous APP traffic to lysosomes was evaluated in the presence or absence of SR3677 by immunofluorescence and confocal microscopy. Similar to BACE1, APP localized more strongly to LAMP1-positive compartments following SR3677 treatment (Fig. 5B, D). Notably, the antibody (C8) recognizes amino acids 676–695; therefore, our APP staining could reflect full-length APP as well as all C-terminal fragments (CTFs; Selkoe et al., 1988). Enhanced traffic of APP and/or APP CTFs to lysosomes may account for the reduction in APP protein level as well as Aβ levels that was observed following exposure to increasing concentrations of SR3677 (Fig. 2C).

**ROCK2 phosphorylates APP at T654, and T654 is critical for Aβ generation in neurons**

Phosphorylation of the APP cytoplasmic tail is reported to play a role in APP processing to Aβ, and [γ-32P] metabolic labeling studies revealed that the ROCK kinase domain can phosphorylate APP (Lee et al., 2003; Amano et al., 2010). Based on these observations, we hypothesized that ROCK2 phosphorylates the cytosolic domain of APP and that this action influences Aβ generation. To identify putative sites of phosphorylation, in vitro kinase assays were performed whereby synthetic peptides comprising the full-length APP cytoplasmic tail were incubated in the presence or absence of recombinant ROCK2 protein. Peptides were independently analyzed by LC-MS/MS, and MS/MS spectra were collected and searched as described above. Our LC-MS/MS data identified a single APP phosphopeptide corresponding to the full-length APP cytoplasmic tail were incubated in the presence or absence of SR3677 (Fig. 3D). These findings indicate that T654, and possibly phosphorylation of T654 by ROCK2, is critical for APP processing by BACE1 and the generation of Aβ. Similar to T654A, a larger reduction in Aβ40 compared with Aβ42 was observed in 5XFAD brain following exposure to SR3677 (Fig. 3B). These distinct changes in Aβ isoforms may allude to a role for T654 phosphorylation in γ-secretase processing of APP C-terminal fragments.

The results above show that ROCK2 inhibition suppresses BACE1 activity and blocks ROCK2 phosphorylation of APP T654 in vitro; thus, we hypothesized that coupling SR3677 treatment with T654A would synergistically reduce Aβ levels. To test this, WT- or T654A-expressing neurons were exposed to a low dose of SR3677 (20 μM), and Aβ levels were measured by ELISA. Substantial decreases in Aβ40 and Aβ42 levels were observed from neurons expressing T654A after treatment with SR3677, indicating an additive effect of ROCK2 inhibition with mutagenesis of T654 (Fig. 6D). These observations suggest that inhibition of ROCK2 activity reduces Aβ production through at least two independent mechanisms.

**ROCK2 protein level is elevated in asymptomatic AD, MCI, and AD brains**

ROCK2 is a cytosolic kinase, and the accumulation of Aβ may cause intracellular cytosolic proteins to lose solubility in AD, effectively altering their function (Xu et al., 2013). To assess whether the ROCK2 protein level is changing in AD brain, frontal cortex tissue homogenates were prepared from 16 AD and 11 age-matched pathology-free control cases (Table 1). Homogenates were subjected to SDS-PAGE and subsequent immunoblot (Fig. 7A–C). Densitometry analysis indicated that ROCK2 levels were elevated in AD brains compared with those of controls (Fig. 7D). To determine whether changes in ROCK2 occur early in disease progression, nine mild cognitive impairment (MCI) and five asymptomatic AD brains were analyzed. MCI is considered a prodromal phase of AD, whereas asymptomatic AD is hypothesized to represent early disease stages between the first appearance of AD neuropathology and the onset of clinical symptoms (Driscoll and Troncoso, 2011; Sperling et al., 2011). ROCK2 protein levels were increased in MCI brains, and asymptomatic AD cases showed markedly elevated ROCK2 levels compared with controls. To determine whether changes in ROCK2 level are due to the accumulation of microglia or astrocytes in AD brain, immunohistochemical analysis was performed on four AD and four

Figure 3. SR3677 reduces Aβ in 5XFAD mouse brain. SR3677 or H2O (mock) was stereotaxic injected into 5XFAD mouse brain. An injection was given on the left and right hemispheres. Each hemisphere was processed separately, and assay measurements for left and right hemispheres were averaged to yield one data point per brain (four mice per condition). Twenty-four hours after injection, brain homogenates were prepared for biochemical assays. A, Representative immunoblot shown. APP and sAPPβ were reduced 32% and 77%, respectively, from brains injected with SR3677 compared with mock. ***p = 0.001; **p = 0.0036. B, Aβ40 and Aβ42 levels were reduced 53% and 33%, respectively, from SR3677 injected brains compared with mock. ***p < 0.0001, *p = 0.0105. All data are expressed as the percentage of the mean ± SEM with respect to mock.
control cases. ROCK2 staining appeared neuronal and was elevated in cortical layers I–II of AD brains compared with controls. ROCK2 colocalization with cells positive for GFAP or Iba1, protein markers for mature astrocytes and microglia, respectively, was minimal (Fig. 7E). Collectively, these results demonstrate that ROCK2 levels are increased in the earliest stages of AD and remain elevated throughout disease progression. On the basis of these findings, coupled with our pharmacologic studies, we propose that ROCK2 is a mechanism-based therapeutic target to combat Aβ production for the treatment of AD progression.

Discussion

Effective therapies that combat the progression of AD do not currently exist. However, studies in animal models of AD provide strong evidence that reducing Aβ production may ease symptoms and slow disease progression. BACE1 cleavage of APP is the rate-limiting step in the generation of Aβ; therefore, constraining this step would decrease Aβ production. While developing small molecules to inhibit BACE1 has met with substantial challenges, the findings herein provide a novel avenue to suppress BACE1...
activity and curb Aβ generation. Our results show that pharmacologic inhibition of ROCK2 activity diminishes sAPPβ production as well as Aβ levels, and the following plausible mechanisms are provided: (1) suppression of BACE1 enzymatic action; (2) altering BACE1 endocytic traffic; and (3) modifying phosphorylation of the APP cytoplasmic tail at T654.

Directly inhibiting the enzymatic activity of the amyloidogenic secretases is fraught with caveats, pertaining to effects on unrelated substrates (De Strooper et al., 2010). One approach to circumvent these issues is to modulate secretase activities as opposed to directly inhibiting them. This approach is being explored for γ-secretase modulators. An alternative strategy is to
regulate post-translational modification of APP in a way that deters amyloidogenic processing but protects α-secretase cleavage of APP. Our findings in neurons indicated that with low-dose (≤10 μM) treatments of SR3677, APP and sAPPα levels were not significantly altered, whereas sAPPβ production was diminished (Fig. 2C). At higher doses (≥20 μM), APP and sAPPα levels were marginally lower, but the ratio of sAPPα to APP remained ≥1:1 (Fig. 2D). Similarly, in 5XFAD mouse brain, exposure to SR3677 slightly reduced APP levels but depleted sAPPβ (Fig. 3A). Furthermore, immunofluorescence and confocal microscopy studies revealed that SR3677 promotes APP traffic to LAMP1-positive compartments (Fig. 5D). Under these conditions, it is possible
that the observed decrease in APP may be accounted for by (1) enhanced APP recycling and subsequent processing by α-secretase, (2) increased APP traffic to degradative pathways, or (3) a combination of both.

We hypothesize that ROCK2 phosphorylates APP at T654 in vivo, and that inhibiting this phosphorylation contributes to the decrease in sAPPβ and Aβ that is observed following treatment with SR3677. Although the functional role of T654 is unknown, T654 is phosphorylated in brain and comprises the YTSI motif, which mediates basolateral sorting of APP (Haass et al., 1995). Nuclear magnetic resonance studies revealed that phosphorylation of T654 induces conformational changes that may stabilize binding of cytosolic factors to the APP C-terminal tail (Ramelot and Nicholson, 2001). Changes in the affinity and/or specificity of APP protein–protein interactions could contribute to the observed effects on Aβ generation. For example, the lipoprotein receptor LR11, or SorLA, complexes with APP through luminal as well as cytosolic interactions and mediates APP traffic; however, whether APP phosphorylation plays a role in this interaction is unknown (Spoelgen et al., 2006). Notably, LR11 and ROCK2 associate in human brain, and it is possible that binding to LR11 facilitates the effects of ROCK2 on APP processing (Herskowitz et al., 2011). Without a phospho-specific antibody, evaluating whether or not LR11 is critical for ROCK2 phosphorylation of APP T654 in cells is a challenging question. In addition, LR11 interacts with BACE1 (Spoelgen et al., 2006); therefore, the association of ROCK2 with LR11 may be important for the observed effects on BACE1 enzyme action following inhibition of ROCK2 activity.

The opposing outcomes of ROCK1 or ROCK2 knockdown on Aβ levels suggest that the net result of inhibiting both kinases is a limited effect, which was observed following treatment with Rhod or Y-27632 (Figs. 1A, 2A). These findings highlight that the effects of ROCK1 and ROCK2 on Aβ production are likely independent of each other, and we hypothesize that selective inhibition of ROCK1 would increase Aβ generation. Previous work demonstrated that the expression of a constitutively active ROCK1 mutant (CA-ROCK1) reduced sAPPα levels, whereas a dominant-negative ROCK1 mutant (DN-ROCK1) increased sAPPα levels in N2a murine neuroblastoma cells overexpressing APP with Swedish mutations (Pedrini et al., 2005). Additional experiments revealed that the expression of CA-ROCK1 or DN-ROCK1 had no effect on Aβ production in human cells (Leuchtenberger et al., 2006). In the present study, we show that RNAi depletion of ROCK1 in human neuroblastoma cells does not significantly alter endogenous full-length APP or sAPPα levels, but substantially increases Aβ40 production (Fig. 1B–E). Deeper analysis of β- and/or γ-secretase processing of endogenous APP following RNAi depletion of ROCK1 may help elucidate how ROCK1 influences the generation of Aβ.

Currently, Fasudil is the only clinically available ROCK inhibitor, but it lacks selectivity to ROCK1 or ROCK2 (Shibuya et al., 2005). Our comparison of Fasudil to SR3677 indicates that an isooform-selective ROCK2 inhibitor may be a more effective approach to reduce amyloidogenic processing of APP in AD (Fig. 2E). The kinase inhibitor profile for SR3677 included an extensive panel of kinases and other non-kinase-relevant enzymes and receptors (Feng et al., 2008). Of 353 kinases, SR3677 hit only 5
kinases with inhibition >50%. Moreover, SR3677 demonstrated an approximately eightfold selectivity of ROCK2 over ROCK1. With an off-target hit rate of 1.4%, SR3677 is considered one of the best known ATP-competitive kinase inhibitors (Karaman et al., 2008). Clearly, more work is needed before isoform-selective ROCK inhibitors reach the clinic; however, the potential for these compounds to combat Aβ production for the treatment of AD raises considerable enthusiasm.

References
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