Response of a Neuronal Model of Tuberous Sclerosis to Mammalian Target of Rapamycin (mTOR) Inhibitors: Effects on mTORC1 and Akt Signaling Lead to Improved Survival and Function

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Tuberous sclerosis (TSC) is a hamartoma syndrome attributable to mutations in either TSC1 or TSC2 in which brain involvement causes epilepsy, mental retardation, and autism. We have reported recently (Meikle et al., 2007) a mouse neuronal model of TSC in which Tsc1 is ablated in most neurons during cortical development. We have tested rapamycin and RAD001 [40-O-(2-hydroxyethyl)-rapamycin], both mammalian target of rapamycin mTORC1 inhibitors, as potential therapeutic agents in this model. Median survival is improved from 33 d to more than 100 d; behavior, phenotype, and weight gain are all also markedly improved. There is brain penetration of both drugs, with accumulation over time with repetitive treatment, and effective reduction of levels of phospho-S6, a downstream target of mTORC1. In addition, there is restoration of phospho-Akt and phospho-glycogen synthase kinase 3 levels in the treated mice, consistent with restoration of Akt function. Neurofilament abnormalities, myelination, and cell enlargement are all improved by the treatment. However, dysplastic neuronal features persist, and there are only modest changes in dendritic spine density and length. Strikingly, mice treated with rapamycin or RAD001 for 23 d only (postnatal days 7–30) displayed a persistent improvement in phenotype, with median survival of 78 d. In summary, rapamycin/RAD001 are highly effective therapies for this neuronal model of TSC, with benefit apparently attributable to effects on mTORC1 and Akt signaling and, consequently, cell size and myelination. Although caution is appropriate, the results suggest the possibility that rapamycin/RAD001 may have benefit in the treatment of TSC brain disease, including infantile spasms.

Key words: tuberous sclerosis; TSC; TSC1; TSC2; rapamycin; RAD001

Introduction

Tuberous sclerosis complex (TSC) is a clinically devastating neurocutaneous syndrome in which benign tumors termed hamartomas develop in multiple organ systems (Gomez et al., 1999; Curatolo, 2003; Crino et al., 2006). Neurological manifestations are a predominant clinical feature and include early-onset epilepsy, mental retardation, developmental delay, and autism (Ess, 2006; Marcotte and Crino, 2006; Holmes and Stafstrom, 2007; Winterkorn et al., 2007). Most neurological symptoms are thought to be attributable to the occurrence of cortical tubers, which typically form at the gray–white matter junction. The laminar structure within these lesions is severely disrupted with occurrence of poorly differentiated giant cells, dysplastic neurons and astrocytes, and a variety of reactive cells (Huttenlocher and Wollmann, 1991; Ess, 2006). The number and location of cortical tubers, as well as more generalized cortical abnormalities, and the timing of onset and duration of infantile spasms all appear to have some relationship to the severity of the neurological manifestations that are seen in TSC patients (Doherty et al., 2005; Marcotte and Crino, 2006; Winterkorn et al., 2007).

TSC is attributable to inactivating mutations in either the TSC1 or the TSC2 gene and segregates in an autosomal dominant manner (Cheadle et al., 2000). TSC1 mutations account for 20–25% of all mutations identified, whereas TSC2 mutations account for the remainder. TSC1 disease is less severe than TSC2 disease in multiple respects (Jones et al., 1999; Dabora et al., 2001; Sancak et al., 2005; Au et al., 2007; Winterkorn et al., 2007), and this appears to be attributable to a reduced frequency of second-hit events in the TSC1 gene compared with the TSC2 gene (Dabora et al., 2001).
The TSC1 and TSC2 proteins form a relatively tight stoichiometric complex in cells, which functions in an ancestrally conserved signaling pathway that regulates the state of activation of mammalian target of rapamycin (mTOR) and thereby cell growth (Goncharova et al., 2002; Kwiatkowski et al., 2002; Inoki et al., 2005; Tee and Blenis, 2005; Avruch et al., 2006; Sabatini, 2006). Loss of either TSC1 or TSC2 leads to elevated rhabdomyosarcoma (Rho-motor) or the mann–Whitney U test, respectively (Sato et al., 2000; Clapp et al., 2007). Scores for these traits were compared using the Mann–Whitney U test.

Materials and Methods

Mouse procedures. As described previously (Meikle et al., 2007), mice with these alleles began in a mixed strain background (129S4/SvJae, C57Bl/6, and CBA) but have been maintained as an inbred population in our colony for > 3 years. Mice bearing the SynICre allele were the generous gift of Jamey Marth (University of San Diego, La Jolla, CA) (Zhu et al., 2001). Mice were generated through matings between parents of Tsc1cwSynICre or Tsc1cc SynICre to denote the conditional (floxed), mouse to denote mice with the genotype Tsc1<sup>e</sup> SynICre<sup>+</sup> or Tsc1<sup>e</sup> SynICre<sup>+/+</sup> or Tsc1<sup>t</sup> SynICre<sup>+</sup> or Tsc1<sup>t</sup> SynICre<sup>+/+</sup>. As reported previously, mice with these two genotypes have identical phenotypes and survival (Meikle et al., 2007). Controls usually had the genotype Tsc1<sup>e</sup> SynICre<sup>−</sup> or Tsc1<sup>e</sup> SynICre<sup>−/−</sup>, but some were Tsc1<sup>e</sup> SynICre<sup>−</sup>, Tsc1<sup>e</sup> SynICre<sup>−/+</sup>, or Tsc1<sup>t</sup> SynICre<sup>−</sup>. littermates were used whenever possible, but all mutants and controls were closely related from this inbred colony.

All procedures were performed in accordance with the Guide for the Humane Use and Care of Laboratory Animals, and the study was approved by the Animal Care and Use Committee of Children's Hospital. Individual mice were killed when weight loss of 20%, greatly reduced movement, or other signs of morbidity were seen.

Neurologic assessment was performed by an observer blinded to genotype and treatment status of the mice. This included the following: assessment of hind leg clasp behavior when suspended by the tail, scored as absent (0) or present (1); whole-body tremor assessed by placing the palm of the hand on the back of the mouse, scored on a scale from absent (0) to severe and persistent (5); kyphosis, scored as absent (0) or present (1); and tail position observed during a 3 min interval during which the animal was allowed to walk freely in a confined space, scored as normal (0), held horizontal (1), held above horizontal (2), or Straub position (elevated highly, 3). Scores for these traits were compared using the Fisher’s exact test (clasp, kyphosis) or the Mann–Whitney U test (tremor, tail position) in Prism (GraphPad Software).

Mice were anesthetized, and total body weight was determined before they were killed. The brains were rapidly removed, snap frozen in liquid nitrogen, and stored as half-brains at −80°C until use. Half-brain weights were measured in milligrams accurate to ±0.1 mg. Brain weight to body weight ratios were calculated, and measurements were compared using the Mann–Whitney U test.

DNA analysis. DNA was prepared from mouse toes/tails by standard procedures for genotyping. Genotyping at the Tsc1 gene was performed using a four-primer system that allows simultaneous analysis of the c, w, and − alleles, followed by agarose gel electrophoresis (Meikle et al., 2005). Primers that amplify a 300 bp portion of the cre recombinase gene were used to assess the presence of the SynICre allele (Meikle et al., 2005).

Antibodies. Antibodies used were as follows: Tsc2 (C20), Akt (C20), extracellular signal-regulated kinase 2 (ERK2) (K231), pCofilin(S3) from Santa Cruz Biotechnology; pS6(S240/244), pS6(S235/236), Tsc1, pAKT(S473), S6, Cofilin, glycogen synthase kinase (GSK) 3β, and pGSK3β(S9) from Cell Signaling Technology; neuronal-specific nuclear protein (NeuN) (MAB 377), neurofilament (MAB4245), MBP (AB980), high- and medium-molecular-weight neurofilament from Millipore Bioscience Research Reagents; nonphosphorylated neurofilament (SMI311), phosphorylated neurofilament (SMI312), and MBP (SMI399) from Sternberger Monoclonals.

Immunoblotting. Mice were killed, and brains were harvested as above. Each frozen half brain was homogenized in 5% volume TBSV lysis buffer (in mM: 20 Tris-Cl, pH 7.5, 140 NaCl, 10 NaF, 1 Na<sub>3</sub>VO<sub>4</sub>, and 1 EDTA) with 0.1% Triton X-100, 10 mg β-glycerophosphate, 1% phosphatase inhibitor cocktail 1 (Sigma), and 1% phosphatase inhibitor cocktail 2 (Sigma). Any solid insoluble matter was removed by centrifugation at 15,000 × g at 4°C for 10 min, and the supernatant was mixed with Laemmli’s SDS-sample buffer. Samples were put through a 22 gauge needle, boiled, and centrifuged before loading. Proteins were separated by electrophoresis on 4–12% Bis-Tris gels (Novex) and transferred onto nitrocellulose membranes. Coomassie staining was performed to confirm that the samples were loaded equally. The membranes were blocked in 5% nonfat dry milk in PBS, pH 7.4, with 0.1% Tween 20 (PBS–Tween) for 1 h at room temperature. Primary antibodies were diluted in blocking solution, and membranes were incubated overnight at 4°C or 1 h at room temperature. The primary antibody was removed, and the blots were washed in PBS–Tween and then incubated for 1 h at room temperature in horseradish peroxidase-conjugated secondary antibodies (Santa Cruz Biotechnology). Reactive proteins were visualized using SuperSignal West Pico chemiluminescence reagent (Pierce) and exposure to x-ray film (BioMax MR; Eastman Kodak). All immunoblots shown in one row of a figure are from the same gel–blot exposure.

Histological preparation, immunohistochemical staining, and microscopy. For immunohistochemistry, mice at ages postnatal day 21 (P21) to P100 were anesthetized, and transcardiac perfusion was performed using PBS, followed by 4% paraformaldehyde (PFA) in PBS (Boston BioProducts). Whole brains were then removed and postfixed overnight at 4°C in 4% PFA solution. Fixed brains were then cryoprotected in 30% sucrose in PBS for 1–14 d at 4°C, and coronal sections were cut at 50 µm using a Microm K400 Fast Freezing Unit and HM 450 Sliding Electronic Microtome and were then stored in PBS at 4°C until use (Meikle et al., 2007). Free-floating sections were incubated in blocking solution (PBS, 0.1% Triton X-100, and 5% normal goat serum) for 1 h at room temperature, followed by incubation with primary antibody overnight at 4°C. After three 20-min PBS washes, sections were incubated in secondary antibody solution containing Alexa Fluor 568 goat anti-rabbit/anti-mouse IgG/ IgM or Oregon Green 488 goat anti-rabbit/anti-mouse IgG (Invitrogen), for 1 h at room temperature. After two washes in PBS, for some sections, 0.5 µg/ml Hoechst 33258 was added to each well for 5 min at room temperature, and the sections were then washed twice in PBS, mounted onto positively charged slides, and coverslipped with antifade medium (Fluoromount-G; Southern Biotechnology Associates). Control sections were done with omission of one or both primary antibodies, adding both secondary antibodies, to establish specificity. Slides were viewed on a Nikon TE2000-E inverted microscope. Illumination wavelength, intensity, and exposure times were selected using a dual filter-wheel apparatus containing an electronically controlled shutter and filters for the different proteins.
fluorophore wavelengths (Lambda 10-2; Sutter Instrument). Digital images were captured using an Orca-II ER cooled CCD camera (Hamamatsu) and MetaMorph (Molecular Devices). Digital images were then processed and analyzed using Adobe Photoshop (Adobe Systems), including pseudocolorization.

Standard histology sections were prepared after Bouins fixation at room temperature. After hematoxylin and eosin staining, slides were viewed on a Nikon Eclipse E400 microscope, and images were captured using Spot Software version 4.0.5.

For all histological and immunostaining observations, a minimum of three (range of 3–12) pairs of mutant and control mice were examined from stereotactically matched brain sections.

Confocal images were captured using a Zeiss LSM510 META two-photon confocal microscope using 20× and 63× objectives. For cell size determination and other images, a Z-stack of confocal images at 0.5 μm intervals were collected from the somatosensory cortex at layer V (Z = −2 to −2.2 mm from bregma and 3.2–3.4 mm lateral to midline) for each of at least two matched mutant and control mice. Images were examined using NIH Image software 1.31v to calculate cell size in square micrometers after manual drawing of cell margins. All SMI311-positive (SMI311+)(+) cells in a 20× field were measured, independent of pS6 staining, for a minimum of eight cells per field, and the largest eight cells were included in the measurements. Measurements were compared using the Mann–Whitney U test.

To quantify dendritic orientation as a measure of neuronal dysplasia, the direction of the apical dendrite of each SMI311+ pyramidal neuron in layer V of a section of somatosensory cortex was assessed. Neurons with apical dendrites oriented within 15° of a vertical line to the pia from the cell center were considered normal. Neurons with apical dendrites oriented outside of this 15° range in either direction were considered to have aberrant orientation.

Drug handling, administration, and pharmacokinetic analyses. Rapamycin was obtained from LC Laboratories, dissolved at 20 mg/ml in ethanol, and stored at −20°C for up to 1 month. Before each administration, rapamycin was diluted in 5% Tween 80, 5% polyethylene glycol 400 (0.5–1.5 mg/ml). Rapamycin was given at 6 mg/kg intraperitoneally every 2 d. RAD001 was provided by Novartis in a proprietary vehicle at 20 mg/ml intraperitoneally every 2 d. Both drugs were given at the appropriate dose to P7–P9, every other day, for up to 92 d (P100).

For pharmacokinetic analyses, each drug was given once or multiple times intraperitoneally to control, nonmutant mice from the Tsc1+/-cre colony. Mice were killed at 1, 4, 12, 24, or 48 h after drug administration, blood was collected in acid–citrate–dextrose solution, and plasma was separated by centrifugation at 10,000 rpm for 5 min. The plasma was then quickly removed and frozen at −80°C. Organ extracts were prepared by homogenization in 5× (w/v) of PBS until a fine suspension was achieved. This solution was clarified by centrifugation at 12,000 × g for 15 min and then frozen at −80°C until ready for assay. Rapamycin levels were determined after solid-phase extraction using liquid chromatography/tandem mass spectrometry (LC/MS/MS) on an API-2000 (Applied Biosystems) instrument in the Clinical Laboratories, Children’s Hospital, Boston. This is a Clinical Laboratory Improvement Amendments-certified laboratory that routinely determines rapamycin levels on pediatric samples. RAD001 levels were determined using LC/MS/MS by Ann Brown at Novartis Biomedical Research Institute (Cambridge, MA). Briefly, tissue lysates, separated plasma, and calibration standards were extracted using the acetonitrile (ACN) protein precipitation technique. Reverse-phase chromatography (column: ACE C18 30 × 2.1 mm, 3 μm) was used using a gradient elution with 0.1% HCOOH/water and 0.1% HCOOH/ACN as mobile phase A and B, respectively. The analyses were performed in electrospay-positive mode using multiple reaction monitoring conditions on a Sciex API 4000 instrument (Applied Biosystems).

DiI staining of neurons and dendritic spine measurements. Spine tracing and analysis was performed as described previously (Murai et al., 2003). Briefly, anesthetized mice were transcardially perfused with PBS and fixed with 4% PFA at 2 ml/min for 5 min. Whole brains were removed and postfix for 30 min in 4% PFA on ice. Brains were then embedded in 3% agarose and sectioned at 150 μm using a vibrating-blade microtome (Leica). Introduction of 1,1’-dioctadecyl-3,3,3,3’-tetramethylindocarbocyanine perchlorate (DiI) (Invitrogen)-coated gold particles was performed using the Helios Gene-Gun (Bio-Rad) at 80 psi. Coated particles were prepared by mixing 20 mg of gold particles with 5 mg of DiI. After gene gun particle injection, sections were placed in 4% PFA for 2 h to allow DiI diffusion along the cell membrane. Sections were then imaged by confocal microscopy. A Z-stack of confocal images at 0.5 μm intervals was collected from the somatosensory cortex in layer V. Spine analysis was performed using MetaMorph software (Molecular Devices). Spine lengths were measured from the tip to the intersection of the spine with the dendrite. Measurements of spine length (n = 856–1742 spines) and spine density (n = 10–19 dendrites) were compared using the Mann–Whitney U test in Prism (GraphPad Software).

Results

Pharmacokinetics of rapamycin and RAD001 in the brain

We explored the pharmacokinetics and pharmacodynamics of rapamycin and RAD001 to help define an effective dose of each compound for treatment of the previously described Tsc1+/–mice (Meikle et al., 2007). These drugs were chosen for study because they are both orally formulated for administration in humans, and both drugs are known to inhibit the kinase activity of mTOR by binding to FK506 binding protein 12 (FKBP12), which then binds to the mTORC1 complex (Brown et al., 1994). In addition, because the drugs are structurally different (Schuler et al., 1997), a direct comparison of both compounds was performed to determine whether they had different pharmacologic properties, particularly their relative penetration into the brain. Although both rapamycin and RAD001 can be given orally to older mice, we administered both drugs intraperitoneally because it is difficult to perform gavage on P7 mice, the age at which we chose to initiate treatment.

A single dose of either drug given at 6 mg/kg intraperitoneally to control mice aged P30–P45 led to substantial drug levels in plasma, liver, and brain (Fig. 1A, D). Brain levels remained markedly lower than systemic levels at all time points, consistent with an effect of the blood–brain barrier in reducing penetration into the CNS. Nonetheless, brain levels of each drug remained above the level required to inhibit mTORC1 (10 ng/ml for either drug when applied to cells in vitro) throughout the 48 h period after administration (rapamycin, 38.6 ng/g; RAD001, 40.5 ng/g; at 48 h after drug administration). Note also that therapeutic trough levels for each drug in humans are 3–20 ng/ml in whole blood. Both drugs displayed higher levels in nearly all cases when given every other day over a 3 week period (12 doses) compared with a single dose (Fig. 1B, E). Brain levels 48 h after the last of 12 doses were rapamycin at 88.4 ng/g and RAD001 at 48.9 ng/g. In an initial pharmacodynamic analysis, pS6(S235/236) and pS6(S240/244) levels were assessed by immunoblot analysis of whole-brain lysates from the Tsc1+/–mice, at 24 h and 48 h after the last treatment at 6 mg/kg intraperitoneally in a 12 dose every other day of treatment regimen. Phosphorylation of S6 at both sites returned to control (or lower) levels 24 h after the last dose (Fig. 1I) and remained low at 48 h (data not shown). However, this normalization of pS6 levels at 48 h after the last dose was not as consistent in mice similarly treated with 3 mg/kg (data not shown).

We also assessed whether the pharmacokinetics of these drugs was different in younger mice. Liver levels were increased 3.5-fold for rapamycin and 4.1-fold for RAD001 in P10 mice 24 h after a single intraperitoneal injection compared with similarly treated P30–P45 mice (Fig. 1C, F). These data indicate that overall clear-
levels in plasma, brain, and liver. Time after a single intraperitoneal injection is shown on the Pharmacokinetic analyses of rapamycin (*). Time after a single intraperitoneal injection is shown on the x-axis, and the y-axis represents drug levels in plasma, brain, and liver.

24 h after the last dose of drug shows reduction of Tsc1 protein levels (*) in all mice are indicated. Treatment began at P7–P9 for all mice. Note that dashed lines reflect a cohort of mice in which treatment was developed at P10. These data indicated that penetration of rapamycin and RAD001 into the CNS was substantial, although it is clearly much higher in younger mice. Although levels were high at P10, we elected to use 6 mg/kg intraperitoneally every other day as our standard dose for several reasons. First, we wanted to ensure that we would have effective mTORC inhibition at the dose used throughout the period of treatment, to have maximal potential therapeutic effect. Second, although levels clearly rose with repeat dosing, we were concerned that these levels might be misleading in reflecting retention of drug in a lipid compartment in the brain or drug bound to protein, which would not be free to enter into a complex with FKBP12, required for mTORC1 inhibition. Finally, as noted above, mTORC1 inhibition in the brain, as assessed by pS6 immunoblotting, was more effective at this dose than at 3 mg/kg for either drug.

Discontinuation of either drug treatment at P30 led to sustained clinical improvement for 1–2 weeks, followed by a gradual clinical deterioration (Fig. 2D, compare with C), which led to death at a median age of 79 d for rapamycin and 77 d for RAD001 (Fig. 1G,H). At P60, 30 d after stopping drug, rapamycin-treated mice had significant weight gain and significant phenotypic improvement in clasping behavior, tremor, and tail position (all p ≤ 0.03) compared with untreated P30 mutant mice. Both drugs had a similar effect in suppressing the growth and weight gain of control mice at P30 (Table 1).

Effects of treatment on cortical organization, dysplasia, cell size enlargement, and myelination We examined several aspects of cortical organization and neuronal morphology to provide insight into the mechanism of action of rapamycin/RAD001, following up previous observations in the Tsc1null-neuron mice (Meikle et al., 2007). Because these two compounds had equivalent therapeutic effects on survival (Fig. 1G,H) and in phenotypic improvement at both P30 and P100 (Table 1, all pairwise comparisons of phenotypic measures are
nonsignificant), we focused these studies on mice treated with rapamycin. Analysis of cortical sections from the rapamycin-treated Tsc1null-neuron mice showed several aspects of improvement (Fig. 2 E–L). First, treatment of the mutant mice with rapamycin led to a marked reduction in the extent of expression of pS6(S235/236) compared with untreated controls (Fig. 2E–G). This was apparent throughout the cortex but was most marked in a subset of enlarged pS6+ cells seen at the base of the cortex (Fig. 2F, *) and in cortical layer V in the mutants. A considerable decrease in levels of pS6(S235/236) was also noted in the thalamus and CA3 region of the hippocampus in the treated mutant when compared with untreated controls (see further below). Interestingly, these abnormal pS6–positive cells reappeared within 2 weeks of discontinuation of rapamycin (Fig. 2H, *). Treatment of control mice with rapamycin suppressed pS6(S235/236) levels even lower than in untreated controls (Fig. 2I). Standard histological sections also showed that these enlarged cells were markedly diminished in the rapamycin-treated mutants (Fig. 2J–L, note * in K). However, the mild overall cortical disorganization noted in the untreated mutants was not affected by rapamycin treatment.

We also examined neuronal morphology in the treated mice, using the SM1311 antibody against nonphosphorylated neurofilament to identify a restricted neuronal population in lateral somatosensory cortex, as done previously (Meikle et al., 2007) (Fig. 3A–J, see N for orientation). We observed that these cells were considerably enlarged in Tsc1null-neuron mice compared with SM1311+ cells from the identical region of control mice (Fig. 3A, B, F, G). Quantitative measurement showed that there was a considerable reduction in cell size after treatment (reduced from 1.81-fold to 1.18-fold when compared with untreated controls, p < 0.0001). However, this was a reversible phenomenon as the enlarged SM1311+ pyramidal cells reappeared when rapamycin treatment was discontinued for 2 weeks (Fig. 3D, I, M) (increased to 1.77-fold compared with untreated controls, p = 0.02).

Thus, rapamycin was very effective at reducing cell size in Tsc1null-neuron mice. However, despite this drastic reduction in cell size, rapamycin treatment appeared to have little effect on the dysplastic features of the neurons in this model (Fig. 3K, L). To examine this quantitatively, we assessed the orientation of the apical dendrite in SM1311+ layer V neurons in somatosensory cortex (Table 2). In control mice, nearly all (96%) neurons were polarized with a long apical dendrite that was oriented directly (±15°) toward the pial surface. In contrast, Tsc1null-neuron neurons often had major dendrites that extended tangentially and diagonally to the pia. In addition, rapamycin treatment initiated
at P7 did not reduce the percentage of SMI311+ neurons with abnormally oriented dendrites in Tsc1null-neuron mice (Table 2, 38.8% without treatment, 43.5% with rapamycin treatment). Tsc1null-neuron mice have impaired myelination to the extent that the cerebral cortex of a P30 mouse had only a faint patchy myelin stain, consistent with reduced myelin synthesis by oligodendrocytes (Meikle et al., 2007) (Fig. 4C). Rapamycin treatment effectively restored myelination in the Tsc1null-neuron brain (Fig. 4A–I; compare C with A, B; compare H with F, G). Although restoration of myelination was seen throughout the brain, the most dramatic improvement was seen in the cortex in which MBP+ myelin sheaths were evident coating radiating fibers extending out from the base of the cortex and in the pericallosal portion of the retrosplenial granular region (Fig. 4D). An improvement in myelination was also seen in the hippocampus (Fig. 4H). Double staining with pS6(S235/236) and MBP showed that there was a clear concordance between reduction in pS6 levels and restoration of myelin expression, as seen in the CA3 region of the hippocampus (Fig. 4K–O). Despite reducing pS6 levels to a subnormal level, rapamycin appeared to have little effect on myelination in the treated controls (Fig. 2E,F,O).

Biochemical and signaling effects of rapamycin treatment in the Tsc1null-neuron mice
Recent studies indicate that an important signaling effect in cells lacking Tsc1 or Tsc2 is a reduction in activation of Akt in response to normal stimuli (Zhang et al., 2003; Harrington et al., 2004; Shah et al., 2004). There has been speculation that this effect might have significant pathophysiological consequences in addition to that of mTORC1 activation in cells lacking Tsc1/Tsc2 (Manning et al., 2005). We assessed this possibility in brain extracts from the Tsc1null-neuron mice (Fig. 5A). Whereas pS6(S235/236) and pS6(S240/244) levels were significantly increased in the mutant mice, pAKt(S473) levels were reduced compared with controls. Moreover, rapamycin treatment led to restoration of pAKt(S473) levels, just as it decreased pS6 levels at both phosphorylation sites. Both of these effects were reversed when rapamycin treatment was discontinued. We also explored downstream signaling effects of this suppression of Akt signaling in the Tsc1null-neuron mice. pGSK3β(S9) levels were also reduced in the Tsc1null-neuron mice and were reversed by treatment with rapamycin (Fig. 5B), consistent with diminished signaling downstream of Akt.

Cytoskeletal abnormalities have also been reported in cells and neurons lacking Tsc1/Tsc2 (Lamb et al., 2000; Goncharova et al., 2004). Levels of phosphorylated neurofilament, neurofilament heavy chain, and neurofilament medium chain were all increased in the Tsc1null-neuron mice compared with controls (Fig. 5C). Furthermore, these effects on the neuronal cytoskeleton were effectively reversed by rapamycin treatment. In contrast, we found no consistent evidence of significant alterations in pCofilin(S3) levels in the Tsc1null-neuron mice compared with controls (Fig. 5C).

Dendritic spine abnormalities in the Tsc1null-neuron mice
Because of a previous report of significant effects of loss of Tsc1 or Tsc2 on dendritic spine density, shape, and length in in vitro hippocampal slice cultures (Tavazoie et al., 2005), we examined dendritic spine morphology in the Tsc1null-neuron mice including in response to rapamycin treatment, using bioscopic with Dil to label a small subset of cortical neurons. Confocal microscopy demonstrated that robust staining was achieved in select neurons (Fig. 6A). Quantitative analysis of spine density and length indicated that dendrites of cortical neurons from Tsc1null-neuron mice had a significant (p < 0.05), 22% reduction in spine density compared with neuronal dendrites from control mice (Fig. 6B,C, Table 3). However, there was no significant difference in spine length in neurons from these two types of mice. In response to rapamycin treatment of the Tsc1null-neuron mice, there was a small (10%) increase in spine density (p value was nonsignificant) toward a normal density. In addition, there was an 9% increase in spine length in the rapamycin-treated Tsc1null-neuron mice compared with both mutant and control mice (Table 3, p ≤ 0.002).

Discussion
The Tsc1null-neuron mice studied here replicate several of the clinical and pathologic features seen in TSC patients (Huttenlocher and Wollmann, 1991; Crino, 2004; Zikou et al., 2005; Ess, 2006; Marcotte and Crino, 2006; Holmes and Stafstrom, 2007). There are enlarged and ectopic cells, with prominent dysplasia, and high
level expression of pS6, as well as reduced myelination (Meikle et al., 2007). The mice show a progressive neurologic phenotype with tremor, hyperactivity, poor weight gain, seizure tendency, and limited survival. The current work demonstrates the marked therapeutic benefit of both rapamycin and RAD001 to effect both dramatic clinical and substantial histologic improvement in this TSC model. Mice treated at 6 mg/kg intraperitoneally every other day with either drug enjoyed survival out past 100 d in the vast majority of mice, with persistent improvement in clinical pheno-
weeks when drug treatment was discontinued. Whereas cell enlargement and biochemical and signaling profiles reverted to their pretreatment patterns within 2 weeks (Figs. 2H, 3I, 4N, 5A), myelination remained intact. It is therefore likely that improved myelination played a major role in the reduction in tremor, ataxia, and spasticity seen in the treated mutant mice. As shown previously (Meikle et al., 2007), this defect in myelination is not attributable to abnormal oligodendrocyte number or distribution, but rather there is a neuronal inductive defect, which as shown here is responsive to rapamycin/RAD001 treatment. Although the precise mechanism requires additional study, it is likely attributable to overactive mTORC1.

In contrast to the many features that were improved in this model in response to treatment, neuronal dysplasia and neuronal migration were both unchanged (Fig. 3K, L, Table 2). This is consistent with completion of neuronal differentiation and cortical migration before institution of rapamycin/RAD001 therapy at P7–P9. It is possible that earlier treatment with either compound might reduce neuronal dysplasia, but any benefit might be offset by other growth and developmental consequences of mTORC1 blockade.

Although spine density was significantly reduced (22%) in the Tsc1null-neuron mice, there was no significant change in spine length or morphology (qualitative observations only) in these mice compared with controls. In response to rapamycin treatment, there was only a modest increase (10%) in spine density and a corresponding increase (9%) in spine length above normal, suggesting that these dendritic morphologic abnormalities had little direct importance for neuronal function in this model. Conversely, phosphorylated neurofilament, neurofilament heavy chain, and neurofilament medium chain levels were all increased in the Tsc1null-neuron mice and were reversed by rapamycin treatment (Fig. 5B). In contrast to a previous report from in vitro slice cultures (Tavazoie et al., 2005), we saw no significant change in pCofilin(S3) levels in brain extracts from the Tsc1null-neuron mice, suggesting that this actin regulatory protein had little to do with the in vivo phenotype produced by loss of Tsc1 in neurons.

Our pharmacokinetic studies demonstrate that rapamycin/RAD001 brain levels are approximately one-third that of systemic levels at 48 h after the last dose, in both acute and chronic treatment paradigms (Fig. 1A, B, D, E). These findings are consistent with the clear treatment benefit noted. The buildup of each drug that is seen over time in the brain may serve as a reservoir for slow release when treatment is discontinued. This phenomenon may help to explain the prolonged symptom-free interval and survival seen after drug withdrawal at P30 in the treated mice. Although rapamycin/RAD001 levels achieved in these mice were significantly higher than are typically sought in patients, it is notable that a lower dose of drug could have been used to achieve both low therapeutic range brain levels (~5 ng/g) and concurrent...
high therapeutic range plasma levels (15–20 ng/ml). This is consistent with more limited studies we have performed, in which both rapamycin and RAD001 at 1 or 3 mg/kg given intraperitoneally three times per week led to clear therapeutic benefit in this model.

Loss of TSC1/TSC2 is now well known to lead to constitutive elevation of rheb-GTP levels and consequent constitutive activation of mTORC1, which causes transcriptional effects to influence cell size increase and growth by phosphorylation and activation of S6 kinase and phosphorylation and inactivation of eIF-4E binding protein 1. In addition to these direct or “downstream” effects, TSC1/TSC2 loss also leads to indirect (also termed “feedback” or “upstream”) effects that limit the phosphorylation and activation of AKT (Zhang et al., 2003; Harrington et al., 2004; Shah et al., 2004). This is the first work to demonstrate these complex effects of loss of Tsc1/Tsc2 in the brain, with strong AKT downregulation seen concurrent with activation of mTORC1. Rapamycin/RAD001 had prominent activity in blocking both direct and indirect effects of mTORC1 activation, restoring Akt phosphorylation. Reduced AKT expression has been engineered in mice, although it is difficult because of the existence of three different AKT isoforms with varying expression levels in various tissues. Mice with significant (50% or more) reduction in brain AKT expression have a significant phenotype with microcephaly and reduced numbers and size of neurons, although neurologic and behavioral abnormalities have not been investigated in detail (Easton et al., 2005; Tschopp et al., 2005; Dummler et al., 2006). Reduced AKT expression leads to a major (80–90%) reduction in pAKT(S473) levels in these brains, whereas pTsc2 and pGSK3β levels were near normal (Dummler et al., 2006). As we see the opposite clinical phenotype in the Tsc2null-neuron mice, with brain and neuronal enlargement, it is clear that these potential effects of AKT downregulation are overcome by the effects of mTORC1 activation in the Tsc1null-neuron mice. Nonetheless, as levels of pGSK3β were more reduced in the Tsc1null-neuron brains than in AKT-deficient brains (Dummler et al., 2006), it is possible that restoration of Akt function contributed significantly to the improvement in neurologic function seen in the Tsc1null-neuron mice in response to treatment. Significant concern has been raised by the possibility that elevation in pAKT(S473) might occur because of rapamycin/RAD001 treatment of malignancy, leading to a growth effect that could negate the potential benefits of mTORC1 blockade (O’Reilly et al., 2006). In this model, elevation of pAKT(S473) did occur in response to these drugs, concurrent with a marked phenotypic and histologic improvement, suggesting that it contributed to rather than impeded the clinical response.

Finally, given the similarities between the cellular and pathological abnormalities seen in this model and cortical tubers (Meikle et al., 2007), these findings suggest the possibility that rapamycin/RAD001 might have clinical benefit in the treatment of TSC patients. Indeed, rapamycin has been shown to have significant benefit, with shrinkage in size of TSC subependymal
c
trol, P30 mutant (Tsc1null-neuron), and P30 rapamycin-treated mutant and control (B) mice are shown. B, Note reduction in elevated pS6(240/244) levels as well as an increase in reduced pS473–AKT and pS9 –GSK3B levels of the mutant mice with treatment. C, In addition, abnormal increases in phosphorylated neurofilament (pNF) and neurofilament heavy (NF-H) and medium (NF-M) chains are reduced in response to rapamycin. Note that pCofilin(S3) levels are not consistently different among mutant and control mice. The untreated and treated controls were all Tsc1null-neuron; untreated mutants were all Tsc1null-neuron; treated mutants were one Tsc1null-neuron + and two Tsc1null-neuron –.
giant cell tumors (Franz et al., 2006). In addition, the brain penetration shown here in P10 mice suggests that rapamycin would also penetrate the CNS at high levels in infants. Therefore, these drugs may possibly have benefit in the treatment of TSC-associated infantile spasms, often a difficult clinical problem (Curatolo et al., 2006; Holmes and Stafstrom, 2007). Because similar drugs may possibly have benefit in the treatment of TSC-associated infantile spasms, often a difficult clinical problem (Curatolo et al., 2006; Holmes and Stafstrom, 2007). Because similar

Table 3. Spine density and length measurements

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Treatment</th>
<th>n cells/spines</th>
<th>Density (μm⁻¹)</th>
<th>Length (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>None</td>
<td>16/1105</td>
<td>1.73 ± 0.14a</td>
<td>1.38 ± 0.02c</td>
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<tr>
<td>Control</td>
<td>Rapamycin</td>
<td>19/1742</td>
<td>1.73 ± 0.12b</td>
<td>1.38 ± 0.02d</td>
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<tr>
<td>Mutant</td>
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<td>17/1524</td>
<td>1.35 ± 0.10c</td>
<td>1.39 ± 0.02e</td>
</tr>
<tr>
<td>Mutant</td>
<td>Rapamycin</td>
<td>10/856</td>
<td>1.49 ± 0.12</td>
<td>1.51 ± 0.05f</td>
</tr>
</tbody>
</table>

*Paired comparison p < 0.05 by both unpaired t and Mann–Whitney tests. All other comparisons in this column are not significant.

Note added in proof. We draw the reader’s attention to concurrent work published recently by Zeng et al. (2008). These investigators report that rapamycin is very effective in the treatment of a mouse astrocyte model of TSC based on this same Tsc1 conditional allele with marked improvement in survival, epilepsy, and brain pathology.

References


