Everolimus in the treatment of subependymal giant cell astrocytomas, angiomyolipomas, and pulmonary and skin lesions associated with tuberous sclerosis complex

David Neal Franz

Department of Pediatrics, Tuberous Sclerosis Clinic, Cincinnati Children’s Hospital Medical Center, Cincinnati, OH, USA

Abstract: Tuberous sclerosis complex (TSC) is an autosomal dominant genetic disorder caused by inactivating mutations in either the TSC1 or TSC2 genes. It is characterized by the development of multiple, benign tumors in several organs throughout the body. Lesions occur in the brain, kidneys, heart, liver, lungs, and skin and result in seizures and epilepsy, mental retardation, autism, and renal and pulmonary organ system dysfunction, as well as other complications. Elucidation of the molecular pathways and etiological factors responsible for causing TSC has led to a paradigm shift in the management and treatment of the disease. TSC1 or TSC2 mutations lead to constitutive upregulation of the mammalian target of rapamycin pathway, which affects many cellular processes involved in tumor growth. By targeting mammalian target of rapamycin with everolimus, an orally active rapamycin derivative, clinically meaningful and statistically significant reductions in tumor burden have been achieved for the main brain (subependymal giant cell astrocytoma) and renal manifestations (angiomyolipoma) associated with TSC. This review provides an overview of TSC, everolimus, and the clinical trials that led to its approval for the treatment of TSC-associated subependymal giant cell astrocytoma and renal angiomyolipoma.

Keywords: everolimus, subependymal giant cell astrocytoma, angiomyolipomas, lymphangioleiomyomatosis, facial angiofibromas, tuberous sclerosis complex

Epidemiology and etiology of subependymal giant cell astrocytomas, angiomyolipomas, skin lesions, and lymphangioleiomyomatosis associated with tuberous sclerosis complex

Tuberous sclerosis complex (TSC) is an autosomal dominant genetic disorder with an approximate birth incidence of 1:6,000 newborns.1–3 Up to 1 million people worldwide are affected.4 Although TSC can be inherited, approximately two-thirds of all cases are sporadic.5 Patients with TSC have a severely debilitating disorder that is associated with multiple benign tumors (hamartomas), most commonly in the brain, kidney, lungs, and skin.5,6 Common symptoms include seizures (90%); renal tumors, also known as angiomyolipomas (up to 80%); skin lesions, ie, ash leaf macules and cutaneous fibromas (~70%); pulmonary involvement, specifically lymphangioleiomyomatosis (LAM) (~40%); autism and autism spectrum disorders (30%–40%); and neurocognitive impairments (50%–60%), which often occur in individuals with a normal intelligence quotient.5,7,8
In the brain, up to 20% of TSC patients develop subependymal giant cell astrocytomas (SEGAs)—slow-growing glioneuronal tumors that typically arise near the foramen of Monro. SEGAs may grow and obstruct cerebrospinal fluid and increase cerebral pressure, causing headaches, blurred vision, or sudden death from acute hydrocephalus (Figure 1). The extent of elevated intracranial pressure and hydrocephalus is proportional to tumor diameter.

In the kidney, angiomyolipoma is one of the leading causes of death in TSC patients. Angiomyolipomas occur in up to 80% of patients and typically are multiple and located bilaterally in the kidneys. They tend to increase in size over time, with some becoming $>25-30$ cm in their longest dimension (Figure 2). Patients are at risk of serious complications such as life-threatening hemorrhage (25%–50%) and often present in the emergency room with shock (~20%–30%). As angiomyolipomas enlarge, they often develop micro- and macroaneurysms. Angiomyolipomas $\geq$4 cm in diameter are at increased risk for spontaneous bleeding as a result of ruptured vessels. The presence of aneurysms $>5$ mm is a strong predictor of spontaneous hemorrhage. Renal failure resulting in end-stage renal disease is largely due to encroachment of angiomyolipomas on normal renal parenchyma. Many patients require dialytic replacement therapy.

Skin lesions are detected in $>70%$ of patients with TSC and include hypomelanotic macules (ash leaf spots), shagreen patches, confetti-like lesions, forehead fibrous plaque, facial angiofibromas, and periungual and ungual fibromas (Figure 3A and B). Facial angiofibromas can result in decreased quality of life for patients because they affect appearance, may cause disfigurement, and are prone to bleeding, which increases the possibility of infection.

Pulmonary LAM occurs in up to 40% of TSC patients and predominantly affects women. LAM occurs more rarely in patients without TSC (sporadic LAM). It is characterized by cystic destruction of the lung caused by infiltration of smooth muscle cells (Figure 4).

The molecular pathogenesis of TSC is thought to be caused by mutations in either the TSC1 (encoding hamartin) or TSC2 (encoding tuberin) genes, both of which are integral to the normal functioning of the mammalian target of rapamycin (mTOR) pathway. The mTOR pathway integrates numerous cellular inputs to affect a multitude of

Figure 1 Computed tomography scan demonstrating acute presentation of bilateral subependymal giant cell astrocytomas with marked obstructive hydrocephalus. After emergency surgical resection, the patient with tuberous sclerosis complex was left blind and mentally retarded.

Figure 2 Renal computed tomography scan demonstrating significant angiomyolipoma burden in both kidneys of a patient with tuberous sclerosis complex.

Figure 3 Examples of tuberous sclerosis complex-associated skin lesions: (A) angiofibromas and (B) shagreen patches and ash leaf macules.
downstream signaling cascades that are involved in cellular processes such as cellular metabolism, growth, proliferation, angiogenesis, and survival.\(^1,6,8,21\) Hamartin and tuberin interact to form a dimer that activates a GTPase, preventing phosphorylation of the GAP protein—Ras homolog enriched in brain (Rheb)—and inhibiting activation of mTOR complex 1 (mTORC1), a serine threonine kinase.\(^22,23\) In patients with TSC, the hamartin/tuberin dimer is unable to form because of mutations in TSC1 and TSC2. As a result, mTORC1 is constitutively activated, which leads to hyperactive mTOR signaling of the downstream kinase signaling cascade and consequent abnormalities in many cellular processes, including cell cycle progression, transcription, translation, and metabolic control (Figure 5).\(^22,24\)

TSC is thought to develop from complete loss of functional TSC1 or TSC2 and follows Knudson’s two-hit hypothesis whereby the first “hit,” causing a mutation in one allele of the TSC1 or TSC2 gene, is followed by a second “hit,” somatic mutation, in the corresponding TSC wild-type allele.\(^8\) Disease severity can be variable, even within families, and may reflect differential expression of normal and mutant TSC alleles.\(^25\)

TSC2 mutations, which occur in 70%–80% of cases, are more common than TSC1 mutations, which occur in 20%–30% of cases.\(^26\) Studies of TSC patients have found
that the clinical manifestations of TSC are more frequent and often more severe in patients with TSC2 mutations.\textsuperscript{18,19} Data from genotype/phenotype studies suggest that different types and locations of TSC germline mutations may be associated with distinct neurocognitive phenotypes.\textsuperscript{19,27,28}

**Management and current therapeutic strategies**

Recently, a subcommittee at the Tuberous Sclerosis Complex Clinical Consensus Conference (sponsored by the Tuberous Sclerosis Alliance) reviewed and updated the recommendations for surveillance and management of patients with TSC.\textsuperscript{29} Specific guidance on screening and follow-up evaluations, as well as clearly defined time intervals for clinical evaluations, is expected in the TSC guidelines and will help to standardize and improve the clinical care of TSC patients. These guidelines will soon be published and available to clinicians. Historically, treatment of TSC-associated hamartomas has focused on a combination of active and watchful waiting, palliative treatments with drug therapy, and surgical procedures to reduce tumor burden (Table 1).

**SEGA**

Active surveillance with serial neuroimaging is recommended even in asymptomatic patients, because SEGAs have the potential to grow over time.\textsuperscript{30,31} Surgical resection has been the standard treatment option for symptomatic SEGAs; however, scientific evidence for the effectiveness of surgical resection has been used, but its role in treating SEGA has not been clearly defined because efficacy and safety data are limited.\textsuperscript{32–35} Postoperative morbidity is variable and may occur in ∼50% of patients.\textsuperscript{36} Complications include increased prevalence of seizures, hydrocephalus, vision disorders, headaches, stroke, hemiparesis, and autism.\textsuperscript{32,33,35,37}

Tumor recurrence invariably is seen in the absence of gross total resection. This may occur with very large, locally invasive, or bilateral lesions.\textsuperscript{32,33} The timing of surgical intervention for SEGA is controversial, with some surgeons advocating early surgery, and others waiting for symptomatic lesions or development of hydrocephalus, which can facilitate a transcortical approach to the lesion.\textsuperscript{30,32,38,39} If surgery is chosen, gradual formation of an operative corridor using balloon dilation is an example of a minimally invasive technique.\textsuperscript{40} This approach has resulted in gross total resection with resolution of ventricular dilation and minimal cortical disruption.\textsuperscript{40} Clinical recommendations made by a panel of European experts who met in Rome, Italy, in March 2012, endorse surgery in symptomatic patients and asymptomatic patients with documented tumor growth.\textsuperscript{41} The panel also recommends everolimus (Afinitor\textsuperscript{4}, Novartis, East Hanover, NJ, USA) for the treatment of adults and children ≥3 years of age with SEGA associated with TSC who require therapeutic intervention but are not amenable to surgery.\textsuperscript{41} Contraindications to surgery include an inability to tolerate anesthesia and cases in which the risks of surgery outweigh the benefits (total resection is unachievable or poses significant risk to the patient).\textsuperscript{41} Endoscopic procedures are associated with lower morbidity, but they are limited to lesions with a diameter ≤2 cm.\textsuperscript{32} Gamma Knife stereotactic radiosurgery has been used, but its role in treating SEGA has not been clearly defined because efficacy and safety data are limited.

**Table 1 Common treatment modalities for SEGA, angiomyolipoma, skin lesions, and pulmonary LAM**

<table>
<thead>
<tr>
<th>Clinical issue</th>
<th>Treatment</th>
<th>Additional comment</th>
</tr>
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| SEGA\textsuperscript{30–32,34,41,45} | • Active surveillance  
• Surgery  
• Gamma Knife stereotactic radiosurgery  
• Pharmacotherapy with mTOR inhibitors | • Periodic neuroimaging is recommended to monitor lesion size  
• Timing of surgery is debatable  
• Incomplete excision of SEGAs results in tumor recurrence  
• Postoperative complications include limit to the efficacy of surgery |
| Angiomyolipoma\textsuperscript{7,43,40} | • Arterial embolization  
• Partial/total nephrectomy  
• Pharmacotherapy with mTOR inhibitors | • Repeat embolization required  
• Postembolization syndrome |
| Skin lesions\textsuperscript{5,14} | • Cryosurgery  
• Curettage  
• Dermabrasion  
• Chemical peeling  
• Excision  
• Laser therapy | • Variable recurrence rate  
• Repeat procedures needed |
| Pulmonary LAM\textsuperscript{1,45} | • Bronchodilators  
• Oxygen support  
• Estrogen antagonists | • Treatments are palliative only |

**Abbreviations:** LAM, lymphangioleiomyomatosis; mTOR, mammalian target of rapamycin; SEGA, subependymal giant cell astrocytoma.
sufficient to support its use. Standard chemotherapy is not advised owing to the fact that therapeutic efficacy data are lacking and risk of remote, second malignancies is increased.

**Angiomyolipomas**

Appropriate management of angiomyolipomas is determined by lesion size and symptoms. An association between angiomyolipoma size (>4 cm), aneurysm size (≥5 mm), and risk of hemorrhage has been noted. Embolization is preferred in the setting of recent or active hemorrhage, or when large or multiple aneurysms are present. Embolization may require multiple procedures over time and is associated with modest reductions in tumor bulk. Complications of embolization include a condition called postembolization syndrome, which is characterized by pain, fever, and malaise resulting from the presence of necrotic tissue in the retroperitoneum. This can be prevented by the use of prophylactic steroids.

**Skin lesions**

Currently, treatments for skin lesions such as facial angiobromas include dermabrasion, cryosurgery, curettage, chemical peeling, electrodesiccation, excision, and laser therapy; however, repeated treatment is often needed because lesions may recur.

**Pulmonary LAM**

Bronchodilators are used to help control symptoms in patients with reversible air flow obstruction. Other treatment approaches include oxygen support and treatment with estrogen antagonists; however, effective treatment options for this condition are lacking.

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**Pharmacology and mode of action of everolimus**

**Pharmacodynamics**

The pivotal role of the mTOR pathway in the etiology of TSC and its associated conditions provides a strong rationale for the use of mTOR inhibition as a targeted therapy. Everolimus is a rapamycin derivative that inhibits the mTOR pathway by acting on mTORC1 (Figure 5). It binds to FKBP-12, forming an inhibitory complex with mTORC1, thereby inhibiting mTOR kinase activity and downstream pathways. Everolimus reduces the phosphorylation of downstream effectors of mTOR, such as the translational repressor eukaryotic elongation factor 4E binding protein 1, which are involved in protein translation.

**Pharmacokinetics**

In healthy volunteers, everolimus was rapidly absorbed within 30 minutes and reached a maximum blood concentration of 44.2 ± 13.3 µg/L after a single 4 mg oral dose (time to reach maximal concentration [T\(_{max}\)] = 0.5 hour [range 0.5–1 hour]) (Table 2). In adult patients with solid tumors, once-daily dosing with 5 mg or 10 mg of everolimus resulted in rapid absorption within 1 hour. The mean (standard deviation) maximum steady-state blood concentration was 32 ng/L (9 ng/L) and 61 ng/L (17 ng/L) after the 5 mg and 10 mg dose, respectively. The median (range) time to maximum serum concentration at the 5 mg and 10 mg dose was 1 hour (1 hour) and 1 hour (1–6 hours), respectively. For these same doses, the mean (standard deviation) area under the curve at steady state was 238 ng⋅h/mL (77 ng⋅h/mL) and 514 ng⋅h/mL (231 ng⋅h/mL), respectively. In healthy

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**Table 2** Pharmacokinetic properties of everolimus in various patient populations

<table>
<thead>
<tr>
<th>Patient population</th>
<th>(T_{max}) (h)</th>
<th>(C_{max}) (ng/mL)</th>
<th>AUC (ng/mL⋅h)</th>
<th>Half-life (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy subjects(^a) (n = 24) 2.0 mg dose(^4)</td>
<td>0.5 (0.5–2.0)</td>
<td>17.9 ± 5.9</td>
<td>122 ± 52</td>
<td>31.5 ± 6.4</td>
</tr>
<tr>
<td>Healthy subjects(^a) (n = 12) 4.0 mg dose(^6)</td>
<td>0.5 (0.5–1.0)</td>
<td>44.2 ± 13.3</td>
<td>219 ± 69</td>
<td>32.2 ± 6.1</td>
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<td>Refractory advanced solid tumor patients aged ≥18 years (n = 4)(^8) 5.0 mg dose(^3)</td>
<td>1 (1)</td>
<td>32 ± 9</td>
<td>238 ± 77</td>
<td>—</td>
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<tr>
<td>Refractory or recurrent solid tumor patients aged ≥3 years to &lt;22 years (n = 6)(^b) 2.1 mg/m(^2)</td>
<td>1 (0.5–1.2)</td>
<td>21.5 (16.1–78.5)</td>
<td>139.8 (94.6–197.7)</td>
<td>18.0 (15.2–19.9)</td>
</tr>
<tr>
<td>Refractory or recurrent solid tumor patients aged ≥3 years to &lt;22 years (n = 8)(^b) 3.0 mg/m(^2)</td>
<td>1.1 (0.6–2.3)</td>
<td>43.5 (18–58)</td>
<td>197.6 (59.6–249.4)</td>
<td>15.9 (13.9–22.5)</td>
</tr>
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Notes: *Values are mean ± standard deviation, except for \(T_{max}\) which is median (range); all values are median (range).

Abbreviations: AUC, area under the concentration–time curve from zero to infinity; \(C_{max}\), maximum concentration; \(T_{max}\), time to reach maximum concentration.
human subjects, a high-fat meal delayed the median time to maximum concentration by 1.25 hours (median) and reduced the maximum serum concentration by 60% and the area under the concentration–time curve by 16%. Therefore, everolimus should be consistently administered with or without food to avoid fluctuations in therapeutic drug.

Everolimus is a substrate of CYP3A4 and P-glycoprotein. Therefore, administration of CYP3A4 inhibitors or inducers affects its metabolism. A comparison of the relative amount of parent compound in blood after 1.5 mg/kg oral and 2-hour 1.0 mg/kg intravenous administration of everolimus suggests that intestinal metabolism (first-pass effect) accounts for a large percentage of metabolism.

Everolimus is eliminated mainly in bile and, to a lesser extent, in urine. The mean elimination half-life is approximately 30 hours. In healthy subjects, the elimination half-life after a 4 mg dose of everolimus was 32.2 ± 6 hours. Hepatic impairment increases exposure of everolimus. The extent of exposure depends on whether hepatic impairment is mild, moderate, or severe. Dose reductions should be implemented in TSC patients with SEGA and/or renal angiomyolipoma and mild to moderate hepatic impairment. If the benefit outweighs the risk, reductions in doses of everolimus are warranted in TSC patients with severe hepatic impairment and renal angiomyolipoma; however, in TSC patients with SEGA and severe hepatic impairment, everolimus should be avoided.

**Clinical efficacy**

The efficacy of everolimus in treating TSC patients was first demonstrated in an open-label Phase I/II clinical trial in 28 patients with SEGA associated with TSC (NCT00411619). The core treatment phase lasted 6 months, after which patients could transition into an extension phase. This trial led to US Food and Drug Administration approval of everolimus for the treatment of SEGA in TSC patients; it was the first trial to show marked reductions in SEGA volume within the first 3 months of treatment that were sustained as long as therapy continued. Patients (median age 11 years [range 3–34 years]) received everolimus initiated at a dose of 3 mg/m² and subsequently adjusted to whole blood trough concentrations of 5–15 ng/mL. The median duration of treatment was 21.5 months (range 4.7–34.4 months). Clinically meaningful reductions in the volume of primary SEGA, as assessed by independent central review, were achieved, and the median reduction was 0.80 cm³ (95% confidence interval [CI]: 0.4–1.2; P < 0.001). No new lesions, evidence of increased intracranial pressure, worsened hydrocephalus, or need for surgical resection or any other therapy was reported. Everolimus was associated with a significant reduction in the overall frequency of clinical and subclinical seizures (median change −1 seizure; P = 0.02). Quality of Life in Childhood Epilepsy scores improved over time (58.74 ± 14 at baseline; 63.4 ± 12.4 at 3 months; 62.1 ± 14.2 at 6 months). This trial continues in the extension phase, and results for patients treated up to 3 years were recently reported by Krueger et al. Of the original 28 patients enrolled, 25 were continuing treatment at the data cutoff for the 3-year extension study. After a median exposure of 34.2 months, the primary SEGA volume was reduced from 1.74 cm³ at baseline to 0.97 cm³ at 36 months. Reductions ≥30% from baseline at 24 months, 30 months, and 36 months were seen in 79.2%, 64.7%, and 77.8% of patients, respectively, and reductions ≥50% were seen in 50.0%, 41.2%, and 55.6% of patients, respectively. At 24 months, improvements in facial angiofibromas compared with the previous visit were observed in eight of nine patients (88.9%). A subgroup analysis of patients enrolled in this trial reported a significant change in white matter diffusion in patients treated with everolimus, suggesting that treatment alters the genetic defects in normal-appearing white matter in the brain of TSC patients. Significant changes in fractional anisotropy and radial diffusivity were observed in the corpus callosum, internal capsule, and geniculocalarine regions of patients treated with everolimus, and no changes were observed in the age- and gender-matched control group of patients with TSC. Mean (95% CI) fractional anisotropy significantly increased by 0.04 (0.019–0.062) in all three brain regions after 12–18 months of everolimus treatment (P < 0.05), and this was driven by a decrease in radial diffusivity in the corpus callosum and the geniculocalarine tract (P < 0.01 for each).

A Phase III, randomized, double-blind, placebo-controlled trial, A Randomised, Double-blind, Placebo-controlled Study of Everolimus in the Treatment of Patients With Subependymal Gaint Cell Astrocytomas (SEGA) Associated With Tuberous Sclerosis Complex (TSC) (EXIST-1)(NCT 00789828), investigated the efficacy of everolimus versus placebo in 117 TSC patients with SEGA (Table 3). Patients were randomized to everolimus 4.5 mg/m² per day (n = 78) or placebo (n = 39), and baseline demographics and disease characteristics were generally well balanced, with the exception of a higher proportion of males and the presence of hydrocephalus in the everolimus group. The median age of patients was 9.5 years (range 0.8–26.6 years), and 51% and 64% of patients in the everolimus and placebo arms, respectively, had one target SEGA lesion; 44% and 36% of patients in the everolimus and
placebo arms, respectively, had two SEGAs lesions.\textsuperscript{36} One or more skin lesions were present in 110 patients (92\% [n = 72] of patients in the everolimus arm and 97\% of patients in the placebo arm [n = 38]), and 38\% (n = 30) of patients in the everolimus arm and 36\% (n = 14) of patients in the placebo arm had one or more angiomyolipomas.\textsuperscript{38} After a median follow-up of 9.7 months, the SEGAs response rate—the proportion of patients with a reduction in sum of volumes of target SEGAs lesions ≥50\% relative to baseline in the absence of nontarget lesion worsening, new lesions ≥1 cm in diameter, or new/ worsening hydrocephalus—was 35\% (95\% CI: 24\%–46\%) in the everolimus arm and 0\% (95\% CI: 0\%–9\%) in the placebo group (difference in response rate 35\%, 95\% CI: 15\%–52\%; P < 0.0001).\textsuperscript{36}

The median time to SEGAs progression was not reached in either treatment arm, and estimated progression-free rates at 6 months were 100\% for everolimus and 86\% for placebo.\textsuperscript{36} Analysis of change in seizure frequency was inconclusive—a fact that was attributed to the method of assessment, a single 24-hour video electroencephalography, which likely limited the number of evaluable patients. The median change from baseline to week 24 in seizure frequency was 0 in both treatment arms (P = 0.2004), yielding an inconclusive analysis, as most patients had no seizures at baseline or follow-up.\textsuperscript{36}

Thirty patients (42\%) in the everolimus arm and four patients (11\%) in the placebo arm had a skin lesion response.\textsuperscript{36} AEs were mostly grade 1/2 in severity. Most common AEs were mouth ulceration (32\% vs 5\% EVE vs PBO), stomatitis (31\% vs 21\% EVE vs PBO), convulsions (23\% vs 26\% EVE vs PBO), and pyrexia (22\% vs 15\% EVE vs PBO).

### Table 3 Efficacy and safety analyses from EXIST-1 and EXIST-2

<table>
<thead>
<tr>
<th>Trial</th>
<th>Efficacy</th>
<th>Safety</th>
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<tr>
<td><strong>EXIST-1\textsuperscript{36}</strong></td>
<td><strong>Primary end point</strong>&lt;br&gt;SEGAs response rate: 35% vs 0% (EVE vs PBO; P &lt; 0.0001)&lt;br&gt;Key secondary end points&lt;br&gt;Change from baseline to week 24 in the total number of seizures per 24 hours&lt;br&gt;– No significant difference in seizure frequency was found in the everolimus arm relative to placebo (P = 0.20)&lt;br&gt;Time to progression of SEGAs&lt;br&gt;– No EVE patients progressed vs six PBO patients (15.4% (P = 0.0002)&lt;br&gt;Skin lesion response rate (among patients with ≥1 skin lesion at baseline; n = 110)&lt;br&gt;– 42% vs 11% (EVE vs PBO; P = 0.0004)&lt;br&gt;Exploratory end points&lt;br&gt;Angiomyolipoma response rate (among patients with ≥1 angiomyolipoma ≥1 cm in longest diameter at baseline; n = 44)&lt;br&gt;53% vs 0% (EVE vs PBO)</td>
<td>AEs were mostly grade 1/2 in severity&lt;br&gt;Most common AEs were mouth ulceration (32% vs 5% EVE vs PBO), stomatitis (31% vs 21% EVE vs PBO), convulsions (23% vs 26% EVE vs PBO), and pyrexia (22% vs 15% EVE vs PBO)</td>
</tr>
<tr>
<td><strong>EXIST-2\textsuperscript{38}</strong></td>
<td><strong>Primary end point</strong>&lt;br&gt;Angiomyolipoma response rate&lt;br&gt;– 42% vs 0% (EVE vs PBO; P &lt; 0.0001)&lt;br&gt;Key secondary end points&lt;br&gt;Time to progression of angiomyolipoma&lt;br&gt;– EVE was superior to PBO (hazard ratio: 0.08; 95% CI: 0.02–0.37; P &lt; 0.0001)&lt;br&gt;Skin lesion response rate (among patients with ≥1 skin lesion at baseline; n = 114)&lt;br&gt;26% vs 0% (EVE vs PBO; P = 0.0002)</td>
<td>AEs were mostly grade 1/2 in severity&lt;br&gt;Most common AEs were stomatitis (48% vs 8% EVE vs PBO), nasopharyngitis (24% vs 31% EVE vs PBO), and acne-like skin lesions (22% vs 5% EVE vs PBO)</td>
</tr>
</tbody>
</table>

**Abbreviations:** AEs, adverse events; CI, confidence interval; EVE, everolimus; EXIST, EXaamining everolimus In a Study of TSC; PBO, placebo; SEGAs, subependymal giant cell astrocytomas; TSC, tuberous sclerosis complex; vs, versus.
everolimus and placebo arms were 57% and 49%, respectively. Of note, lesions ≥4 cm are at risk for hemorrhagic rupture. SEGAs were noted as a major feature in a higher percentage of everolimus patients (54%) compared with placebo patients (36%) at baseline, but patient demographics were well balanced between the treatment arms and were generally representative of disease epidemiology. Skin lesion response was determined for the 114 patients with one or more skin lesions at baseline.

The angiomyolipoma response rate, defined as the proportion of patients with confirmed ≥50% reductions in the sum of volumes of all target angiomyolipomas relative to baseline, was 42% (95% CI: 31%–53%) for everolimus compared with 0% (95% CI: 0%–9%) for placebo, translating to a significant difference of 42% (95% CI: 23.5%–58.4%; P < 0.0001). A higher proportion of patients in the everolimus arm had a ≥50% reduction from baseline in the sum of volumes of target angiomyolipoma lesions at week 24 (55% versus 0%). Similarly, a higher proportion of patients in the everolimus arm had a reduction in the sum of volumes of target angiomyolipoma lesions ≥30% at this same time point (80% versus 3%). The median time to angiomyolipoma response with everolimus was 2.9 months.

The everolimus arm had a significantly higher skin lesion response rate: response rates were 26% (95% CI: 17%–37%) in the everolimus arm and 0% (95% CI: 0%–10%) in the placebo arm (P = 0.0002). Time to angiomyolipoma progression was significantly longer with everolimus versus placebo (hazard ratio: 0.08 [95% CI: 0.02–0.37]), and progression-free rates for everolimus versus placebo, respectively, were 98% (95% CI: 95% CI: 89%–100%) versus 83% (95% CI: 65%–93%) at 6 months, and 92% (95% CI: 65%–98%) and 25% (95% CI: 1%–64%) at 12 months. The median time to angiomyolipoma progression was 11.4 months for placebo and was not reached for everolimus.

**Safety and tolerability**

In all three clinical trials, everolimus yielded a comparable safety profile: adverse events (AEs) were generally mild and grade 1 or 2 in severity, as assessed by the National Cancer Institute Common Terminology Criteria for Adverse Events. Stomatitis (79%) and upper respiratory tract infection (79%) accounted for the majority of AEs in the Phase I/II trial; grade 3 AEs were reported in ten patients, and one grade 4 event (convulsion) occurred in one patient. No consistent laboratory abnormalities were noted, with the exception of increases in the concentrations of total cholesterol, low-density lipoprotein cholesterol, and triglycerides. The safety profile in the extension phase continues to be consistent with that reported in the original analysis, and no patient discontinued treatment because of an AE. No patients discontinued because of AEs in EXIST-1, and few patients discontinued because of AEs in EXIST-2. Reasons for discontinuation in the everolimus arm included one case of grade 2 blood phosphorus decrease; one patient with concurrent grade 3 hypersensitivity, grade 3 angioedema, and grade 3 bronchospasm; and one patient with convulsion due to intractable seizures deemed not related to study drug. In the EXIST-1 study, the most common AEs in the everolimus versus placebo groups, respectively, were mouth ulceration (32% versus 5%), stomatitis (31% versus 21%), convulsion (23% versus 26%), and pyrexia (22% versus 15%). The most common grade 3 AEs in the everolimus versus placebo group were stomatitis (8% versus 3%), pyrexia (6% versus 0%), and convulsion (5% versus 5%). In the EXIST-2 study, the most common AEs in the everolimus versus placebo group, respectively, were stomatitis (48% versus 8%), nasopharyngitis (24% versus 31%), acne-like skin lesions (22% versus 5%), and headache (22% versus 18%). The most common grade 3 AEs in the everolimus versus placebo group were aphthous stomatitis (3% versus 0%) and mouth ulceration (3% versus 0%). Grade 4 events were rare in both studies. Secondary amenorrhea was observed in EXIST-1 and EXIST-2. In EXIST-1, three of eight everolimus-treated and no placebo-treated female patients experienced amenorrhea. In EXIST-2, seven of 52 everolimus-treated females and one of 26 placebo-treated females experienced amenorrhea. As a result, amenorrhea is being further investigated as a potential AE associated with everolimus.

**Biomarkers**

Loss of the TSC1 or TSC2 gene has been associated with increased production of vascular endothelial growth factor (VEGF), and inhibitors of mTOR have been shown to have an inhibitory effect on production of VEGF, tumor growth, and angiogenesis, both in vitro and in vivo.

The effect of everolimus on several angiogenic biomarkers is being assessed in the EXIST-1 and EXIST-2 trials. Compared with placebo, a sustained ~30% and ~60% increase in VEGF-A was observed in the everolimus arm of EXIST-1 and EXIST-2, respectively. A concomitant decrease in collagen type IV (~25% EXIST-1; ~45% EXIST-2) and soluble VEGF receptor (sVEGFR2) (~25% both trials) was observed in the everolimus arm. A sustained decrease
(~60%) in VEGF-D was observed in the everolimus arm of EXIST-2, but not in EXIST-1.64 In both studies, no change was observed in placental growth factor, sVEGF-R1, or c-Kit plasma concentrations in the everolimus arm or in any biomarkers evaluated in the placebo arm.64 Baseline sVEGF-R2 and VEGF-D were ~40% and ~four-fold higher, respectively, while VEGF-A was ~50% lower in EXIST-2 compared with in EXIST-1.64 A similar baseline plasma concentration for the other biomarkers was noted in both studies.64 The inclusion of biomarkers in future clinical trials may provide new predictive or prognostic biomarkers when evaluating disease burden and everolimus efficacy and safety.

Conclusion

Everolimus is approved for the treatment of adults with renal angiomylipoma and TSC not requiring immediate surgery, and of adults and children ≥3 years of age with SEGA associated with TSC who require therapeutic intervention but are not candidates for curative surgical resection. The therapeutic efficacy and safety of everolimus have been demonstrated in multicenter, international, randomized, double-blind, placebo-controlled trials. These studies continue in their extension phases with the goal of establishing the long-term safety and efficacy profile of everolimus in patients with TSC. Everolimus, an orally administered mTOR inhibitor, is the first pharmacological therapy approved for the treatment of TSC. Targeted systemic therapies such as everolimus are particularly useful in that they treat the multiple manifestations of TSC and offer patients both a means of symptom relief and a treatment alternative to invasive surgical procedures.

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