Cardiovascular Manifestations of Tuberous Sclerosis Complex and Summary of the Revised Diagnostic Criteria and Surveillance and Management Recommendations From the International Tuberous Sclerosis Consensus Group

Robert B. Hinton, MD; Ashwin Prakash, MD; Robb L. Romp, MD; Darcy A. Krueger, MD, PhD; Timothy K. Knilans, MD

Tuberous sclerosis complex (TSC) is a genetic syndrome with a highly variable phenotype that may affect several organ systems. The central nervous system findings were the first to be described, and the classic triad of cognitive impairment, facial angiofibromas, and seizures was delineated shortly thereafter.1,2 As the variability and extent of organ involvement were appreciated, diagnostic criteria evolved to include major and minor criteria that taken together would lead to a definite, probable, or possible clinical diagnosis.3,4 Since the most recent refinement of the diagnostic criteria, dramatic advances have been made in understanding the genetic basis and pathogenesis of TSC, and new treatment strategies have been established, significantly affecting all aspects of coordinated care for TSC patients.

The Tuberous Sclerosis Alliance (www.tsalliance.org) convened a Consensus Conference composed of 8 working groups that generated Revised Diagnostic Criteria5 and new Surveillance and Management Guidelines6 with the intention of creating “living documents” to accommodate rapid advances and the need for coordination of care. The conference was informed in part by a recent constituency survey of key opinion leaders, which summarized interim progress, areas in need of further research, unmet medical needs, and barriers to progress.7 The goals of this report are to highlight the new diagnostic criteria and management guidelines as they pertain to cardiology and to expand consideration of the issues relevant to optimal cardiac care of patients with TSC.

TSC is characterized by widespread hamartomas, or abnormal growth of normal tissues. Cardiac rhabdomyomas are hamartomatous growths or benign tumors composed of cardiac myocytes, and they represent the classic neonatal manifestation of cardiac disease in TSC. Additional cardiac diseases such as arrhythmia occur later in life, underscoring the importance of ongoing cardiology care. Here, we review what is known about the natural history of cardiac manifestations in TSC with an emphasis on diagnostic testing, surveillance, and treatment.

The Revised Diagnostic Criteria Include Clinical Genetic Testing

Significant changes have been implemented in the revised diagnostic criteria.5 For example, clinical genetic testing has been added as an independent criterion, sufficient to make the diagnosis of TSC. Since TSC1 and TSC2, the genes that encode hamartin and tuberin, were identified as the cause of TSC,8,9 substantial strides have been made in defining the pathogenesis of TSC. Mutations in the genes TSC1 and TSC2 cause 75% to 90% of cases (Figure A). Given the increasing appreciation for disease variability and an assortment of mild disease phenotypes that may be on the TSC spectrum, the inclusion of a molecular test represents an important change in the approach to diagnosis. While approximately one-third of cases have a positive family history, this has not been included as diagnostic criteria but remains informative given the various challenges with performing genetic testing. Importantly, the designation of a definite, probable, or possible clinical diagnosis has been simplified to either “definite” or “possible.” Additional changes were made in several of the clinical criteria (Table 1), and changes regarding cardiovascular features are considered next in detail.
Overall Recommendations Have Shifted to Careful Surveillance and Early Intervention

Guidelines for the management and surveillance of TSC patients were comprehensively addressed in a companion article to the revised diagnostic criteria. Given the successful clinical trials establishing mammalian target of rapamycin (mTOR) inhibition as a new pharmacologic treatment strategy, a variety of surveillance issues have been considered (Tables 2 and 3). The addition of genetic testing to the diagnostic criteria has implications for screening that were addressed as well. These recommendations affect cardiologists directly with respect to surveillance and potentially in rare circumstances with respect to medical therapy. There is an increasing appreciation for latent cardiovascular phenotypes, indicating a need for continued surveillance of these patients. As the natural history of disease in the cardiovascular system is better understood, continued care in adulthood needs to be defined, underscoring efforts to transition care from pediatric to adult cardiology and to maintain surveillance vigilance in adulthood.

The Natural History and Diagnosis of TSC

The Natural History of Cardiac Rhabdomyomas

Cardiac tumors are rare, and ascertaining incidence is difficult. Based on clinical studies and autopsy series, primary cardiac tumors occur in 0.2% of children. Cardiac rhabdomyomas are by far the most common primary cardiac tumor in childhood. After the advent of echocardiography, but before clinical genetic testing was available, studies estimated that up to 70% to 90% of children with rhabdomyomas have TSC, and at least 50% of children with TSC have rhabdomyomas, representing a significant increase in the proportion of cardiac rhabdomyomas attributed to TSC compared with historic clinical data. In 1 study, Allan et al analyzed 52 cardiac tumor cases, among which 44 (86%) were associated with TSC.
Table 1. Revised Diagnostic Criteria for TSC

<table>
<thead>
<tr>
<th>Genetic diagnostic criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>The identification of either a TSC1 or TSC2 pathogenic mutation in DNA from normal tissue is sufficient to make a definite diagnosis of TSC. A pathogenic mutation is defined as a mutation that clearly inactivates the function of the TSC1 or TSC2 proteins (eg, out of frame indel or nonsense mutation), prevents protein synthesis (eg, large genomic deletion), or is a missense mutation whose effect on protein function has been established by functional assessment (<a href="http://www.lovd.nl/TSC1">www.lovd.nl/TSC1</a>, <a href="http://www.lovd.nl/TSC2">www.lovd.nl/TSC2</a>). Other TSC1 or TSC2 variants whose effect on function is less certain do not meet these criteria and are not sufficient to make a definite diagnosis of TSC. Note that 10% to 25% of TSC patients have no mutation identified by conventional genetic testing, and a normal result does not exclude TSC, or have any effect on the use of Clinical Diagnostic Criteria to diagnose TSC.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Clinical diagnostic criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major features</td>
</tr>
<tr>
<td>1. Hypomelanotic macules (≥3, at least 5-mm diameter)</td>
</tr>
<tr>
<td>2. Angiofibromas (≥3) or fibrous cephalic plaque</td>
</tr>
<tr>
<td>3. Ungual fibromas (≥2)</td>
</tr>
<tr>
<td>4. Shagreen patch</td>
</tr>
<tr>
<td>5. Multiple retinal hamartomas</td>
</tr>
<tr>
<td>6. Cortical dysplasias*</td>
</tr>
<tr>
<td>7. Subependymal nodules</td>
</tr>
<tr>
<td>8. Subependymal giant cell astrocytoma</td>
</tr>
<tr>
<td>9. Cardiac rhabdomyoma</td>
</tr>
<tr>
<td>10. Lymphangioleiomyomatosis (LAM)*</td>
</tr>
<tr>
<td>11. Angiomyolipomas (≥2)*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Minor features</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. “Confetti” skin lesions</td>
</tr>
<tr>
<td>2. Dental enamel pits (&lt;3)</td>
</tr>
<tr>
<td>3. Intraoral fibromas (≥2)</td>
</tr>
<tr>
<td>4. Retinal achromic patch</td>
</tr>
<tr>
<td>5. Multiple renal cysts</td>
</tr>
<tr>
<td>6. Nonrenal hamartomas</td>
</tr>
</tbody>
</table>

Definite diagnosis: 2 major features or 1 major feature with ≥2 minor features. Possible diagnosis: either 1 major feature or ≥2 minor features. TSC indicates tuberous sclerosis complex.

*Includes tubers and cerebral white matter radial migration lines.

A combination of the 2 major clinical features LAM and angiomyolipomas without other features does not meet criteria for a definite diagnosis.

Reproduced with permission from Northrup et al.5

were rhabdomyomas.23 Tumors are diagnosed more frequently in fetal series than in postnatal series, resulting in an increased sensitivity when examining fetal echocardiograms.18,19 Cardiac rhabdomyomas tend to appear at 20 to 30 weeks’ gestation, with the earliest diagnosis having been made at 15 weeks at the current technical limits of ultrasoundography, suggesting rhabdomyomas may be present earlier in development. The frequency of fetal detection is increasing dramatically; therefore, it is reasonable to anticipate that the rate of fetal diagnosis will increase significantly.

Fetal cardiac tumors may present in utero as a mass on ultrasonography, irregular heart rhythm, hydrops fetalis, or pericardial effusion. Cardiac rhabdomyomas can increase in size during the second half of gestation, and this has been attributed to maternal hormonal changes associated with pregnancy. When larger tumors result in hemodynamic compromise in utero, intrauterine fetal demise may occur. Fetal loss has been reported to be ≈11% in 1 small series of 44 cases.23 Cardiac rhabdomyomas do not cause symptoms or hemodynamic compromise in the vast majority of patients but may become symptomatic shortly after birth or in the first year of life. Tumors may obstruct inflow or outflow, which can cause ventricular dysfunction and heart failure, as well as redirection of flow across the foramen ovale.19,25,26 Nearly 100% of fetuses with multiple rhabdomyomas have TSC, underscoring the practical importance of identifying additional tumors at the time of fetal assessment for diagnosis and prognosis.17,27 In light of emerging human genetic and molecular knowledge, it is a possibility that the underlying pathogenesis of all rhabdomyomas is a result of a spectrum of TSC disease.

Cardiac rhabdomyomas are typically well circumscribed and nonencapsulated (FigureB). Micropathologic examination demonstrates abnormal myocyte architecture, including vacuolization and pathognomonic “spider cells” (FigureC). The individual cardiac rhabdomyomas range in size from a few millimeters to several centimeters and are multiple in number in 90% of cases. There is an equal predilection for left, right, and septal ventricular myocardium.26,28 Tumors are typically located in the ventricles, where they can compromise ventricular function and on occasion interfere with valve function or result in outflow obstruction. Tumors may be located in the atria, where they can compress the coronary arteries, leading to myocardial ischemia.29

Diagnosis of Cardiac Rhabdomyoma

Echocardiography is the imaging modality of choice for assessing cardiac involvement in TSC. Cardiac rhabdomyomas can be detected prenatally or postnatally. In prenatal life, ultrasound detection of multiple cardiac tumors is often the first sign of TSC.30 Typically, cardiac rhabdomyomas are visible as multiple, echogenic, nodular masses embedded in the ventricular myocardium, sometimes protruding into the involved chamber (FigureD). They are homogeneous and hyperechoic compared with normal myocardium. Diagnosis of cardiac rhabdomyomas is made easily when these typical features are present, but differentiation from other cardiac tumors may be difficult when there is a large solitary tumor or when tumors are located in an atypical location, such as the atria. Doppler echocardiography is also useful in assessing the presence of obstruction to ventricular inflow or outflow.
Echocardiography is also used to assess ventricular function, which may be impaired by multiple confluent tumors. Cardiac rhabdomyomas are seen readily in fetal life after 20 weeks of gestation and are seen in a majority of infants with TSC. Rhabdomyomas can enlarge significantly in size during gestation and may be seen later in gestation when they are not visible prior to 20 weeks. Cardiac rhabdomyomas regress spontaneously in a large majority of patients during the first year of life and as a result are seen with decreasing frequency in patients with TSC after 2 years of age. There is some suggestion that the incidence of identifiable cardiac rhabdomyomas in TSC increases during adolescence, but this observation has not been validated in additional studies.

Sensitivity and Specificity of Echocardiography to Identify Cardiac Rhabdomyomas

This varies with patient age, related to the previous discussion. In fetal life, of patients diagnosed with cardiac rhabdomyomas by echocardiography, 75% to 80% are found to satisfy criteria for TSC postnatally. The presence of multiple ventricular tumors seems to be the finding best associated with TSC. The presence of a family history of TSC also increases the likelihood of TSC. In fetuses with a single ventricular tumor, only 30% satisfy criteria for TSC. Because a diagnosis of TSC during fetal life is often prompted by the presence of cardiac rhabdomyomas, the negative predictive value of fetal echocardiography is not established.

In early infancy, the predictive value of echocardiography is similar to that in fetal life, with ≈80% of infants with cardiac rhabdomyomas eventually satisfying a diagnosis of TSC. Conversely, 80% to 85% of children with confirmed TSC have identifiable rhabdomyomas when younger than 2 years. Beyond 2 years of age, the incidence of identifiable rhabdomyomas is significantly lower (≈20% to 25%), although if they are readily seen on echocardiography, the...
Table 3. Surveillance and Management Recommendations for Patients Already Diagnosed With Definite or Possible TSC Summary Table

<table>
<thead>
<tr>
<th>Organ System or Specialty Area</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genetics</td>
<td>● Offer genetic testing and family counseling, if not done previously, in individuals of reproductive age or newly considering having children</td>
</tr>
</tbody>
</table>
| Brain                         | ● Obtain MRI of the brain every 1 to 3 years in asymptomatic TSC patients under the age of 25 years to monitor for new occurrence of SEGA. Patients with large or growing SEGA, or with SEGA causing ventricular enlargement but still asymptomatic, should undergo MRI more frequently and the patients and their families should be educated regarding the potential of new symptoms. Patients with asymptomatic SEGA in childhood should continue to be imaged periodically as adults to ensure there is no growth.  
● Surgical resection should be performed for acutely symptomatic SEGA. Cerebrospinal fluid diversion (shunt) may also be necessary. Either surgical resection or medical treatment with mTOR inhibitors may be used for growing but otherwise asymptomatic SEGA. In determining the best treatment option, discussion of the complication risks, adverse effects, cost, length of treatment, and potential impact on TSC-associated comorbidities should be included in the decision-making process  
● Perform screening for TAND features at least annually at each clinical visit. Perform comprehensive formal evaluation for TAND at key developmental timepoints: infancy (0 to 3 years), preschool (3 to 6 years), pre-middle school (6 to 9 years), adolescence (12 to 16 years), early adulthood (18 to 25 years), and as needed thereafter. Management strategies should be based on the TAND profile of each patient and should be based on evidence-based good practice guidelines/practice parameters for individual disorders (eg, autism spectrum disorder, attention-deficit/hyperactivity disorder, anxiety disorder, etc). Always consider the need for an individual educational program (IEP). Sudden change in behavior should prompt medical/clinical evaluation to look at potential medical causes (eg, SEGA, seizures, renal disease, etc.)  
● Obtain routine EEG in individuals with known or suspected seizure activity. The frequency of routine EEG should be determined by clinical need rather than a specific defined interval. Prolonged video EEG, 24 hours or longer, is appropriate when seizure occurrence is unclear or when unexplained sleep, behavioral changes, or other alteration in cognitive or neurological function is present  
● Vigabatrin is the recommended first-line therapy for infantile spasms. ACTH can be used if treatment with vigabatrin is unsuccessful. Anticonvulsant therapy of other seizure types in TSC should generally follow that of other epilepsies. Epilepsy surgery should be considered for medically refractory TSC patients, but special consideration should be given to children at younger ages experiencing neurologic regression and is best if performed at epilepsy centers with experience and expertise in TSC |
| Kidney                        | ● Obtain MRI of the abdomen to assess for the progression of angiomyolipoma and renal cystic disease every 1 to 3 years throughout the lifetime of the patient  
● Assess renal function (including determination of GFR) and blood pressure at least annually  
● Embolization followed by corticosteroids is first-line therapy for angiomyolipoma presenting with acute hemorrhage. Nephrectomy is to be avoided. For asymptomatic, growing angiomyolipoma measuring >3 cm in diameter, treatment with an mTOR inhibitor is the recommended first-line therapy. Selective embolization or kidney-sparing resection is acceptable second-line therapy for asymptomatic angiomyolipoma |
| Lung                          | ● Perform clinical screening for LAM symptoms, including exertional dyspnea and shortness of breath, at each clinic visit. Counseling regarding smoking risk and estrogen use should be reviewed at each clinic visit for individuals at risk of LAM  
● Obtain HRCT every 5 to 10 years in asymptomatic individuals at risk of LAM if there is no evidence of lung cysts on their baseline HRCT. Individuals with lung cysts detected on HRCT should have annual pulmonary function testing (PFT and 6-minute walk) and HRCT interval reduced to every 2 to 3 years  
● mTOR inhibitors may be used to treat LAM patients with moderate to severe lung disease or rapid progression. TSC patients with LAM are candidates for lung transplantation, but TSC comorbidities may affect transplant suitability |
| Skin                          | ● Perform a detailed clinical dermatologic inspection/examination annually  
● Rapidly changing, disfiguring, or symptomatic TSC-associated skin lesions should be treated as appropriate for the lesion and clinical context, using approaches such as surgical excision, laser(s), or possibly topical mTOR inhibitor |
| Teeth                         | ● Perform a detailed clinical dental inspection/examination at minimum every 6 months and panoramic radiographs by age 7 years, if not performed previously  
● Symptomatic or deforming dental lesions, oral fibromas, and bony jaw lesions should be treated with surgical excision or curettage when present |

Continued
Cardiac Arrhythmia Is a Significant Problem in TSC

While arrhythmia is relatively common in individuals with TSC, the range of arrhythmic substrates is wide and not sufficiently specific to form a specific diagnostic criterion. Reported cases of arrhythmia associated with TSC from slow to irregular to fast heart rhythms. Bradycardia mechanisms have been associated with both sinus and atrioventricular (AV) nodal dysfunction. Tachycardia mechanisms have been related to atrial, accessory AV connection reentrant, and ventricular tachycardia. Ventricular preexcitation associated with abnormal AV connections has also been commonly reported and has been noted to participate in rapid potentially life-threatening anomalous AV conduction during atrial fibrillation. The mechanisms of arrhythmia have often been directly linked to the location of specific cardiac rhabdomyomas. Indeed, abnormal AV connections associated with TSC have been shown histologically to be directly related to rhabdomyomas tumor tissue connecting the atrium to the ventricle, rather than a “typical” accessory pathway.

In addition to the wide range of mechanisms of arrhythmia in these individuals, the effects of the arrhythmias can be extremely varied. Isolated atrial or ventricular ectopy may remain without symptoms for a lifetime. Bradyarrhythmia, depending on its severity, may also remain without symptoms but may result in fatigue or syncope. Differentiation of fatigue related to bradycardia from other organ system dysfunction associated with tuberous sclerosis may be challenging. Syncope may also have similar presentation to “drop attacks” and other neurologic events seen with TSC. Sustained tachyarrhythmia may result in palpitations or, in some instances, in syncope or cardiac arrest and sudden death. Developmentally delayed individuals may not report symptomatic palpitations associated with hemodynamically stable sustained tachyarrhythmia and may present with signs and symptoms of heart failure due to tachycardia-mediated cardiomyopathy. Recurrent syncope may be mistaken for seizures or “drop attacks,” and thus the warning signs of impending cardiac arrest may not be attended.

The presence of a diagnosis of TSC does not alter treatment recommendations for any arrhythmia. Observation and treatment of episodes of tachycardia as they occur, antiarrhythmic medications, catheter and surgical ablation, and implanted pacemakers and defibrillators remain options for treatment as they do in all individuals. Catheter ablation appears to have less frequent success than in those without TSC, probably due to the size of the tumor and possible participation of the entire tumor in the arrhythmia mechanism.

Table 3. Continued

<table>
<thead>
<tr>
<th>Organ System or Specialty Area</th>
<th>Recommendation</th>
</tr>
</thead>
</table>
| Heart                         | - Obtain an echocardiogram every 1 to 3 years in asymptomatic pediatric patients until regression of cardiac rhabdomyomas is documented. More frequent or advanced diagnostic assessment may be required for symptomatic patients
- Obtain an ECG every 3 to 5 years in asymptomatic patients of all ages to monitor for conduction defects. More frequent or advanced diagnostic assessment such as ambulatory and event monitoring may be required for symptomatic patients |
| Eye                           | - Perform annual ophthalmologic evaluation in patients with previously identified ophthalmologic lesions or vision symptoms at the baseline evaluation. More frequent assessment, including those treated with vigabatrin, is of limited benefit and not recommended unless new clinical concerns arise |

TSC indicates tuberous sclerosis complex; MRI, magnetic resonance imaging; SEGA, subependymal giant cell astrocytoma; mTOR, mammalian target of rapamycin; TAND, TSC-associated neuropsychiatric disorder; EEG, electroencephalography; ACTH, adrenocorticotropic hormone; LAM, lymphangioleiomyomatosis; HRCT, high-resolution chest computed tomography; PFT, pulmonary function tests; GFR, glomerular filtration rate.

Reproduced with permission from Krueger et al.6

Cardiac magnetic resonance imaging (MRI) can also be used to detect cardiac rhabdomyomas; however, its strength lies in more specific tissue characterization. It can be a useful adjunct to echocardiography in situations where it is unclear whether a cardiac tumor represents a rhabdomyoma (eg, in patients with a large solitary tumor). In addition, MRI is more accurate than echocardiography in delineating the proximity of cardiac tumors to normal myocardium and the great vessels and therefore may be a useful adjunct to surgical planning once a decision to operate has been made. It can also provide a more reliable and reproducible estimate of ventricular systolic function. Cardiac MRI in infants and young children (<8 years of age) requires general anesthesia or sedation and, hence, its use should be limited by necessity.

Alternative Imaging Modalities for Cardiac Rhabdomyomas

Cardiac magnetic resonance imaging (MRI) can also be used to detect cardiac rhabdomyomas; however, its strength lies in more specific tissue characterization. It can be a useful adjunct to echocardiography in situations where it is unclear whether a cardiac tumor represents a rhabdomyoma (eg, in patients with a large solitary tumor). In addition, MRI is more accurate than echocardiography in delineating the proximity of cardiac tumors to normal myocardium and the great vessels and therefore may be a useful adjunct to surgical planning once a decision to operate has been made. It can also provide a more reliable and reproducible estimate of ventricular systolic function. Cardiac MRI in infants and young children (<8 years of age) requires general anesthesia or sedation and, hence, its use should be limited by necessity.

The presence of a diagnosis of TSC does not alter treatment recommendations for any arrhythmia. Observation and treatment of episodes of tachycardia as they occur, antiarrhythmic medications, catheter and surgical ablation, and implanted pacemakers and defibrillators remain options for treatment as they do in all individuals. Catheter ablation appears to have less frequent success than in those without TSC, probably due to the size of the tumor and possible participation of the entire tumor in the arrhythmia mechanism.
Cardiology Changes to the Diagnostic Criteria

The presence of cardiac rhabdomyomas remains a major criterion (Table 4). There is no longer a need to specify 1 versus >1 rhabdomyoma. Importantly, because cardiac rhabdomyomas are often the presenting manifestation of TSC, it is important to emphasize the need for pediatric cardiologists to initiate and facilitate the TSC evaluation. Specifically, the pediatric cardiologist making a new diagnosis of rhabdomyomas should obtain clinical genetic testing and make the appropriate subspecialty referrals; typically human genetics and neurology, depending on available local resources. Genetic testing practices may vary by center, prompting a need to be familiar with local processes and the closest tertiary center with specialized care for patients with TSC. Clinical genetic testing should be obtained in all pediatric cardiac rhabdomyomas and most isolated or possible rhabdomyomas. Because there is benefit to early diagnosis and potential added morbidity to late diagnosis, a proactive approach is warranted.

The Management of Cardiac Manifestations of TSC

Medical Treatment for Heart Failure

Cardiac rhabdomyomas can lead to hemodynamic compromise and congestive heart failure, and while this occurrence is rare, it remains one of the most frequent causes of death among TSC children <10 years old. Heart failure occurs in 2% to 5% of infants and children with TSC-associated rhabdomyomas. Pharmacology-based therapy for congestive heart failure due to TSC-associated rhabdomyomas is typically not needed; however, on occasion, medical management for CHF, including digitalis, angiotensin-converting enzyme inhibition, and diuresis, may be indicated. When the cause of heart failure is arrhythmia, then the appropriate antiarrhythmic treatment is indicated and effective. However, when the cause of heart failure is inflow or outflow obstruction, typically “watchful waiting” is used with the anticipation that most cardiac rhabdomyomas will spontaneously regress over a period of months. If the heart failure is refractory, then surgery is indicated. When there is hemodynamic compromise in the neonate, prostaglandin E may be initiated to stabilize the critically ill newborn. A critically ill neonate with hemodynamic compromise due to cardiac rhabdomyomas at the time of diagnosis should be transferred to a tertiary center with cardiac intensive care infrastructure and the ability to perform surgery if needed.

New Treatment Modalities May Have a Role in Cardiology Management

mTOR inhibitors have been successfully used for TSC-associated tumors in different organ systems, and limited observations to date suggest they may also be efficacious in reducing the size of cardiac rhabdomyomas. Because mTOR inhibitors are not benign drugs and rhabdomyomas tend to regress, therapy should be considered only in situations of hemodynamic compromise where there is the potential to avoid surgery with their use. Given the low frequency of surgical resection for cardiac rhabdomyomas, it will be challenging to study in a controlled manner. However, there may be opportunities to use mTOR inhibitors, such as sirolimus (rapamycin) or everolimus, to induce tumor regression. Based on limited observations, there do not appear to be significant cardiovascular side effects associated with mTOR inhibitors. In general, side effects are considered manageable in adults, but because mTOR inhibitors affect the immune system and cells’ ability to grow and proliferate, there may be an increased risk for infections and malignant tumors over the long term. Case reports using sirolimus in the context of an infant with cardiac rhabdomyomas and refractory heart failure and everolimus in the context of hemodynamic instability demonstrate benefit and avoided surgery, suggesting that application in cases of malignant arrhythmia may also be therapeutic and avoid the need for invasive intervention. More studies are needed to define the role of mTOR inhibitors in this situation, as well as for arrhythmias, aneurysms, and latent left ventricular dysfunction.

Surgical Intervention for Complete or Partial Cardiac Rhabdomyoma Resection is Indicated in Rare Circumstances

Because most cardiac rhabdomyomas are asymptomatic and the natural history is spontaneous regression, surgical resection is not required for the vast majority of infants with

<table>
<thead>
<tr>
<th>Table 4. New Cardiology-Specific Recommendations for Tuberous Sclerosis Complex</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cardiac rhabdomyomas remain a major diagnostic criterion</strong></td>
</tr>
<tr>
<td>Echocardiogram at the time of diagnosis</td>
</tr>
<tr>
<td>If fetal diagnosis, then serial observation and at least 1 postnatal echocardiogram</td>
</tr>
<tr>
<td>Surveillance studies until regression demonstrated</td>
</tr>
<tr>
<td>Electrocardiogram at the time of diagnosis</td>
</tr>
<tr>
<td>Surveillance studies every 3 to 5 years</td>
</tr>
<tr>
<td>Holter monitor as indicated for appropriate signs and symptoms</td>
</tr>
<tr>
<td>Cardiology consultation at time of diagnosis</td>
</tr>
<tr>
<td>Ongoing cardiology surveillance as indicated</td>
</tr>
<tr>
<td>Medical and surgical intervention as indicated</td>
</tr>
<tr>
<td>Referral to genetics and neurology when cardiology makes initial diagnosis</td>
</tr>
<tr>
<td>Pediatric to adult transition plan with ongoing cardiology surveillance</td>
</tr>
</tbody>
</table>
Tuberous Sclerosis Cardiology Consensus Guidelines

Hinton et al

DOI: 10.1161/JAHA.114.001493

Journal of the American Heart Association

TSC. Among the 2% to 5% of cases that do present with heart failure and/or hemodynamic instability, only a small proportion require surgery.\(^\text{26}\) Surgical series have reported a rate as high as 20%, but this is likely a reflection of referral bias.\(^\text{26,59}\) Because the infant with heart failure requiring surgery may be critically ill, these patients are relatively high-risk surgical candidates. However, partial resection is typically adequate if complete excision would sacrifice vital structures or myocardial mass. Orthotopic heart transplantation can be considered in extreme cases, such as in the rare event that tumor replaces myocardium; however, the necessary medical regimen is associated with significant medical risks. Specifically, the seizure threshold is lowered, and the risk of infection and malignant cancer is increased. Excellent short- and long-term results have been reported in multiple series, but cases of early death have been reported.\(^\text{19,25,26}\) To date, late fetal surgical resection has not been reported, but ex utero intrapartum treatment technology suggests this may be feasible in select situations.

**Recommendations Expand Surveillance Efforts**

**Cardiology Changes to Surveillance Recommendations**

Due to the rise of diagnosis on fetal echocardiography, serial imaging during gestation is now indicated to monitor disease severity and postnatal imaging is indicated to confirm anatomy and determine the status of disease after birth. Surveillance is now recommended until regression is demonstrated (Table 4). Because cardiac rhabdomyomas are often the presenting manifestation of TSC, it is important to emphasize the need for pediatric cardiologists to make the appropriate subspecialty referrals, typically human genetics and neurology, depending on available local resources. Given the increasing appreciation for cardiology issues later in life, including arrhythmias, ECGs are now recommended every 3 to 5 years. A lower index of suspicion is required during adolescent ages. Importantly, increasing efforts are required to facilitate transition from pediatric to adult care.\(^\text{60}\)

**Recommendations for Imaging Surveillance**

In fetal life, echocardiography is recommended if there is a positive family history of TSC in a first-degree relative or if there is suspicion for TSC based on other criteria. If cardiac rhabdomyomas are identified, evaluation should include assessment for inflow or outflow obstruction which may lead to hemodynamic compromise postnatally, evaluation for arrhythmias and ventricular dysfunction, and evidence of hydroops fetalis. The presence of a complicating factor requires close follow-up during the pregnancy along with careful coordination and planning of prenatal and postnatal care with involvement of specialists from maternal-fetal medicine, cardiology, and cardiac surgery. Even in the absence of complicating factors, if cardiac rhabdomyomas are diagnosed or suspected on fetal echocardiography, consultation with maternal-fetal medicine and genetics is recommended to counsel the family regarding the likelihood of TSC and long-term prognosis. Because rhabdomyomas can enlarge during gestation, follow-up imaging later during gestation (30 to 35 weeks) is recommended. After birth and in the first 2 years of life, echocardiography is recommended for any child with a suspected diagnosis of TSC, because of the high correlation between the presence of cardiac rhabdomyomas and TSC in this age group. In addition, hemodynamic compromise due to outflow or inflow obstruction is most likely in this age group and can be easily assessed on echocardiography. If echocardiography is conclusive of the diagnosis of rhabdomyomas, no further imaging is recommended. In patients who are suspected but not confirmed to have TSC and have a cardiac tumor on echocardiography but the diagnosis of rhabdomyoma is uncertain, referral to a tertiary pediatric cardiac center for cardiac MRI under sedation or general anesthesia should be considered for tissue characterization. However, this decision should be made jointly by experts in cardiology, neurology, and/or genetics to consider the risk.

In typical cases, and in the absence of inflow/outflow obstruction and ventricular dysfunction, follow-up echocardiography is not recommended in the first year of life but may be considered once between 1 and 3 years of age to document regression of the tumors. Once regression of tumor size has been documented, follow-up echocardiography is not recommended, unless new cardiac concerns such as arrhythmia or syncope arise, and, in these cases, should be performed in consultation with a pediatric cardiologist. Closer follow-up should be considered in atypical cases. In patients with inflow/outflow obstruction, ventricular dysfunction, or large solitary tumors, more frequent repeat echocardiography may be necessary and should be coordinated in consultation with a pediatric cardiologist. In patients suspected with TSC beyond 2 years of age, echocardiography should be considered although the yield is significantly lower. Echocardiography is recommended if the physical examination is consistent with outflow tract obstruction (rare in this age group) or if there is concern for arrhythmia or syncope.

**Recommendations for Electrophysiologic Surveillance**

All individuals with tuberous sclerosis, regardless of age, should have a 12- to 15-lead ECG performed at the time of diagnosis. Subsequently, in an individual with tuberous sclerosis and no cardiac symptomatology, a repeat study
Tuberous Sclerosis Cardiology Consensus Guidelines

Hinton et al

Future Research Directions and Unresolved Clinical Issues

Animal Models of TSC Provide Potential Insight Into Mechanisms of Tumor Regression

Both TSC1- and TSC2-deficient mice are embryonic lethal with ventricular dysfunction potentially contributing to death.61,62 Heterozygous and conditional mice appear to recapitulate some of TSC phenotypes, with the Tsc2+/− mouse demonstrating more severe overall disease. Importantly, these mice are responsive to sirolimus.63 Meikle et al examined ventricular myocytes of mice with Tsc1 insufficiency (haploinsufficiency) conditionally restricted to the myocardium and demonstrated cardiomyopathy with cell findings reminiscent of human cardiac rhabdomyomas.64 However, most preclinical studies have not focused on cardiac findings, so evaluating the cardiac phenotype in these mice may provide special insight into early disease processes. For example, general mechanisms of hypertension may be elucidated.65 In addition, because rhabdomyomas tend to regress spontaneously, mechanistic insights into regression may be elucidated, potentially identifying new therapeutic targets. The mice would also provide a mechanism to preclinically test the benefit of mTOR inhibitors controlling for regression, which may underscore limited observations in human studies.

Unresolved Issues Warrant Consideration for Future Investigation

Several unresolved issues have been identified and require careful examination (Table 5). These research questions require substantial organization. Strategies that may enhance these efforts include future clinical studies examining mTOR inhibitor effects on the cardiovascular system. By adding cardiac end points to longitudinal clinical studies, we will gain insight into various natural history questions. Some of these issues may be addressed by using the TSC Alliance Clinical Registry, which has collected comprehensive clinical data on >1000 TSC patients from 17 centers (Table 6). The TSC Alliance is organized to function as a network for this purpose but requires subspecialty commitments from investigators at large centers where TSC patients are cared for (not necessarily participating already within the Alliance), highlighting the importance of coordinated multidisciplinary care and standardized approaches to care. Transitioning the patient from pediatric to adult cardiology care remains a challenging and important goal, with a need for careful monitoring and a low index of suspicion for latent manifestations of cardiovascular disease, including arrhythmias.

TSC is a multisystem genetic disorder characterized by variable abnormalities, such that patients carrying mutations may not fulfill clinical criteria for diagnosis, raising questions regarding familial screening. For example, does the presence of fetal cardiac rhabdomyomas warrant a recommendation for family screening, which is not presently indicated? In mutation-positive children, parents and siblings can undergo specific mutation testing as screening, but in parents and

Table 5. Cardiology-Specific Future Research Directions

<table>
<thead>
<tr>
<th>Question</th>
<th>mTOR indicates mammalian target of rapamycin; TSC, tuberous sclerosis complex.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Why do cardiac rhabdomyomas regress and other hamartomas do not?</td>
<td></td>
</tr>
<tr>
<td>Do cardiac rhabdomyomas completely resolve?</td>
<td></td>
</tr>
<tr>
<td>What is the incidence of sudden death? Malignant arrhythmia?</td>
<td></td>
</tr>
<tr>
<td>Do TSC1 and TSC2 genotypes predict cardiac phenotype or outcome?</td>
<td></td>
</tr>
<tr>
<td>Does treatment with mTOR inhibitors decrease the long-term risk of</td>
<td></td>
</tr>
<tr>
<td>arrhythmia?</td>
<td></td>
</tr>
<tr>
<td>What is the incidence of latent left ventricular hypertrophy and/or</td>
<td></td>
</tr>
<tr>
<td>dysfunction?</td>
<td></td>
</tr>
<tr>
<td>What is the incidence and natural history of lipidemia in TSC?</td>
<td></td>
</tr>
</tbody>
</table>

Table 6. Cardiology Variables Maintained in the TSC Alliance Clinical Registry

<table>
<thead>
<tr>
<th>Variable</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medical history, physical examination, family history</td>
</tr>
<tr>
<td>Current medications</td>
</tr>
<tr>
<td>ECG, CXR, echocardiogram, MRI, CT</td>
</tr>
<tr>
<td>Pathology if available</td>
</tr>
<tr>
<td>Other cardiac conditions (malformation, hypertension, lipidemia,</td>
</tr>
<tr>
<td>aneurysm)</td>
</tr>
<tr>
<td>Number, size, and location of cardiac rhabdomyomas</td>
</tr>
</tbody>
</table>

TSC, tuberous sclerosis complex; CXR, chest radiography; MRI, magnetic resonance imaging; CT, computed tomography.

every 3 to 5 years may be prudent. Evaluation of symptomatic palpitations should include cardiac event monitoring as appropriate. Episodes of “drop attacks” or “seizures” that cannot definitively exclude cardiac syncope should be evaluated with monitors with “looping” memory, either external or implanted. Particularly concerning cases of syncope or episodes in individuals with other concerning cardiac manifestations should be evaluated with invasive cardiac electrophysiology study. Sudden deaths in individuals with TSC are reported at all ages and have potentially diverse etiologies, including not only arrhythmia but also epilepsy, intratumor hemorrhage, obstructive hydrocephalus, and aneurysmal rupture. It remains unclear whether surveillance with ambulatory ECGs for occult arrhythmia but also epilepsy, intratumor hemorrhage, obstructive hydrocephalus, and aneurysmal rupture. It remains unclear whether surveillance with ambulatory ECGs for occult arrhythmia will be able to predict and prevent the arrhythmic deaths. Periodic ambulatory ECGs seem prudent until the question can be answered definitively.
siblings of phenotypically diagnosed children, it may be prudent to perform ECG in parents and both ECG and echocardiography in children <3 years old. Some studies have demonstrated that cardiac rhabdomyomas are more frequent in those with TSC2 (54%) versus TSC1 (20%) mutations. Currently, there is insufficient evidence of absolute risks to recommend surveillance by TSC1- and TSC2-associated cardiac disease. Variability in pathology or natural history based on presentation with TSC1 and TSC2 mutations is unclear but potentially clinically significant. Genetic testing will facilitate early identification and provide opportunities for disease stratification and early intervention.

Author Contributions
The authors participated in the Cardiology Working Group for the TSC Alliance Consensus Conference (Drs Hinton, Prakash, Romp, and Knilans), from which the concept and design for this report were conceived (Drs Hinton, Prakash, Romp, and Knilans). The manuscript was drafted by Drs Hinton, Prakash, and Knilans and revised by Drs Hinton, Prakash, Romp, Knilans, and Krueger. All authors approved the final manuscript.

Acknowledgments
The authors would like to thank the TS Alliance Consensus Conference organizers and Jo Anne Nakagawa, director of clinical projects for the TS Alliance, for sharing information and insights about the TSC Natural History Database.

Sources of Funding
This manuscript was supported in part by the Tuberous Sclerosis Alliance.

Disclosures
None.

References

DOI: 10.1161/JAHA.114.001493

Downloaded from http://aha.ahajournals.org/ by guest on August 27, 2017
43. Mas C, Penny DJ, Menahem S. Pre-excitation syndrome secondary to cardiac rhabdomyomas presenting as fetal supraventricular tachycardia. Jpn Heart J. 1997;38:133–137.

Key Words: rhabdomyoma • cardiac tumor • pediatrics • genetics
Cardiovascular Manifestations of Tuberous Sclerosis Complex and Summary of the Revised Diagnostic Criteria and Surveillance and Management Recommendations From the International Tuberous Sclerosis Consensus Group
Robert B. Hinton, Ashwin Prakash, Robb L. Romp, Darcy A. Krueger and Timothy K. Knilans

J Am Heart Assoc. 2014;3:e001493; originally published November 25, 2014;
doi: 10.1161/JAHA.114.001493
The Journal of the American Heart Association is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
Online ISSN: 2047-9980

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://jaha.ahajournals.org/content/3/6/e001493

Subscriptions, Permissions, and Reprints: The Journal of the American Heart Association is an online only Open Access publication. Visit the Journal at http://jaha.ahajournals.org for more information.