Cardiac rhabdomyomas and their association with tuberous sclerosis

David W Webb, R D Thomas, John P Osborne

Abstract
A search for children presenting with signs or symptoms of cardiac rhabdomyomas was made through members of the paediatric section of the British Cardiac Society in order to establish their birth incidence, presenting features, clinical course, and the frequency of a concurrent diagnosis of tuberous sclerosis. Fifteen children were identified and 12 had tuberous sclerosis (80%). Heart failure was the presentation in six, five of whom died; six presented because of a murmur and three because of arrhythmias. The prevalence of echocardiographic evidence of cardiac rhabdomyomas in a population of patients with tuberous sclerosis was established. Twenty individuals had echocardiography and eight had echodensities consistent with cardiac rhabdomyomas.

It is concluded that the minimum birth incidence for children presenting because of the effects of cardiac rhabdomyomas is 1/326 000 and a minimum of 80% have tuberous sclerosis. In a population of patients with tuberous sclerosis a minimum of 60% under 18 years have cardiac rhabdomyomas.

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Symptomatic primary heart tumours are rare in childhood. In several reviews cardiac rhabdomyomas have been the commonest lesion.1,4 Until recently these hamartomas were usually diagnosed at necropsy but with improved non-invasive imaging they are now diagnosed with reasonable certainty in infancy and childhood. Their natural history is still unclear.

They represent the earliest detectable hamartoma in tuberous sclerosis where they have been diagnosed in fetal life as early as 22 weeks' gestation. As they may be the only manifestation of tuberous sclerosis at this stage the probability that other features of the disease will develop with time is important for genetic counselling and patient management. A review of all published cases of cardiac rhabdomyomas up to 1990 found at least 50% to be associated with tuberous sclerosis.7 However the majority of cases reported are in young infants when excluding a diagnosis of tuberous sclerosis can be extremely difficult even at postmortem examination.4 A nationwide search for children presenting with symptoms and signs due to cardiac rhabdomyomas could give some indication of their minimum birth incidence and with improved survival and more widespread use of modern imaging techniques might also give a more accurate estimate of their true association with tuberous sclerosis.

Estimates of the frequency of cardiac rhabdomyomas in patients with tuberous sclerosis have been made8–10 but an estimate of their prevalence in a population based study has not previously been reported.

Methods
Members of the paediatric section of the British Cardiac Society were asked to identify infants that they had seen presenting because of the effects of cardiac rhabdomyomas. Case histories were reviewed and where possible contact with the family was made to assess recent progress. Individuals with tuberous sclerosis resident in the Bath health district were identified by a large prevalence study11 and were offered transthoracic cross sectional echocardiography. This was performed by a consultant cardiologist (RDT) on an ATL Ultramark 9 machine with a 3 MHz phased array scanner. Two dimensional examinations were made from parasternal, subcostal, and apical windows and the images recorded on videotape. Echodensities greater than 5 mm in greatest diameter that were discreet and clearly more echodense than the adjoining myocardium were considered significant.

Results
Seventy five per cent of cardiologists replied to the search, including members from all the supraregional centres in England and regional centres in Scotland. Fifteen cases (10 boys, five girls) were identified with presentation due to cardiac rhabdomyomas. Five individuals were diagnosed in utero after routine ultrasound. One was hydropic and died. After birth the remaining four each had a murmur. Of those found in the neonatal period, five had heart failure, four of whom died, while two presented because of a murmur and one because of an arrhythmia. The two postneonatal presentations were with arrhythmias (table 1). Of the five infants who died, two had left ventricular outflow tract obstruction and three had huge intramural tumours involving most of the left ventricle. The clinical features at presentation are shown in table 2. Arrhythmias were common but were less frequently the presenting feature. Three infants

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Table 1  Reason for presentation in 15 children with cardiac rhabdomyomas

<table>
<thead>
<tr>
<th>Anatomical diagnosis*</th>
<th>Neumal</th>
<th>Postnatal</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n=8)</td>
<td>(n=2)</td>
<td>(n=2)</td>
</tr>
<tr>
<td>Hydrops (1)</td>
<td>Heart failure (5)</td>
<td>Arrhythmias (2)</td>
</tr>
<tr>
<td>Murmur (4)</td>
<td>Murmur (2)</td>
<td></td>
</tr>
</tbody>
</table>

*Diagnosis was made at routine antenatal ultrasound; features given are those present at birth.
had Wolff-Parkinson-White syndrome. Twelve had tuberous sclerosis: eight were sporadic and the four familial cases were the family proband. All of the children with tuberous sclerosis who survived developed seizures and two are reported to be developing normally at 3-5 and 4 years. The remainder have some degree of mental retardation. The three cases in whom a diagnosis of tuberous sclerosis was not established were a newborn girl with a single rhabdomyoma who had no evidence of tuberous sclerosis at postmortem examination, a 6 year old boy who had not been specifically investigated for evidence of tuberous sclerosis, and a 3-5 year old girl who had normal cranial and renal ultrasound in infancy.

Tumours were multiple in 13 (87%) and involved the left ventricle in 11 (73%), right ventricle in 11, interventricular septa in eight (54%), and atria in two (13%). The diagnosis was confirmed histologically in seven cases. Three individuals had surgery, two for removal of lesions obstructing the right ventricular outflow tract and one for involvement of the mitral valve which required replacement. No lesions increased in size and at follow up in five children significant reduction in tumour size had been documented.

Twenty three individuals with tuberous sclerosis resident in the Bath health district were identified by a prevalence study.‡ Twenty con-
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Table 3 Abnormal echocardiographic findings in eight patients from a population of 20 patients with tuberous sclerosis. There were no atrial lesions seen

<table>
<thead>
<tr>
<th>Age</th>
<th>Sex</th>
<th>No of echodense areas</th>
<th>Site and size (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Left ventricle</td>
</tr>
<tr>
<td>1 day</td>
<td>M*</td>
<td>3</td>
<td>14</td>
</tr>
<tr>
<td>2-5 years</td>
<td>M*</td>
<td>1</td>
<td>8</td>
</tr>
<tr>
<td>4 years</td>
<td>M</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>16 years</td>
<td>M</td>
<td>2</td>
<td>19</td>
</tr>
<tr>
<td>18 years</td>
<td>M</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>46 years</td>
<td>M</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>9 years</td>
<td>F</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>39 years</td>
<td>F</td>
<td>2</td>
<td>23</td>
</tr>
</tbody>
</table>

*Spontaneous and complete resolution documented.

Figure 2 (A) Long axis parasternal views taken on day 1 (left) and at 6 weeks of age (right) in an infant whose mother had tuberous sclerosis. An echodense lesion is seen at the apex of the left ventricle which disappears on the second view. (B) A four chamber view of the same infant shows multiple echodensities (left) that have resolved at 6 weeks of age (right).

Sonically. While most rhabdomyomas are multiple, even a single lesion in infancy is very likely to be a rhabdomyoma and if other features of tuberous sclerosis are found the diagnosis is beyond reasonable doubt.

Infants with cardiac rhabdomyomas present with features related to the size and anatomical location of their tumour. Large obstructive lesions and those interfering with cardiac motility have a poor prognosis. In this study five of six cases presenting in cardiac failure died and all six had large left ventricular lesions. The surviving child is now 4 years old and asymptomatic but did require a period of ventilatory support at 2 weeks of age and had mitral valve replacement at 18 months for a lesion involving the valve. Three infants presented with symptoms related to supraventricular tachycardia and two of these had Wolff-Parkinson-White syndrome. Neonatal or fetal arrhythmias should alert the clinician to the possibility of a cardiac tumour. Lesions presenting with an asymptomatic neonatal murmur are those most likely to be missed. There were two infants identified who presented in this way and another who presented with a supraventricular tachycardia and Wolff-Parkinson-White syndrome, who were only diagnosed as having cardiac rhabdomyomas after they developed seizures and were found to have tuberous sclerosis. For this reason they have not been included in the analysis.

It has been difficult to establish the true relationship between tuberous sclerosis and this hamartoma. Early death, inadequate follow up, or incomplete evaluation may result in failure to diagnose tuberous sclerosis. In this study an association could not be confirmed in three cases. Two of these have not been completely investigated. The third case died on day one and had an otherwise normal postmortem examination including normal brain histology. Interestingly another infant with no clinical evidence of the disease and with normal brain histology had a mother and grandfather with tuberous sclerosis confirming the diagnosis.15 Several other cases highlight diagnostic pitfalls. Woods lamp examination was normal at 1 month in an infant who was later discovered to have multiple hypomelanic macules. Another infant without hypomelanic macules has developed a shagreen patch at 19 months. The external surface of the cerebral hemispheres appeared normal in one infant who had complete malformation of several gyri with cortical tubers on histological examination.

The prevalence of cardiac rhabdomyomas in patients with tuberous sclerosis has been estimated from hospital based samples. This
population study found 60% of such children and 20% of such adults to have echocardiographic evidence of tumours and the prevalence for the whole group is 40%. These figures support the work of Smith et al who found tumours in 58% of children and 18% of adults who had attended clinical genetic or paediatric neurology outpatient departments. We would also agree with the difficulty in interpretation of small reflectice areas and bulky papillary muscles, particularly in adults.

Our study suggests that the birth incidence for cardiac rhabdomyomas in patients with tuberous sclerosis is likely to be higher than 60% as asymptomatic tumours can diminish in size and disappear echocardiographically within weeks of birth. This potential for spontaneous regression has been noted by several authors before but not to such a dramatic degree in such a short time. It is likely that an increasing number of cardiac rhabdomyomas are going to be diagnosed antenatally. This lesion has never been shown to grow postnatally and it may be justified, in the light of our findings, to consider premature induction of labour in a fetus developing signs of cardiac failure because of a cardiac rhabdomyoma. While most rhabdomyomas appear to regress spontaneously, some infants may benefit from surgery for obstructive lesions and this should be considered at an early stage. The prevalence of mental handicap in an unbiased sample of tuberous sclerosis patients is only 38% so despite a likely diagnosis of tuberous sclerosis in children presenting because of the effects of cardiac rhabdomyomas the long term outlook is that they are more often intellectually normal than not.

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