Syncope in childhood is very common. The vast majority of episodes are benign, and are due to neurocardiogenic syncope. Only a minority are due to something potentially more serious or life threatening. The diagnosis and differentiation of benign from more serious causes of syncope is made primarily by the history. Investigations are often unfruitful. The mainstay of management in neurocardiogenic syncope is reassurance. An increase in dietary fluid and salt can be helpful. Drug treatment is reserved for those with more frequent and severe attacks. Cardiac pacemakers should be reserved for those with very severe symptoms who are refractory to drug therapy.

**Neurally mediated syncopes**

Neurally mediated syncopes are a heterogeneous group of autonomic disorders, which result in orthostatic intolerance.\(^4\) They can be divided into four main groups (table 1). Postural orthostatic tachycardia syndrome, pure autonomic failure, and multiple system atrophy are all types of chronic autonomic failure, whereas reflex syncopes are transient disturbances in autonomic control of heart rate and blood pressure.\(^5\)

Neurocardiogenic syncope can be considered as a form of reflex syncope and is by far the commonest type of syncope in childhood. It can occur at any age, but the peak age groups are toddlers and older children between the ages of 9 and 14 years. A typical history in an older child is of syncope that occurs when the child is upright, either sitting or standing. Characteristically there is a prodrome such as dizziness, nausea, and pallor, before loss of tone and consciousness. Depending on the duration and severity of cerebral hypoxia secondary to hypotension and/or profound bradycardia, the child may have an anoxic seizure and may be incontinent. An anoxic seizure is different from an epileptic seizure in that the electroencephalogram (EEG) is flat, and rather than tonic-clonic movements, there tends to be stiffening, opisthotonus, and fine twitching.\(^6\) On recovery the child may feel tired and “washed out” for some time. In toddlers, reflex syncope tends to manifest as so called “reflex anoxic seizures” or “pallid breath holding spells” with reflex asystole followed by an anoxic seizure in response to a noxious stimulus (fig 1A).\(^7\) Of note is that the distinction between “pal- lid breath holding” and “blue breath holding” is more blurred than often realised. Both are types of reflex syncope, but the mechanism behind the cyanosis is poorly understood.\(^8\)

**Table 1 Classification of syncope in childhood**

<table>
<thead>
<tr>
<th>Category</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neurally mediated syncopes</td>
<td>Reflex syncope, Postural orthostatic tachycardia syndrome (POTS), Pure autonomic failure, Multiple system atrophy</td>
</tr>
<tr>
<td>Cardiovascular causes</td>
<td>Arrhythmic, Structural, Vascular</td>
</tr>
<tr>
<td>Non-cardiovascular pseudo-syncopes</td>
<td>Epileptic, Psychogenic</td>
</tr>
</tbody>
</table>

**Diagnosis of syncope**

Investigations for syncope in childhood will almost always be normal, and the key to the diagnosis is a detailed and careful history.\(^2\) When a careful history is taken, it will usually be clear to which of the three categories the syncope belongs. In the vast majority of cases, the syncope will be neurally mediated. There are, however, a
number of “warning bells” from the history that should indicate a potentially more serious or life threatening cause (table 2).

INVESTIGATIONS FOR RECURRENT SYNCOPE

Probably the most important investigation for recurrent syncope is a 12 lead electrocardiogram (ECG), primarily to exclude a long QT interval. Pre-excitation, heart block, or ventricular hypertrophy can also be diagnosed from an ECG. Although the ECG will almost always be normal in children with recurrent syncope, a normal ECG is an important finding. In older children, if symptoms are related to exercise, an exercise test should be performed in the hope of inducing symptoms. In reality, symptoms rarely occur during the test. Holter monitoring is usually unhelpful, as symptoms almost never occur in the 24–48 hour period while the monitor is worn. Cardiac event monitoring has a theoretical advantage over Holter monitoring in that the monitors can be worn for a longer period of time, such as weeks or months. The aim is to try to record the ECG during symptoms, but inevitably the monitor never seems to be worn at the time an event occurs. Unless the child has other cardiac signs or symptoms, or any of the warning bells from the history, an echo will almost certainly be normal. Although an EEG is often performed on children with syncope to “exclude epilepsy”, this is rarely helpful. Even in children with epilepsy the EEG will usually be normal between attacks, and a percentage of children without epilepsy will have a frankly “epileptic” EEG. Intracerebral causes of syncope are very rare in childhood and would usually be associated with other neurological signs or symptoms; thus magnetic resonance imaging or computed tomography scan is usually an expensive waste of time.

A suggested approach to the investigation of the child with recurrent syncope is as follows. A 12 lead ECG should be

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Table 2  Syncope “warning bells”

| Syncope in response to loud noise, fright, or extreme emotional stress |
| Syncope during exercise |
| Syncope while supine |
| Family history of sudden death in young person <30 years old |
| Syncope with an “odd” history |
performed primarily to exclude a long QT. If there are any of the “warning bells” from the history or if there are any other cardiac or neurological signs or symptoms, appropriate cardiac or neurological investigations should be undertaken. If there is a good history for neurally mediated syncope and the ECG is normal, usually no further investigation is required. Further investigation may be considered for those who have very severe or frequent attacks, who are in need of reassurance, or where the history is not entirely clear. Where there is a good history for neurally mediated syncope, tilt testing is probably the most productive investigation for older children, and cardiac event monitoring for the toddler.

TILT TESTING
Tilt testing can be successfully performed in children from the age of 6 years. There is no standardised protocol for tilt testing and protocols vary in terms of duration of tilt, degree of tilt, and whether or not drugs such as isoprenaline are given. Our own protocol is very simple and tolerated by most children. The child rests supine for 15 minutes and is then tilted to 60 degrees head-up for a maximum of 45 minutes. During this time the blood pressure is continuously but non-invasively monitored using the Finapres system, and a three lead ECG continuously recorded. The main “side effect” of the test is boredom, and younger children especially have a tendency to become restless. Using this protocol, approximately 50% of children with a good history for neurally mediated syncope will have a positive tilt test. The use of drugs increases the sensitivity of the test, but reduces its specificity and makes the test more unpleasant for the child. Whatever tilt test protocol is chosen, the three important requirements appear to be a period of supine rest before tilting, to tilt to between 60 and 80 degrees to reduce false positive and negative responses, and to tilt for at least 40 minutes of drug free period.

In those who have a positive tilt test, the most common response is a combination of hypotension and bradycardia prior to syncope. Hypotension with no significant change in heart rate is the next most common positive response. The least common positive response is asystole prior to syncope, often termed a “cardioinhibitory” response.

Another type of reflex syncope, which has only been recognised relatively recently, is cerebral vasoconstrictive syncope. In this condition syncope is caused by cerebral vasoconstriction without hypotension and bradycardia. Recording an EEG or measuring cerebral blood flow during a tilt test in individuals who have symptoms despite normal blood pressure and heart rate should help make the diagnosis.

In situations where it is claimed that syncope is occurring several times a day every day, admission to hospital for observation with continuous ambulatory ECG and EEG monitoring is usually the best approach. Almost always the ECG, EEG, and measured blood pressure will be found to be normal during the “syncopes” and a diagnosis of psychogenic pseudosyncope can be confirmed.

For toddlers with reflex asystolic syncopes, cardiac event monitoring can help confirm diagnosis, although an ECG recording during symptoms is only likely to be obtained in those few who have very frequent attacks (fig 1A).

In infants who have recurrent and severe episodes of syncope that only have their onset in the presence of a particular parent or guardian, factitious or induced illness should be considered. This is particularly likely if some of the syncopes are shown to other adults—relatives or hospital staff—before the infant has recovered. The child should be admitted to hospital for continuous ambulatory EEG monitoring together with cardiac event monitoring and overt video surveillance. The parent or guardian should be asked to activate the cardiac event monitor at the onset of any syncopal event. Typically in the situation of induced syncope e.g. by smothering, there tends to be a significant delay between the onset of EEG and ECG changes and activation of the monitor (fig 1B). This is different from most spontaneous syncope episodes, where activation of the monitor tends to occur within seconds of the onset of ECG and EEG changes (fig 1A).

Occasionally the distinction between neurally mediated syncope and potentially serious arrhythmia remains unclear, for example, in a child with a borderline QT interval or slightly worrying history. In this situation, an implantable Reveal monitor (Medtronic Inc., USA) can be useful (fig 2).

MANAGEMENT OF NEURALLY MEDIATED SYNCOPE
The mainstay of therapy is reassurance, specifically that the episodes are not caused by epilepsy or a cardiac problem. Advice should be given to drink plenty and to increase dietary salt. When prodromal symptoms are experienced, manoeuvres such as crossing the legs and folding the arms, especially when standing, help to maintain blood pressure. Often with the above simple measures of reassurance, fluid, posture, and salt, symptoms will improve significantly. In toddlers with reflex asystolic syncope, parents should be advised to place the child in the recovery position and avoid the natural tendency to pick up the child.

The likelihood of further syncopal attacks depends on the number of episodes of syncope prior to presentation. For those who present with frequent syncope or continue to have syncope despite the above simple measures, drug therapy should be considered. There are many pharmacological agents available, but no drug has been adequately evaluated by randomised clinical trials. Probably all have a degree of placebo effect. Fludrocortisone and β blockers are the most favoured first line drugs, with relatively few side effects. A reasonable first approach is fludrocortisone at a dosage of 100 µg/day. This is effective in most children in reducing frequency and severity of syncope. Occasionally the drug is not tolerated as a result of fluid retention and weight gain. If symptoms continue despite fludrocortisone, addition of a β blocker can be helpful. Serotonin uptake inhibitors such as fluoxetine hydrochloride, and vasoconstrictors such as midodrine, a α agonist, are currently under clinical review, but their safety has not been established in children.

It would seem sensible to reserve them for those who continue to have symptoms despite first line therapies. An alternative or adjunct to drugs is biofeedback techniques, including tilt training and active tension. For toddlers with neurocardiogenic syncope, iron therapy might be helpful. Iron deficiency may be more important in cyanotic rather than pallid breath holding spells,
but there is often overlap between the two. Nevertheless it is
reasonable to look for and treat any iron deficiency in toddlers
with neurocardiogenic synapses. Atropine can be effective in
reducing severity and frequency of syncope, but side effects,
including loss of concentration, dry mouth, gastrointestinal
upsets, and behavioural problems are common.24

CARDIAC PACING

The use of cardiac pacing for neurocardiogenic syncope
remains controversial.21 The idea behind pacing is that it
should prevent any significant bradycardia or asystole that
might contribute to the hypotension that ultimately results in
syncope. It would not be expected to prevent hypotension that
can be caused by vasodilatation. There is little evidence, however,
that profound bradycardia occurs during most spontaneous
attacks, even if it occurs during tilt testing.

A number of studies have shown a benefit from pacing for
neurocardiogenic syncope, but only one of the studies
successfully addressed the issue of placebo.20-28 As cardiac pac-
ing is a significant commitment in a young person, it should
be reserved for those who have severe, frequent attacks and in
whom drug therapy has failed or is declined. While the ques-
tion remains as to whether children with neurocardiogenic syncope but without demonstrable asystole will benefit from pacing, it would seem sensible to reserve pacing for those who
have a recorded asystole or profound bradycardia during a
typical attack. One approach is to implant a Reveal monitor in
any child with frequent neurally mediated syncope in whom a
pacemaker is being considered. The Reveal allows a more
accurate assessment both of the frequency of syncope and
whether asystole or profound bradycardia occurs during
spontaneous syncope. It has been our experience that implanta-
tion of the Reveal can in itself have an astonishingly curative
effect, perhaps by working as a placebo or as a form of
biofeedback, and may even circumvent the need for a
pacemaker.

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