Abstract

Supraventricular tachyarrhythmia (SVT) is the most common tachyarrhythmia in pediatric patients and accounts for more than 90% of pediatric arrhythmias. As many as 16 different mechanisms of pediatric SVT exist; we focused in this review article on the three most common groups of SVT mechanisms in pediatrics: atrioventricular re-entry tachyarrhythmia (AVRT), atrioventricular nodal re-entry tachyarrhythmia (AVNRT), and atrial tachyarrhythmia (AT). Many methods have been implicated in the diagnosis of SVT like full history taking regarding onset and frequency of palpitation, syncope, chest pain, dyspnea, heart failure, drug intake and physical examination. Children with suspected SVT must be subjected to resting 12-lead surface ECG and ECG during the attack. Echocardiography must be done for detection of any structural heart diseases. Moreover, recent techniques like Electrophysiologic study (EPS) is the most definitive diagnostic procedure. Many therapeutic strategies like drugs and radiofrequency ablation (RFA) have been succeeded to provide a significant reduction of the risk of SVT, but RFA offers the prospect of cure of the arrhythmia and avoidance of drug-associated side effects. It is often becoming the first line of therapy for many children with SVT, as it has a relatively low morbidity and mortality, and it results in a low rate of recurrence of SVT.
After the impulse is travelled down via the slow pathway, the fast pathway is now unblocked and the impulse can spread retrograde via the fast pathway. So, a re-entrant circuit is formed [7]. (Figure 2).

AVNRT accounts for 15% of cases of pediatric SVT, mostly present over the age of 5 years and almost completely absent in infants. Females were more than males in cases diagnosed as AVNRT or AT [6].

Atrial tachyarrhythmia (AT): AT is mostly due to abnormal automaticity, may be formed by remnant embryonic cells with automatic qualities, causing abnormal impulses that come away from normal sinus node but still within the atria. Although it is a rare condition, but still the third common mechanism of tachyarrhythmia occurring in children, if remains undetected can result in a dilated cardiomyopathy [7].

History and clinical presentations of svt in pediatrics

The clinical presentation varies with the age of the patient; generally, it differs in infancy from that in an older child. In infancy, most attacks occur before 6 months of age, and may often less than 4 months. They may be undetected or associated with pallor. The parents may take their child to the doctor with a complaint of poor feeding, or vomiting. The rapid breathing or ashen color may not be observed by them. By the time, the pediatrician sees them: there is usually significant cardiorespiratory distress. The infant will often have signs of heart failure. Occasionally frank circulatory shock will be present [8].

Occasionally, a young patient may have palpitations, chest or abdominal pain, occasionally syncope, particularly with exertion may occur, but the presence of congestive heart failure beyond infancy is extremely uncommon [9].

Physical examination of svt in pediatrics

Following the history, the physical examination should be performed: unexplained hypothermia, with poor perfusion, a picture of sepsis but with negative cultures. In the older infants, irritability associated with vomiting and sometimes diarrhea (uncommon). In approximately 20% of infants, SVT is detected during the routine examination without symptoms [10].

Auscultation should be done to reveal the presence of structural heart disease, heart rate and rhythm should be noted. Further evaluation will depend on the severity and frequency of symptoms, the age of the child, and the presence of structural heart disease. Infants with SVT may have a heart rate range from 220 to 300 BPM. Older children generally have slower rates which may count from 180 to 240 Bpm. Untreated SVT can result in congestive heart failure (CHF) within 24 to 48 hours. Termination of the attack by vagal maneuvers may suggest a reentrant tachyarrhythmia involving AV node [5].

Diagnostic investigations of svt in pediatrics

Recording a 12- Lead Electrocardiography (ECG) at rest should be done and examined for any abnormal rhythm, delta waves, abnormal QT interval, sinus tachyarrhythmia, or any sign of underlying structural heart disease (Figure 3) [11]. ECG can clearly diagnose about 80% of AVNRT and AVRT, but incorrectly categorize approximately 20% of cases of SVT; therefore, the ECG cannot serve as the sole means for determining the mechanism of tachyarrhythmia [12].

Another diagnostic tool is worn by the patient named Holter monitor. It can record continuous ECG tracing for 24 to 48 hours, having the same idea as classic ECG wires and electrodes but it is a portable device that provides information about symptoms that can occur during the day [7].

In selected patients with occasional complaints (i.e. less than two episodes of SVT per month) and associated with disabling symptoms (i.e. hemodynamic instability) can be diagnosed by implantable loop recorders. If the clinical history is not enough or other measures have failed to identify the SVT mechanism, then recordings and stimulation can be done via Transesophageal atrial loops for these patients for diagnosis or to provoke paroxysmal tachyarrhythmia [13].

The possibility of structural heart disease should be excluded by Echocardiography examination in children with identified attacks of SVT, which is not possible to be detected via physical examination or resting 12-lead ECG [14].

Other investigations can be done to exclude other causes included in the differential diagnosis of SVT. Chest x-ray with lateral and anteroposterior views is done to diagnose cardiomyopathy and CHF. Laboratory tests which include; serum electrolytes to diagnose any imbalance leading to abnormal cardiac rhythm, complete blood cell count (CBC) with differential to exclude infections or anemia, also, screening of...
Management of SVT in pediatric age group

The dealing with SVT can be done in two ways: acute termination of the attack of tachyarrhythmia and prevention of recurrences (chronic therapy). The acute treatment begins with documentation of the arrhythmia with a 12 lead ECG, assessment of haemodynamic status and recognition of the arrhythmia mechanism. Chronic therapy is based on the SVT mechanism, the patient’s age, and the frequency and patient’s complaint during the attack as well as access to medical care and sophistication of the patient caretaker (Table 1) [15].

Acute management of SVT: For the haemodynamically tolerated episode, vagal physical maneuvers are to be used first. The patient may be taught to do Valsalva maneuver (the most potent physical maneuver), induce vomiting, dive the face in the iced water for 10–20 seconds, or carotid sinus massage may be done, an ocular pressure is not recommended for children. These physical maneuvers are potent in terminating re-entry SVT; mostly AVRT and to less extent AVNRT [11].

An ultra–short–acting drug (Adenosine) is highly effective in terminating AV node–dependent SVT, but it is of limited value in the diagnosis of AT. Vagal maneuvers and adenosine produce transient AV node block and result in sudden termination of SVT in any tachyarrhythmia involving AV node but not in AT (Figure 4) [16].

The use of Beta blockers or long–acting calcium channel blockers is of value especially for patients with frequent atrial or ventricular premature beats which may act as a cause of recurrence of SVT but they must be used with great caution as they may potentiate hypotension [11].

In a haemodynamically unstable child with severe hypotension, electrical cardioversion by synchronized DC shocks is the treatment of choice. It is a safe and effective procedure in the majority of patients [2].

Long term (chronic) management of SVT: The decision to initiate chronic therapy in infants and children is based on symptoms, efficacy rates, safety issues, expected response to therapy and frequency of episodes. The range of therapeutic options aimed at treating disorders of heart rhythm has expanded tremendously over the past decade; the complexity of actions produced by antiarrhythmic agents occasionally leaves the clinician frustrated by attempted at predicting the clinical results [2].

A child with SVT should be viewed globally while putting a long-term therapeutic plan, as this may affect him through his life–long. A pharmacologic therapy is used to provide symptom relief. The usual selection of drugs (in order) is propranolol, verapamil and amiodarone.

Digoxin is especially considered in cases with AVNRT, while amiodarone in combination with propranolol or/and verapamil is effective in cases with AT [11].

Conditions that warrant consideration for referral to an electrophysiologist include symptomatic SVT with debilitating or severe symptoms (e.g., syncope, dyspnea), and intolerance of or resistance to drug therapy. Patients who also benefit from a referral to an electro–physiologist include those who require recurrent hospital admissions or emergency room visits for tachycardia termination, those who require treatment with drugs other than beta blockers or calcium–channel blockers, and those who are simply not interested in drug therapy [15].

Radiofrequency ablation (RFA) using the traditional radiofrequency energy to heat and destroy the site of origin of the SVT mechanism has been more widely applied in children since the 1990s [3]. It is now considered as the first line of management for many children with SVT and offering the prospect of cure of the arrhythmia and avoidance of drug–associated side effects. Success rates for ablation are as high 95% and characterized by low morbidity, low mortality, and a low rate of SVT recurrence [17] RFA may have some complications resulting from the vascular access (i.e. hematomas, perforation of the aorta, deep venous thrombosis), or during catheter manipulation (i.e. valvular damage, perforation of the coronary sinus or myocardial wall), or inaccurate ablation (resulting in i.e. AV node block, myocardial perforation, coronary artery spasm, transient ischemic attacks) [18].
Successful elimination of SVT substrate results in improving the pediatric quality of life regarding physical, emotional, social, school and psychosocial functions [19].

Conclusion

SVT is a common pediatric age group problem. Many methods have been implicated in the diagnosis of SVT like detailed history taking and physical examination. ECG and echocardiography must be done to children suspected to have SVT. Electrophysiologic study (EPS) is a recent technique for definitive diagnosis. RFA is a definitive therapy for many children with SVT. However, further studies are mandatory in order to provide more novel therapeutic agents for complete protection of SVT and decreasing the complications of already used management procedures.

Core tip

This manuscript discusses the most common types, recent strategies in diagnosis and management of supraventricular tachycardia in pediatric age group.

References


Copyright: © 2016 Abo-Haded HM, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.