Appropriate Initial Adenosine Dosage for Children with Supraventricular Tachycardia

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Abstract: SVT can be defined as an abnormally rapid heart rhythm originating above the ventricles, often (but not always) with a narrow QRS complex on ECG. Re-entrant SVT is the most frequent mechanism of tachycardia in childhood (>90%). Adenosine is widely recognized as the first line of pharmacologic treatment for SVT. Since, the initial clinical trials in late 1980s, the adequate dosage of adenosine have been questioned and current guidelines offer confusing advice about adenosine dosage in children, with very heterogeneous dosage recommendations. In this review of the pediatric literature we show that in several studies the optimal dose of adenosine is higher than the 0.1 mg/kg recommendation. Because adenosine is a drug with few adverse effects owing to its very short half-life, it seems advisable to recommend a higher initial dose, at least 200-300 mcg/kg.

Keywords: Supraventricular tachycardia; Adenosine; Children; Antiarrhythmic drugs.

1. SUPRAVENTRICULAR TACHYCARDIA (SVT)

SVT can be defined as an abnormally rapid heart rhythm originating above the ventricles, often with a narrow QRS complex on ECG. SVT is the most common tachyarrhythmia requiring treatment in childhood, with an estimated incidence of 1/250 to 1/1000 children[1,2]. Re-entrant SVT is the most frequent mechanism of tachycardia in childhood (>90%)[3]. In infants and children it is most commonly the result of electric conduction through accessory pathways between the atria and the ventricles (atrioventricular re-entrant tachycardia, mostly Wolff-Parkinson-White syndrome)[2]. In other cases, it is secondary to intranodal re-entry at the AV node, which is more common in adolescents and adults (atrioventricular nodal re-entrant tachycardia). Both mechanisms utilize AV node to allow the electrical impulse to go on repetitively between the ventricles and the atrium, resulting in tachycardia[1](Figure 1).

Reentry Paroxysmal Supraventricular Tachycardia

Figure 1. Mechanism for reentrant paroxysmal supraventricular tachycardia. (A) A premature atrial complex (PAC) occurs and is blocked in a fast pathway, but it can propagate down the slower pathway. (B) By the time the electrical signal reaches the end of the slow pathway, the fast pathway has repolarized, and retrograde conduction of the wave occurs. (C) The wave then returns down the slow pathway, setting up a closed circuit that is self-sustaining.
The first step in the management of SVT is to determine whether the patient is hemodynamically stable[4]. If SVT is sustained and there is any indication of instability electrical cardioversion should be performed urgently. If not, conservative measures should be applied that include both non-pharmacologic (vagal manoeuvres) and pharmacologic measures[2]. Vagal manoeuvres are a number of physical manoeuvres increase the resistance of the AV node to transmit impulses (AV nodal block), principally through activation of the parasympathetic nervous system, conducted to the heart by the vague nerve.

2. ADENOSINE

Adenosine is widely recognized as the first line of pharmacologic treatment for SVT[4,5]. It is an endogenous purine nucleotide (6-amino-9-beta-D-ribofuranosyl-9-H-purine) present in all cells of the body (Figure 2).[6,7] that has an important role in regulating the myocardial oxygen supply-demand balance. Adenosine achieves this through coronary vasodilation, decreasing myocardial contractility, and depressing automaticity and conduction within the sinus and atrioventricular (AV) nodes. [6,7].

As an antiarrhythmic agent, adenosine has several unique properties as it is an intermediate metabolite, it has a very short half-life (less than 1.5 seconds), its effects are mediated by specific membrane purine (cell surface A1 and A2 adenosine) receptors coupled to guanine nucleotide binding proteins (G proteins), and it has site specific actions in the myocardium with important differential effects in supraventricular and ventricular tissue (Figure 3). Activation of cardiac A1 receptors has a myocardial depressant effect. A1 receptor activation mediates inhibition of cAMP-mediated calcium influx and enhances potassium conduction. As a result, there is (1) hyperpolarisation and resultant slowing of the sinus node, (2) depression of action potential in the AV node, (3) mild inhibition in inward calcium current (ICa) in atrial myocytes and (4) suppression of ICa current in ventricular myocytes, leading to negative dromotropic, chronotropic and inotropic effect. These mechanisms explain how adenosine terminates SVTs after intravenous infusion[6,7]. A2 receptors which are found in endothelial cells and vascular smooth muscle and are responsible for the enhancement of adenylylcylase activity and increased cyclic AMP which relaxes smooth muscle and produces vasodilation (Figure 3)[6,7].
The above cardiovascular effects make adenosine a very useful and effective agent in controlling SVT, and it is widely accepted as the first line treatment for rapid termination of the short-term episode of hemodynamically stable SVT in infants and children which have failed to respond to vagal manoeuvres.

3. INITIAL DOSAGE OF ADENOSINE IN CHILDREN

Adenosine is very effective in reverting SVT to sinus rhythm, terminating SVT in 90-99% of patients when administered [5,8,9]. However this response is usually achieved at various incremental doses, not at the initial dosage in children. Thus the initial dosage of adenosine in SVT is a highly controversial issue in pediatric patients. Early experimental work and guidelines suggested adenosine doses of 37.5–50 µg/kg[5,10,11]. Later studies supported evidence of higher doses of adenosine for optimum results in children and showed that the mean effective dose was around 150 mcg/Kg[8,12,13]. Current guidelines offer confusing advice about adenosine dosage in children, with very heterogeneous dosage recommendations. The starting recommended dosage varies between 50-100 mcg/kg increasing by 50-100 µg/kg every 2 min to a maximum dose of 300-500 µg/kg if no effect is seen.[4,14].

The pediatric literature shows that these current recommended adenosine initial doses for acute management of SVT are ineffective in the vast majority of cases. Thus the reported response to initial recommended dosage of 50 mg/Kg and 100 mg/Kg is only 6-16% and 22-36% respectively[9,13,15]. This minimum effective dose approach is designed to minimize to minimize side effects, but it is important to bear in mind that if multiple doses are required to control SVT, it may result in increased exposure to the drug and possibly increased side effects. Consistent with this Losek et al. found that a higher total dose was needed when the first administered dose was low[15]. The most important implication of this too low starting dosage is that paediatricians could lose confidence in what is a very effective drug and also the delay of control for a potentially life-threatening complication. For the daily clinical practice it is important to know that the maximum recommended dose for children of 500 mg/kg and neonates of 300 mg/kg is for a singly administered dose, not for the resulting cumulative dose after summing each dose of adenosine administered, so the cumulative dose should not precludes further administration.

A high initial adenosine dosage approach constitutes probably the best management for SVT in children. In all pediatric adenosine studies the average effective dose was consistently reported to be >150 µg/kg, so it seems that this could be the initial dosage[8,9,12,13,16]. The response to doses up to 200 µg/kg has been reported to be 53%-64%, and the mean effective dose of adenosine to control SVT is reported to be 160 mcg/Kg[8,12,13,16]. Also a reduction in risk of unsuccessful cardioversion of 35% (95% CI 28% to 43%) using the higher adenosine dose has been demonstrated[9].

The initial adenosine dosage differs in small infants and children[1,13,16]. Also the average dose of adenosine required for tachycardia termination in children has been reported to be higher than in adults[13]. A dose of 50 mg/kg is effective in 10% of infants and children and 100 mg/kg is effective in 25% of infants and 50% of children. A dose of 150 mg/kg is more effective in children (around 80%) than in infants (around 35%)[9,13,16]. This difference may be explained by the difference in weight to body surface area ratio. Other possible explanations for the lower response in infants include smaller cannulae limiting the injection rate and the fact that babies tend to be more ill at presentation and may have prolonged circulation times[1]. Adenosine has a very short half-life of only 8–10 s, so it is imperative that the drug is rapidly delivered to its intended site of action, the atrioventricular node and therefore adenosine has to be administered through a vein of good calibre, preferably in the upper limbs, as close as possible to the heart. The infusion rate will be the fastest possible, and the subsequent flushing with physiological saline must be performed to accelerate the arrival of the drug in the coronary circulation. The smaller gauged cannulae, which are frequently chosen when treating children in view of their smaller veins and in order to reduce the pain associated with cannula insertion, do not permit rapid delivery of adenosine and do not facilitate sufficient flow rates and therefore reduce the chance of cardioversion, necessitating repeated and higher dosing.

Finally high initial dosage of adenosine is a safe practice[8,9,12,13,16]. After intravenous (IV) administration, adenosine is almost completely eliminated after a single pass through the
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Coronary circulation. It is rapidly taken up from plasma into red blood cells and blood vessel endothelial cells where it is converted to adenosine-5′-monophosphate, inosine, or both. In physiological concentrations, adenosine has a half-life of 0.6-1.5 seconds, but in therapeutic doses the effects persist for 20-30 seconds[2,13]. Thus although side effects are common (cumulative clinical side effects occur in about 30% of patients) they are usually transient and mild, and do not require clinical intervention. Side effects include flushing, headache, chest discomfort, cough, and occasionally hypotension. No significant or sustained adverse effects when giving adenosine in therapeutic doses are usually reported in children. Short autolimited runs of nonsustained ventricular tachycardia, sinus tachycardia, or atrial fibrillation may also be observed after termination of tachycardia, but only sporadic life-threatening tachyarrhythmia generated or accentuated by adenosine are reported[17]. In addition, transient sinus bradycardia and AV block may also occur. Prolonged sinus arrest or AV block is uncommon but is dose-dependent and is seen in patients with underlying conduction disturbances.

4. CONCLUSIONS
Although most patients with SVT have a mild course, a few may develop cardiac failure and cardiogenic shock if prolonged and/or not diagnosed. Adenosine is an effective medicine in treating SVT in children and the current recommended starting doses of adenosine seems to be low.

A review of the pediatric literature shows that in several studies the optimal dose of adenosine is higher than the 50-100 mcg/kg recommendation. Because adenosine is a drug with few adverse effects owing to its very short half-life, it seems advisable to recommend a higher initial dose, at least 200-300 mcg/kg.

The type of venous access; the speed of administration; the flushing of the line; age; and cardiovascular status, are important factors influencing in the efficacy of adenosine to abort SVT.

REFERENCES


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