Pediatric patients’ head-up-tilt-test with pharmacological challenge, it is safe?

Mesa inclinada en pacientes pediátricos con reto farmacológico, ¿es seguro?

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Abstract

Syncope in pediatrics represents an important cause of visits to the emergency units. For this reason, excluding a cardiac or malignant origin is essential at the time of the initial approach to determine what is the next step in management, or if they need to be referred to a pediatric cardiologist and/or electrophysiologist. Vasovagal syncope is the most frequent cause of syncope in pediatrics, in which a detailed clinical history is enough to make the diagnosis. If no diagnosis is concluded by the history, or if it is necessary to define the hemodynamic response of the patients, the head-up-tilt-test is indicated; this will trigger syncope due to an orthostatic stress caused by the angulated table (passive phase). If a negative response remains, it can be followed by a pharmacologic challenge to trigger the hemodynamic response, which is still controversial in pediatrics. The pharmacologic challenge increases the sensitivity with a slight reduction in test specificity. Although there is not a specific drug for the challenge in pediatric patients yet, the most commonly drugs used are nitrates and isoproterenol, the latter related to a great number of adverse effects. Sublingual administration of nitrates in the challenge has been proven to be ideal, effective, and safe in this specific age group. The aim of this article is to make a literature search to demonstrate the effectiveness and safety of the pharmacologic challenge during the head-up-tilt-test in pediatrics, emphasizing a study conducted at the National Institute of Cardiology with isosorbide dinitrate.

Key words: Head-up-tilt-test (tilt-test). Isosorbide dinitrate. Syncope.
Introduction

Syncope is a common problem in pediatrics that accounts for 1% of visits to the Pediatric Emergency Department. Annual incidence is calculated at 1.25/1000 pediatric patients, and it is more common in children older than 10 years. Recurrent syncope can markedly affect patient quality of life due to the stage of development he/she is in.

Syncope is generally defined as transient loss of consciousness secondary to cerebral hypoperfusion, characterized by its sudden nature, and sometimes it is preceded by short-lived prodromes (pre-syncope symptoms), almost always with spontaneous and complete recovery without post-event states of confusion. However, cerebral hypoperfusion can generate seizure-like movements, especially of the tonic-clonic or myoclonic type, followed by short periods of amnesia; this can confuse the doctor during interrogation, particularly when the episode was not witnessed by someone else that helps clarify the clinical scenario of the syncope (Fig. 1).

On the other hand, pre-syncope is defined as a sensation of loss of consciousness without reaching it and is characterized by secondary symptoms of both hypoperfusion and parasympathetic stimulation, such as sweating, paleness, nausea, vomiting, blurred vision, and among others. Although the definition of syncope is not clearly established when talking about a neuro-cardiogenic origin secondary to a transient dysfunction of the autonomic nervous system that occurs in the pediatric population, there are still several terms for refer to it, including “fainting,” “vasovagal syncope (VVS),” “reflex syncope,” and “neuro-cardiogenic syncope.”

Among the causes of syncope, vasovagal etiology affects 15%-25% of children and adolescents and is the most common cause of syncope in the pediatric population (61-80% of cases). Due to the complexity of syncope, it is essential for a detailed medical history to be obtained during patient initial approach, which can be diagnostic for VVS without any other complementary study being required. Similarly, an extensive approach is necessary to rule out other causes such as primary origin, including heart conditions of obstructive or arrhythmogenic types, or neurological, and metabolic diseases (Table 1).

<table>
<thead>
<tr>
<th>Causes of syncope ordered by frequency</th>
<th>Example</th>
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<tbody>
<tr>
<td>Vasovagal syncope</td>
<td>Typical: prodrome symptoms</td>
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<tr>
<td></td>
<td>Atypical: no prodrome symptoms</td>
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<tr>
<td>Breath-holding spells</td>
<td>Brief periods in which the children stop breathing, with duration of up to 1 min, secondary to emotions such as anger, surprise, or injury</td>
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<tr>
<td>Heart-related</td>
<td>Primary electrical disturbances</td>
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<td></td>
<td>Long QT syndrome</td>
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<td></td>
<td>Short QT syndrome</td>
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<td></td>
<td>Brugada syndrome</td>
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<td></td>
<td>Wolff-Parkinson-White (WPW) syndrome</td>
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<td></td>
<td>Catecholaminergic polymorphic ventricular tachycardia</td>
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<td></td>
<td>Heart structural alterations</td>
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<td></td>
<td>Hypertrophic cardiomyopathy</td>
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<td></td>
<td>Coronary artery abnormalities</td>
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<td></td>
<td>Right ventricular arrhythmogenic dysplasia</td>
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<td></td>
<td>Aortic valve stenosis</td>
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<td></td>
<td>Dilated cardiomyopathy</td>
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<td></td>
<td>Pulmonary hypertension</td>
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<td></td>
<td>Acute myocarditis</td>
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<tr>
<td>Neurological</td>
<td>Seizures</td>
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<td></td>
<td>Panayiotopoulos syndrome</td>
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<td></td>
<td>Vascular episodes</td>
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<td></td>
<td>Subclavian steal phenomenon</td>
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<tr>
<td></td>
<td>Vertebrobasilar insufficiency</td>
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<td></td>
<td>Cerebrospinal fluid impaired circulation</td>
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<td></td>
<td>Third ventricle colloid cyst</td>
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<td></td>
<td>Tumors of the posterior fossa</td>
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<td></td>
<td>Vertigo crisis</td>
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<td></td>
<td>Basilar migraine</td>
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<td></td>
<td>Narcolepsy/catataplex</td>
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<td></td>
<td>Bleeding, dehydration, hypoglycemia, electrolytic abnormalities. Conversion syndrome, somatization, Munchausen syndrome/pretending the disease (malingering)</td>
</tr>
<tr>
<td></td>
<td>Anxiety and hyperventilation syndrome</td>
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<tr>
<td>Unknown origin</td>
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</tbody>
</table>

Table 1. Common causes of transient loss of consciousness in children

Palabras clave: Dinitrato de isosorbida (DNIS). Prueba de mesa inclinada. Síncope.
When there is diagnostic doubt and other causes of syncope have been previously ruled out, it is necessary to perform a head-up-tilt-test to verify the vasovagal origin through a vasodepressor or cardioinhibitory response\(^2,9\).

**Head-up-tilt-test**

Head-up-tilt-test is a useful, low cost tool that allows the diagnosis of reflex syncope to be established when it is uncertain. Although most studies are carried out in adults, its safe use has been documented in the pediatric population\(^10\). This test should not be used with the purpose to induce or assess the response to pharmacological or interventional treatment in the patient with VVS due to low reproducibility of the test for triggering the same hemodynamic response in a second test\(^2,6\).

Initially, head-up-tilt-test only included a passive phase, and orthostatic stress was triggered by tilting the table. Subsequently, the so-called “method of symptoms” was used, which through carotid massage increased vagal tone until culminating in syncope secondary to a vasodepressor or cardioinhibitory response\(^11\). Finally, the provocation or pharmacological challenge phase was developed, and the “Italian protocol” is one of the mainly used: it uses glycerol trinitrate to demonstrate a positive response for syncope more quickly and to shorten not only the duration of the test, but it also increases its sensitivity with a slight specificity reduction\(^12\).

Subsequently, different protocols modified some factors such as duration of the phases, both the passive phase and the pharmacological challenge, and they also used different drugs in the latter looking to increase the sensitivity and specificity of the test\(^13\).

**Head-up-tilt-test methodology**

Head-up-tilt-test should be carried out in a comfortable, relaxed environment and away from noise to avoid false positives and negatives. Before starting the procedure, it is important to know if the patient is allergic to the drugs used in the pharmacological challenge if necessary. The patient must be under continuous monitoring through an electrocardiogram before starting the test and cardioinhibitory response being evaluated, since some may experience syncope due to the puncture. On the other hand, a plethysmograph placed on a finger or cuff-type on a limb documents the vasodepressor component.

The head-up-tilt-test has two phases: in the first one, called passive (unprovoked), it is recommended for the table to be tilted at an angle between 60° and 70°, because higher or lower angles have been shown reduce the sensitivity and specificity of the test\(^3,14\). This phase lasts from 20 to 40 min, which is time enough to cause

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Figure 1. In the context of non-traumatic loss of consciousness, the disorder is classified in four groups: syncope, epileptic seizures, transient psychogenic loss of consciousness and diverse origins (rare causes). Non-traumatic loss of consciousness can cause falls with subsequent trauma, in which case loss of consciousness is traumatic and non-traumatic. (With permission of 2018 ESC Syncope Consensus).
orthostatic stress. The passive phase is still the first-line in the tilt test. When it remains negative and concluding a diagnosis is still required, the procedure is continued with a second phase with pharmacological challenge (provoked phase) for 10-20 min, with various nitrates or isoproterenol being used in the vast majority of protocols.

At the National Institute of Cardiology (Mexico), the tilt test is performed at an angle of 70° in the passive phase for 20 min, followed by a provocative phase when a diagnosis is not yet established, and 5 mg isosorbide dinitrate is used for 12 min, which allows enough time to trigger the syncope.

The test is considered positive when the patient reproduces symptoms of syncope or hypotensive or cardioinhibitory response or both are documented, with return of the patient to a Trendelenburg position within 10-15 s and spontaneous recovery. At conclusion, it is classified according to the predominant hemodynamic response, either vasodepressor, cardioinhibitory, or mixed (Table 2); in addition, the test allows diagnosing other causes of syncope, such as orthostatic hypotension (OH) or postural orthostatic tachycardia syndrome (POTS). The classification confers clinical meaning to the tilt test (Table 3), with a variable positivity rate (Fig. 2).

### Table 2. Types of hemodynamic response on the tilt table

<table>
<thead>
<tr>
<th>Type of response on the tilt table</th>
<th>Response characteristics</th>
</tr>
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<tbody>
<tr>
<td>Type 1 or mixed</td>
<td>Heart rate drops at the moment the syncope occurs, but not &lt; 40 bpm; drop lasts &lt; 10 s. BP drops prior than HR</td>
</tr>
<tr>
<td>Type 2A (cardioinhibitory without asystole)</td>
<td>Heart rate drops &lt; 40 bpm, for more than 10 s. BP drops before HR</td>
</tr>
<tr>
<td>Type 2B (cardioinhibitory with asystole)</td>
<td>Asystole with &gt; 3-s duration. Drop in blood pressure coincides with or is subsequent to that of HR</td>
</tr>
<tr>
<td>Type 3 or vasomotor</td>
<td>HR drop does not exceed 10% in comparison with the peak at the moment of syncope</td>
</tr>
<tr>
<td>Chronotropic incompetence</td>
<td>No significant increase in HR during tilt (e.g., &lt; 10% of HR before tilt)</td>
</tr>
<tr>
<td>Postural orthostatic tachycardia syndrome (POTS)</td>
<td>Exercise increases HR (&gt; 130 bpm), at the beginning and during the tilt, before the syncope</td>
</tr>
</tbody>
</table>

HR: heart rate; bpm: beats per minute; BP: blood pressure.

### Table 3. Tilt test clinical Interpretation (warnings regarding interpretation of the result on the tilt table)

<table>
<thead>
<tr>
<th>Clinical meaning according to the result on the tilt table</th>
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<tbody>
<tr>
<td>– A negative head-up-tilt-test does not exclude reflex syncope diagnosis</td>
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<tr>
<td>– Despite tilt test sensitivity and specificity with regard to the presence of susceptibility to hypotension, which may not only be present in reflex syncope but also in other causes of syncope, including those of cardiac origin. The concept of susceptibility to hypotension, rather than being a diagnosis, has important clinical usefulness, given that its absence or presence play an important role for directing treatment toward pacemaker in patients affected with reflex syncope and in the control of hypotension, which is more common in elderly patients</td>
</tr>
<tr>
<td>– A positive cardioinhibitory test on the tilt table predicts with high probability the presence of syncope with spontaneous asystole; this is an important finding due to the implication of considering pacemaker in the treatment of these cases. The presence of a vasopressor response or mixed response does not exclude that the patient suffers from asystole during spontaneous syncope periods</td>
</tr>
<tr>
<td>– The tilt test is useful to separate the syncope that occurs with seizures in patients who experience abnormal movements due to epilepsy</td>
</tr>
<tr>
<td>– The tilt test has value to distinguish the syncope of patients who only suffer falls</td>
</tr>
<tr>
<td>– The tilt test is useful to separate the syncope from the psychogenic pseudosympnope. In patients with suspected psychogenic pseudosyncope, the tilt test is performed in conjunction with an EEG to monitor and confirm the diagnosis</td>
</tr>
<tr>
<td>– The head-up-tilt-test should not be used to assess treatment efficacy</td>
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</table>

EEG: electroencephalogram.

Although there is still no reference standard for the diagnosis of syncope, head-up-tilt-test remains the most widely used method for diagnostic purposes.\(^3\,15\) Nevertheless, given that, usually, the documented standards of the test come from studies on adult subjects, there is no specific methodology for pediatric patients. It has been documented that, in the latter, it is enough with 10-min test periods at an angulation of the table from 60° to 70° to precipitate the syncope, and a specificity of 85% has been demonstrated.\(^6\)

### Indications

1. To confirm the syncope diagnosis in patients in whom it has not been confirmed or to reproduce the symptoms referred by the subject, with hemodynamic pattern determined after initial evaluation\(^3\).
2. To assess the autonomic failure causative of OH, either due to suspicion or confirmation of changes in blood pressure during initial evaluation\(^3\).

3. To diagnostically confirm the presence of POTS\(^3\).

4. To differentiate in case of clinical suspicion of psychogenic pseudosyncope with regard to another cause of syncopal origin\(^3\).

**Contraindications**

Contraindications to the test include severe mitral stenosis, diseases that cause obstruction of the left ventricular outflow tract, arterial cerebral disease, and severe coronary artery disease\(^17\).

**Pharmacological challenge on the tilt table in pediatric patients**

One of the fears when performing the head-up-tilt-test in pediatric patients has been the false positive results obtained in some cases, due to patient lack of cooperation, fear of the test, separation from the parents on admission to the test, which arouses anxiety and crying, as well as doctor’s fear of drug-related secondary reactions and asystole periods, which are more alarming the longer they are. Although asystole duration is not related to disease severity and most subjects spontaneously recover the state of alertness after lowering the bed to the supine position, secondary episodes of cardiorespiratory arrest have been documented\(^18,19\).

Pharmacological challenge has been highly useful and has increased the sensitivity and specificity of the test. However, controversy continues regarding which drug is most useful for this age group due to the wide variety of results and the paucity of specific studies; isoproterenol and nitrates are the most widely used drugs.

**Isoproterenol**

Most adverse effects on the tilt test with pharmacological challenge have been attributed to the use of isoproterenol\(^13,20\). Cases of acute myocardial infarction and ventricular arrhythmias have been reported during isoproterenol administration in the pharmacological challenge; however, these were recorded in patients > 60 years\(^21\). Another relevant point regarding isoproterenol administration route is that intravenous instrumentation can trigger syncope and confound the results\(^22\).

Lai et al. carried out a study in 79 pediatric patients (age range, 5.5-18 years) divided in two groups: the first one, with VVS clinical diagnosis (65 patients), and the second, without clinical data consistent with VVS (14 patients). Both groups underwent head-up-tilt-test, which consisted of a first phase without pharmacological challenge, followed by a challenge with isoproterenol infusion in patients who during the first phase did not trigger syncope. Up to 29 patients in Group 1 and two in Group 2 required pharmacological challenge to determine the syncope origin and type of response. Group 2 showed various medically adverse effects: two patients suffered ventricular arrhythmias, one required pacemaker secondary to a second degree blockade and the rest experienced ventricular extrasystoles. It was concluded that isoproterenol infusion increased the sensitivity of the test by up to 45%, and specificity decreased slightly from 93% to 86%\(^23\). However, drug infusion duration (30 min) and dose used in this protocol (0.5-1 µg/min) might have caused an increase in the number of false positives during the test.

A shorter infusion duration and an isoproterenol dose between 0.5 and 5 µg/min have been reported to be enough to precipitate symptoms, which prevents a larger number of erroneous results\(^24,25\).
**Nitrates**

The first studies by Dindar et al. showed an increase in the sensitivity of the test with the use of Integrated Services Digital Network (ISDN) in comparison with the passive phase alone, with an increase in sensitivity to 77.5% with regard to 15%, and a slight change in specificity, 91.6% versus 100%, respectively. Subsequently, Karacan et al. demonstrated similar results in a study carried out in 29 patients where ISDN was compared with the conventional test without pharmacological challenge, an increase in sensitivity was recorded (96.7% vs. 30%), with a slight reduction in sensitivity (93.3% vs. 100%, respectively), without serious side effects being demonstrated, and thus it can be indicated as an effective and safe method.

**Comparative studies**

Sensitivity (50-80%) with the use of nitroglycerin is similar to that demonstrated with isoproterenol administration (60-85%) in various studies. However, despite the specificity reduction in the pharmacological challenge, specificity is 85-95% with nitroglycerin in comparison with isoproterenol, with a specificity reduction from 35% to 83%

Even though Swissa et al., in a comparison of 136 patients undergoing pharmacological challenge in the tilt test with isoproterenol, relative to ISDN, demonstrated that the use of the latter caused a larger number of bradycardiac reflexes and longer asystolic response duration, only one case of cardiorespiratory arrest with recovery after rescue maneuvers was reported. Various comparative studies have shown a larger number of positive responses to the test in up to 12% with nitroglycerin, as well as better tolerance and fewer adverse effects in comparison with the use of isoproterenol.

In another study carried out at the National Institute of Cardiology in Mexico City, which is described below, the safe and effective use of said test (ISDN was used) was demonstrated in a larger patient cohort. Similarly, nitroglycerin sublingual use in the provocation phase of the test, especially in pediatric patients, prevents intravenous administration from increasing both patient psychological stress and false-positive results.

**Study conducted in patients of the National Institute of Cardiology**

A retrospective analysis of head-up-tilt-test results was carried out at Ignacio Chávez National Institute of Cardiology in patients aged ≤ 18 years with a clinical history indicative of VVS or syncope of unknown origin, between 2015 and 2018. There were 220 patients included, who were divided in two age groups: Group 1, < 10 years (36 patients), and Group 2, from 10 to 18 years of age (183 patients). No predominant gender was observed in either group. In 179 patients (83%) of the total sample, pharmacological challenge was necessary in the tilt table test to determine their hemodynamic response. The positive response rate of the entire sample was 66% (Group 1, 50%; Group 2, 70%). In both groups, the hemodynamic responses pattern that predominated was the mixed type (Group 1, 50%, Group 2, 56%).

**Group I**

Thirty-six patients, with an age range of 6-10 years (mean of 7.8). The gender ratio was 1.25:1 (20/16) in favor of the male gender. Main indication for the test was syncpe in 23 patients, followed by pre-syncpe in 13 patients. The test was positive in 18 patients, three showed positive response during phase 1 (without pharmacological challenge) and the rest required ISDN administration. Response time ranged from 4 to 12 min, with an average of 9.3. Predominant hemodynamic response was of the mixed type in 9 patients (50%), followed by dysautonomia (4) and cardioinhibitory 2B response (4) with pauses ranging between 3 and 6 s; only one patient had a cardioinhibitory 2A response.

**Group II**

One hundred and eighty-three patients aged between 11 and 18 years (mean, 14.8). No gender difference was observed (1:1 ratio). The test was positive in 128 patients (70%), 31 during phase 1 and the rest required ISDN sublingual administration. Response time during the test was variable, from 1 to 20 min, with a mean of 8.5 min. The predominant hemodynamic response was of the mixed type in 43 patients (33.7%), followed by dysautonomia in 32 patients (25%), cardioinhibitory 2B in 25 patients (19.5%), vasodepressor in 14 patients (11%), cardioinhibitory 2A in 13 patients (10%), and only one patient experienced POTS (0.8%).

In all asystole cases during the test, recovery was achieved after positioning the patient in the supine position and administering fluid therapy. No complications were recorded during the study. This test demonstrates...
like others already mentioned, the safe use of sublingual ISDN in pediatric patients.

**Treatment**

**NON-PHARMACOLOGICAL MEASURES**

In children with VVS, preventive measurements are still the first-choice treatment and include reassuring the patient (he/she is indicated that the procedure is benign and that it has the purpose to reduce psychological stress), avoiding syncope-triggering factors and increasing salt ingestion (2 g/day); fluids (3 L daily) reduce such episodes. Training through physical maneuvers that increase venous return and orthostatic tolerance have been shown to be effective to revert syncope. Various clinical studies have demonstrated the effectiveness of non-pharmacological treatment and have reduced syncope recurrence from 56% in untreated patients to 39% when fluid and salt consumption was increased ($p = 0.029$). It is essential to emphasize the importance of discipline in dietary changes (water and salt ingestion) and physical training to prevent syncope in pediatric patients. Recently, a cross-sectional study was carried out in 70 patients aged from 5 to 20 years, divided into two groups: Group 1 was made up of 30 patients (13 males/17 females) who received pharmacological and non-pharmacological measures and was compared with Group 2, which included 40 patients (18 males/22 females) treated only with non-pharmacological measures (increase of dietary water and salt, physical maneuvers). Both groups were followed up for 3 years, and a significant difference was observed in pre-syncope and syncope symptoms in those patients only non-pharmacologically treated (Group 2), with a 3-year recurrence rate of 5% in comparison with patients who combine the same treatment with pharmacological measures (Group 1), in whom syncope recurrence was 44% ($p = 0.001$); it should be mentioned that the obtained results may be due to the fact that patients undergoing pharmacological and non-pharmacological treatments tend to reduce follow-up and correct use of the latter; it is thus concluded that correctly-used preventive measures is enough for the treatment of patients with neuro-cardiogenic syncope.

**Pharmacological measures**

The use of drugs for the treatment of VVS should be considered as second line in pediatric patients in whom syncope recurrence continues despite preventive measures correct use. The most widely used medications include beta-blockers, midodrine ($\alpha$ receptor agonist), fluoro-hydrocortisone, and serotonin reuptake inhibitors; however, the ideal drug has not yet been found due to the low response to treatment. Despite this, although there are only few studies of midodrine treatment in pediatric patients, they have shown high effectiveness in syncope reduction. A randomized study was carried out in 26 patients with ages ranging between 6 and 16 years, who were divided into two groups: Group 1, with 13 patients who were treated with midodrine and non-pharmacological measures, and Group 2 (13 patients), which was treated only with non-pharmacological measures, for a short follow-up of 6 months. A reduction in conservative treatment-resistant VVS recurrence was demonstrated, with a recurrence rate of 80% in Group 1, in comparison with 22% in patients treated with midodrine ($p = 0.023$). No supine hypertension was observed in any of the subjects treated with midodrine and only one experienced gastrointestinal discomfort during treatment. In a second study, conducted in 48 patients aged 6-17 years with syncope or pre-syncope symptoms, the participants were randomly assigned to three groups to compare different treatments. Group 1 included individual treated with non-pharmacological measures; Group 2, subjects with cresol-based treatment (placebo); and Group 3, patients treated with midodrine, each group made up of 16 patients, during an approximate follow-up of 9 months. It was concluded that midodrine was effective in reducing syncope when compared to the other treatments used ($p = 0.05$). Despite the results of both studies, the follow-up short duration, as well as treatment response assessment by repeating the head-up-tilt-test, limit the obtained results, since the guidelines indicate the modest usefulness of the latter for therapeutic evaluation. Another possibility for the demonstrated effectiveness might be secondary to the type of predominant hemodynamic response (vasodepressor), which is correlated with a higher effectiveness rate when using an $\alpha$ receptor agonist drug.

Evidence of fluoro-hydrocortisone benefit is limited because there are only few studies conducted in this age group. A randomized, double-blind, and placebo-controlled study was conducted in 32 patients (one was lost to follow-up during the study), all aged < 18 years. Group 1 included 18 patients treated with salt supplements and fluoro-hydrocortisone, and Group 2 (14 patients) received placebo, with an approximate follow-up of 1 year. Conversely to the hypothesis raised...
in favor of the use of fluoroxyhydrocortisone, subjects in the placebo group were found to have lower syncope recurrence (p = 0.04), and only 53\% of total treated patients did not suffer syncope during follow-up; therefore, low effectiveness of the employed drug was observed\(^4\). These results are similar to those of a comparative study between fluoroxyhydrocortisone (0.2 mg/day) and atenolol (100 mg/day) for the treatment of VVS, where the recurrence rate was 52\% (30 patients) of total subjects treated in both groups (a total of 58 individuals)\(^4\).

Beta-blockers remain a treatment with a Class III recommendation (no benefits) for pediatric patients, according to syncope treatment guidelines\(^3\,^6\), despite the fact that a reduction of 60\% in recurrent syncope was initially demonstrated in 21 patients treated with metoprolol\(^5\). Subsequently, in a prospective, randomized study, carried out in patients aged 8-17 years, with a 1-year follow-up, treatment with metoprolol was shown to be related to a higher syncope recurrence rate in comparison with non-pharmacological treatment, 43\% versus 29\%, respectively (p = 0.389)\(^16\). However, a study was recently performed in 38 pediatric patients with VVS, aged 6 to 13 years, in whom 24-h urine norepinephrine (NE) values analysis was carried out to demonstrate that urine NE high figures are related to a higher response to metoprolol treatment. Said patients were initially compared with a control group of 20 healthy subjects. After 24-h urine NE values were measured in patients with VVS (31.62 ± 14.11 µg/24 h), a significantly high dispersion coefficient was recorded (R\(^2\) = 0.0028) in comparison with the control group, in which 24-h urine NE figures were 35.04 ± 7.28 µg/24 h, with a lower dispersion coefficient (R\(^2\) = 0.0002). These results demonstrate individual differences between patients with VVS, in whom urine NE quantification has a possible predictive value for metropolol treatment evaluation. Due to the obtained results, the use of metropolol was evaluated in the 38 patients with the condition, and it was concluded that patients who show an effective response to metropolol are linked to 24-h urine NE high levels in comparison with those without treatment response (40.75 ± 12.86 vs. 21.48 ± 6.49, respectively). Similarly, it was established that patients with 24-h urine NE elevated values have a higher supine blood pressure elevation, both systolic and diastolic and that 24-h urine NE values > 34.84 µg/24 h are indicators that metropolol treatment can be effective in pediatric patients, with 100\% specificity and 70\% sensitivity\(^3\). This opens a door for further studies based on the principles presented in this study.

**Interventional treatments**

The use of pacemaker in pediatric patients remains a controversial topic due to the benign nature of the disease, and even in the guidelines there is still controversy regarding this issue. The AHA/ACC/HRS guidelines (2017) for syncope assessment and treatment classify the use of pacemaker with level of evidence IIb, based on two studies conducted in 22 pediatric patients who experienced periods of apnea that sometimes caused seizures secondary to cerebral anoxia, and in whom prolonged periods of asystole (> 4 s) were subsequently documented; after placement of the pacemaker, syncope episodes were reduced by up to 86\%\(^5\).

There is still insufficient evidence to consider pacemaker as an absolute indication when the patient meets said clinical criteria, and the type of pacemaker to be used has neither yet been studied because both the one-chamber pacemaker with hysteresis and the dual-chamber pacemaker demonstrate similar effectiveness in syncope reduction\(^4\,^8\,^9\). However, studies comparing both pacemakers are still required, as well as long-term follow-up of these patients.

On the other hand, the European Society of Cardiology (ESC 2018) guidelines do not recommend the use of pacemakers in pediatric patients and neither in young adults; subjects aged > 40 years are the ideal patients when they meet the criteria for the procedure\(^3\).

Cardioneuroablation has been proposed as a novel treatment for patients who exhibit a cardioinhibitory-type response. This treatment involves ablation of intra-cardiac parasympathetic ganglionated plexuses, predominantly localized on the posterior wall of the left atrium. The objective is to reduce vagal tone and thus avoid the cardioinhibitory component by increasing heart rate to reduce syncope episodes. Although there are no cohort studies in pediatric patients, some isolated cases such as that reported by Suenega et al.\(^5\) in a 17-year-old female patient, and the one by Debruyne et al.\(^5\) in a 16-year-old individual, show promising results, with patients being free of syncope during a 12- and a 22-month follow-up, respectively, after the intervention. Despite the good results, studies with larger cohorts are required to consider cardioneuroablation as a therapeutic option for pediatric patients.

**Conclusions**

Neurocardiogenic syncope constitutes a diagnostic challenge in pediatrics. On initial evaluation, it is necessary ruling out personal and family-inherited heart
pathology history in search for sudden death and, in case of suspicion or doubt, referring the patient to the pediatric cardiologist or electrophysiologist is necessary once neurological and psychogenic problems have been ruled out as the primary origin of syncope.

The head-up–tilt-test helps to clarify diagnostic suspicion of neuro-cardiogenic syncope, which is the most common cause of syncope. In pediatric patients, there is still controversy regarding the use of pharmacological challenge during head-up-tilt-test, but various studies have documented its safe use, even though cases of cardiorespiratory arrest have been documented in tilt tests, as well as other adverse effects attributed to use of isoprotenerol during pharmacological challenge, although these are mostly studies carried out in adults in whom comorbidities play an important role. The use of nitrates has been shown to be effective and safe in pediatric patients, and sublingual administration facilitates their use and benefits the test, which prevents false-positive results due to stress secondary to the puncture for medication administration. Doctor’s fear of using this compound limits the test results, since pharmacological challenge increases sensitivity with a slight specificity reduction.

First-choice treatment is non-pharmacological measures, which involves educating the patient and changing his/her diet (increase in water and salt intake). Pharmacological treatment has been shown to be effective in few studies, specifically in the case of midodrine. Although some recent studies demonstrate that metoprolol can be effective if its use is based on 24-h urine NE quantification, this study opens a field in pharmacological treatment research.

The use of pacemaker to treat neuro-cardiogenic syncope in pediatric patients remains controversial even in syncope guidelines and the paucity of evidence limits its use given that there are no clearly-established implementation criteria. Cardioneuroablation is a promising treatment in patients that show cardioinhibitory response during syncope due to the success reported in isolated cases, and it has been shown to be effective, safe and without syncope recurrence during follow-up. However, further studies are required in this regard aimed at the pediatric population.

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Conflicts of interest

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Ethical disclosures

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