PART 1. THE INITIAL EVALUATION OF PATIENTS WITH SYNCOPE


The initial evaluation

The ‘Initial evaluation’ of a patient presenting with syncope consists of: careful history, physical examination including orthostatic blood pressure measurements and standard electrocardiogram (ECG).

Three key questions should be addressed during the initial evaluation:

- Is loss of consciousness attributable to syncope or not?
- Are there important clinical features in the history that suggest the diagnosis?
- Is heart disease present or absent?

Question 1. Syncope or non-syncopal condition?

Differentiating true syncope from other ‘non-syncopal’ conditions associated with real or apparent transient loss of consciousness is generally the first diagnostic challenge and influences the subsequent diagnostic strategy (Fig. 1).

Definition of syncope

Syncope is a symptom, defined as a transient, self-limited loss of consciousness, usually leading to falling. The onset of syncope is relatively rapid, and the subsequent recovery is spontaneous, complete, and usually prompt. The underlying mechanism is a transient global cerebral hypoperfusion.

In some forms of syncope there may be a premonitory warning of an impending syncopal event, in another loss of consciousness occurs without warning. Recovery from syncope is usually accompanied by almost immediate restoration of appropriate behaviour and orientation. Retrograde amnesia, although believed to be uncommon, may be more frequent than previously thought, particularly in older individuals. Sometimes the post-recovery period may be marked by fatigue. Typical syncope episodes are brief and usually last no longer than 20 s. Rarely, syncope duration may be longer, even lasting for several minutes. In such cases, the differential diagnosis between syncope and other causes of loss of consciousness can be difficult.

Presyncope or ‘near-syncope’ refers to a condition in which patients feel as though syncope is imminent. The classification of syncope is shown in Table 1.

Non-syncopal conditions

Several disorders resemble syncope in two different ways. In some, consciousness is impaired or lost because of metabolic disorders (including hypoxia, hyperventilation, hypoglycaemia), epilepsy and intoxications. In several other disorders, consciousness is only apparently lost; this is the case in somatization disorders, cataplexy and drop attacks (Table 2).

- Epilepsy can (depending on the type) cause loss of consciousness. When witnessed by an expert, tonic-clonic attacks are easy to recognize. Questions to distinguish between tonic-clonic seizures and syncope should be directed separately at patients and eye witnesses (Table 3). The patient should be asked whether there were any promonitory signs. The typical textbook aura, consisting of a rising sensation in the abdomen, and/or an unusual unpleasant smell, is relatively rare. Aura patterns are usually repetitive over time in patients, who will therefore learn to recognize them as such. The patient should be asked how he/she felt on regaining consciousness: confusion or sleepiness lasting more than a few minutes point to epilepsy, as do tongue biting, or muscle pains lasting for hours or days. Urinary incontinence is not useful in the distinction. Witnesses should be asked to describe any movements. Unconsciousness without any movement makes epilepsy unlikely, but movements certainly do not exclude syncope (also improperly called ‘convulsive syncope’), although the presence of any movement is often interpreted by both medical personnel and laymen as indicative of epilepsy. Syncopal movements are typically asynchronous and limited in scope (called ‘myoclonic’ in neurology), while a tonic posture involves forceful extension of the extremities, and clonic movements (not called myoclonus) involve massive synchronous jerks of the arms.
Real or apparent transient loss of consciousness

**Syncope:**
- Neurally-mediated reflex syncopal syndromes
- Orthostatic
- Cardiac arrhythmias as primary cause
- Structural cardiac or cardiopulmonary disease
- Cerebrovascular

**Non-syncopal:**
- Disorders resembling syncope with impairment or loss of consciousness, e.g. seizure disorders, etc
- Disorders resembling syncope without loss of consciousness, e.g. psychogenic "syncope" (somatization disorders), etc

*Figure 1  Classification of transient loss of consciousness.*

**Table 1  Causes of syncope**

<table>
<thead>
<tr>
<th>Neurally-mediated reflex syncopal syndromes</th>
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<tbody>
<tr>
<td>Vasovagal faint (common faint)</td>
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<tr>
<td>Carotid sinus syncope</td>
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<tr>
<td>Situational faint</td>
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<tr>
<td>— acute haemorrhage</td>
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<tr>
<td>— cough, sneeze</td>
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<tr>
<td>— gastrointestinal stimulation (swallow, defaecation, visceral pain)</td>
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<tr>
<td>— micturition (post-micturition)</td>
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<td>— post-exercise</td>
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<tr>
<td>— others (e.g. brass instrument playing, weightlifting, post-prandial)</td>
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<tr>
<td>Glossopharyngeal and trigeminal neuralgia</td>
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<tr>
<th>Orthostatic</th>
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<tr>
<td>Autonomic failure</td>
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<tr>
<td>— Primary autonomic failure syndromes (e.g. pure autonomic failure, multiple system atrophy, Parkinson's disease with autonomic failure)</td>
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<tr>
<td>— Secondary autonomic failure syndromes (e.g. diabetic neuropathy, amyloid neuropathy)</td>
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<tr>
<td>— Drugs and alcohol</td>
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<tr>
<td>Volume depletion</td>
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<tr>
<td>— Haemorrhage, diarrhoea, Addison's disease</td>
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<tr>
<th>Cardiac arrhythmias as primary cause</th>
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<tr>
<td>Sinus node dysfunction (including bradycardia/tachycardia syndrome)</td>
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<tr>
<td>Atrioventricular conduction system disease</td>
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<tr>
<td>Paroxysmal supraventricular and ventricular tachycardias</td>
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<tr>
<td>Inherited syndromes (e.g. long QT syndrome, Brugada syndrome)</td>
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<tr>
<td>Implanted device (pacemaker, ICD) malfunction</td>
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<tr>
<td>Drug-induced proarrhythmias</td>
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<tr>
<th>Structural cardiac or cardiopulmonary disease</th>
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<tr>
<td>Cardiac valvular disease</td>
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<tr>
<td>Acute myocardial infarction/ischaemia</td>
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<tr>
<td>Obstructive cardiomyopathy</td>
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<tr>
<td>Atrial myxoma</td>
</tr>
<tr>
<td>Acute aortic dissection</td>
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<tr>
<td>Pericardial disease/tamponade</td>
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<td>Pulmonary embolus/pulmonary hypertension</td>
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<table>
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<tr>
<th>Cerebrovascular</th>
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<tr>
<td>Vascular steal syndromes</td>
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and/or legs. Mimicking the movements helps to make witnesses choose between the options. In syncope, movements occur as a result of brain ischaemia, and therefore occur after the patient has slumped to the floor. In epilepsy, clonic movements can occur before the fall, whereas the tonic posture can cause the patient to keel over like a falling log. In other forms of epilepsy, such as absence epilepsy in children and partial complex epilepsy in adults, consciousness is not so much lost as altered, and this does not lead to falls.

There are many neurological reasons to fall, but only rarely are such falls accompanied by loss of consciousness. Cataplexy is an example: a partial or complete loss of muscular control occurs triggered by emotions, usually laughter. Even when the patient appears to be wholly unconscious, there is a later full recollection of all events. Cataplexy most often occurs as part of narcolepsy; in fact, the combination of cataplexy with daytime sleepiness ensures the diagnosis of narcolepsy.

‘Drop attack’ is an unclear entity. Definitions vary from very expansive ones, encompassing syncope and most other causes to more restrictive ones. The clearest use of the term concerns ‘cryptogenic drop attacks’ describing women (very rarely men) who suddenly...
drop on their knees without any apparent reason, after which patients can get up immediately; because of this the disorder is also known as 'maladie des genoux bleus'. There is no loss of consciousness, or this is so short that it cannot be ascertained with certainty by patient or doctor. There are no associated signs or symptoms of any kind. The disorder can exist unaltered for many years. If used in this strict sense, the term has a specific meaning. If it is used in the wide sense, it obscures rather than elucidates a variety of diverse disorders, and hampers understanding.

Syncope-like symptoms may be due to anxiety, hysteria, panic attacks and major depression. Despite the presence of psychiatric disorders, a careful search for other causes of syncope is needed since the attribution of psychiatric disorder to syncope is often difficult. Patients with syncope associated with psychiatric illnesses are young, with low prevalence of heart disease who have frequent recurrent syncope. Patients with conversion reactions may faint in the presence of witnesses and may not have injury. Syncope may be mimicked by somatization disorder.

Question 2. Are there important clinical features in the history that suggest the diagnosis?

Accurate history taking alone may be diagnostic of the cause of syncope or may suggest the strategy of evaluation (Table 4).

Question 3. Is heart disease present or absent?

Heart disease is considered present if it can be suspected or diagnosed on the basis of history, physical examination and/or electrocardiographic abnormalities. The absence of suspected or certain heart disease excludes a cardiac cause of syncope with the exception of syncope accompanied by palpitations which could be due to paroxysmal tachycardia. Conversely, the presence of heart disease at the initial evaluation is a strong predictor of cardiac cause of syncope and virtually includes all cardiac syncopes, but its specificity is low as about half of patients with heart disease have a non-cardiac cause of syncope.

Table 4  Important historical features

<table>
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<tr>
<th>Questions about circumstances just prior to attack</th>
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<tbody>
<tr>
<td>- Position (supine, sitting or standing)</td>
</tr>
<tr>
<td>- Activity (rest, change in posture, during or after exercise, during or immediately after urination, defaecation, cough or swallowing);</td>
</tr>
<tr>
<td>- Predisposing factors (e.g. crowded or warm places, prolonged standing, post-prandial period) and of precipitating events (e.g. fear, intense pain, neck movements);</td>
</tr>
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</table>

Questions about onset of attack:
- Nausea, vomiting, abdominal discomfort, feeling of cold, sweating, aura, pain in neck or shoulders, blurred vision;

Questions about attack (eyewitness):
- Way of falling (slumping or keeling over), skin colour (pallor, cyanosis), duration of loss of consciousness, breathing pattern (snoring), movements (tonic, clonic, tonic-clonic or minimal myoclonus, automatism) and their duration, onset of movement in relation to fall, tongue biting;

Questions about end of attack:
- Nausea, vomiting, sweating, feeling of cold, confusion, muscle aches, skin colour, injury, chest pain, palpitations; urinary or faecal incontinence

Questions about antecedents:
- Family history of sudden death, congenital arrhythmogenic heart disease or fainting
- Previous cardiac disease
- Neurological history (Parkinsonism, epilepsy, narcolepsy)
- Metabolic disorders (diabetes, etc.)
- Medication (antihypertensive, antianginal, antidepressant agent, antiarrhythmic, diuretics and QT prolonging agents)
- (In case of recurrent syncope.) Information on recurrences such as the time from the first syncopal episode and on the number of spells
Specificity increases in some situations. These are:

- In the presence of heart disease, some historical variables such as blurred vision, syncope occurring in the supine position or during effort, or convulsive syncope suggest a cardiac cause of syncope with high specificity. These patients should probably be hospitalized and cardiological examinations should be given priority.
- Even in the presence of heart disease, a long duration (>4 years) of symptoms, syncope preceded by abdominal discomfort or followed by nausea and sweating suggest a neurally mediated cause of syncope with high specificity. In these cases autonomic tests should be given priority and, in case of their positivity, a neurally-mediated mechanism is likely and the diagnostic work-up should be stopped.

When heart disease is absent, a cardiac cause is unlikely unless syncope is preceded by palpitations. In the latter case, the possibility that a tachyarrhythmia is the cause of syncope should be evaluated. In the other cases, autonomic tests should be performed in order to evaluate the possible neurally mediated nature of syncope. Patients do not usually need hospitalization given the benign outcome of this type of syncope.

**Recommendations**

When the mechanism of syncope is not evident, the presence of suspected or certain heart disease is associated with a higher risk of arrhythmias and a higher mortality at one year. In these patients, cardiac evaluation (consisting of echocardiography, stress testing and tests for arrhythmia detection such as prolonged electrocardiographic and loop monitoring or electrophysiological study) is recommended. If cardiac evaluation does not show evidence of arrhythmia as a cause of syncope, evaluation for neurally mediated syndromes is recommended in those with recurrent or severe syncope.

In patients without suspected or certain heart disease, evaluation for neurally mediated syncope is recommended for those with recurrent or severe syncope. The tests for neurally mediated syncope consist of tilt testing and carotid sinus massage. The majority of patients with single or rare episodes in this category probably have neurally mediated syncope. An additional consideration is psychiatric illness. Psychiatric assessment is recommended in patients with frequent recurrent syncope who have multiple other somatic complaints and initial evaluation raises concerns of stress, anxiety and possible other psychiatric disorders.

**Indications**

Class I (evidence for and/or general agreement that the procedure or treatment is useful):

- Basic laboratory tests are only indicated if syncope may be due to loss of circulating volume, or if a syncope-like disorder with a metabolic cause is suspected.
- In patients with suspected heart disease, echocardiography, prolonged electrocardiographic monitoring and, if non-diagnostic, electrophysiological studies are recommended as first evaluation steps.
- In patients with palpitations associated with syncope, electrocardiographic monitoring and echocardiography are recommended as first evaluation steps.
- In patients with chest pain suggestive of ischaemia before or after loss of consciousness, stress testing, echocardiography, and electrocardiographic monitoring are recommended as first evaluation steps.
- In young patients without suspicion of heart or neurological disease and recurrent syncope, tilt testing and, in older patients, carotid sinus massage are recommended as first evaluation steps.
- In patients with syncope occurring during neck turning, carotid sinus massage is recommended at the outset.
- In patients with syncope during or after effort, echocardiography and stress testing are recommended as first evaluation steps.
- In patients with signs of autonomic failure or neurological disease a specific diagnosis should be made.

**The diagnostic value of the initial evaluation**

The initial evaluation may lead to certain or suspected diagnosis or no diagnosis (here termed as unexplained syncope).

**Certain diagnosis**

Initial evaluation may lead to a certain diagnosis based on symptoms, signs or ECG findings. Under such circumstances, no further evaluation of the disease or disorder may be needed and treatment, if any, can be planned. This is the case in the following recommendations:

**Recommendations**

**Diagnosis**

Class I (evidence for and/or general agreement that the procedure or treatment is useful): The results of the initial measures are diagnostic of the cause of syncope in the following situations:

- *Vasovagal syncope* is diagnosed if precipitating events such as fear, severe pain, emotional distress, instrumentation or prolonged standing are associated with typical prodromal symptoms.
- *Situational syncope* is diagnosed if syncope occurs during or immediately after urination, defaecation, cough or swallowing.
Orthostatic syncope is diagnosed when there is documentation of orthostatic hypotension associated with syncope or presyncope. Orthostatic blood pressure measurements are recommended after 5 min of lying supine, followed by each minute, or more often, after standing for 3 min. Measurements may be continued longer, if blood pressure is still falling at 3 min. If the patient does not tolerate standing for this period, the lowest systolic blood pressure during the upright posture should be recorded. A decrease in systolic blood pressure $\geq 20$ mmHg or a decrease of systolic blood pressure to $<90$ mmHg is defined as orthostatic hypotension regardless of whether or not symptoms occur.

Cardiac ischaemia related syncope is diagnosed when symptoms are present with ECG evidence of acute ischaemia with or without myocardial infarction, independently of its mechanism*.

Table 5 Clinical features suggestive of specific causes

<table>
<thead>
<tr>
<th>Symptom or finding</th>
<th>Possible cause</th>
</tr>
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<tbody>
<tr>
<td>After sudden unexpected unpleasant sight, sound, or smell</td>
<td>Vasovagal</td>
</tr>
<tr>
<td>Prolonged standing at attention or crowded, warm places</td>
<td>Vasovagal or autonomic failure</td>
</tr>
<tr>
<td>Nausea, vomiting associated with syncope</td>
<td>Vasovagal</td>
</tr>
<tr>
<td>Within one hour of a meal</td>
<td>Post-prandial (autonomic failure)</td>
</tr>
<tr>
<td>After exertion</td>
<td>Vasovagal or autonomic failure</td>
</tr>
<tr>
<td>Syncope with throat or facial pain</td>
<td>Neuralgia (glossopharyngeal or trigeminal neuralgia)</td>
</tr>
<tr>
<td>With head rotation, pressure on carotid sinus (as in tumours, shaving, tight collars)</td>
<td>Spontaneous carotid sinus syncope</td>
</tr>
<tr>
<td>Within seconds to minutes upon active standing</td>
<td>Orthostatic hypotension</td>
</tr>
<tr>
<td>Temporal relationship with start of medication or changes of dosage</td>
<td>Drug induced</td>
</tr>
<tr>
<td>During exertion, or supine</td>
<td>Cardiac syncope</td>
</tr>
<tr>
<td>Preceded by palpitation</td>
<td>Tachyarrhythmia</td>
</tr>
<tr>
<td>Family history of sudden death</td>
<td>Long QT syndrome</td>
</tr>
<tr>
<td>In course of a migraine attack</td>
<td>Brugada syndrome</td>
</tr>
<tr>
<td>Associated with vertigo, dysarthria, diplopia</td>
<td>Right ventricular dysplasia</td>
</tr>
<tr>
<td>With arm exercise</td>
<td>Hypertrophic cardiomyopathy</td>
</tr>
<tr>
<td>Differences in blood pressure or pulse in the two arms</td>
<td>Migraine</td>
</tr>
<tr>
<td>Confusion after attack for more than 5 min</td>
<td>Subclavian steal</td>
</tr>
<tr>
<td>Tonic-clonic movements, automatism, tongue biting, blue face, epileptic aura</td>
<td>Subclavian steal or aortic dissection</td>
</tr>
<tr>
<td>Frequent attack with somatic complaints, no organic heart disease</td>
<td>Seizure</td>
</tr>
</tbody>
</table>

Table 6 ECG abnormalities suggesting an arrhythmic syncope

<table>
<thead>
<tr>
<th>Abnormality</th>
<th>Suggestive conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bifascicular block (defined as either left bundle branch block or right bundle branch block combined with left anterior or left posterior fascicular block)</td>
<td>Orthostatic hypotension</td>
</tr>
<tr>
<td>Other intraventricular conduction abnormalities (QRS duration $\geq 0.12$ s)</td>
<td>Other intraventricular conduction abnormalities</td>
</tr>
<tr>
<td>Mobitz II second degree atrioventricular block</td>
<td>Long QT syndrome</td>
</tr>
<tr>
<td>Asymptomatic sinus bradycardia ($&lt;50$ bpm) or sinoatrial block</td>
<td>Brugada syndrome</td>
</tr>
<tr>
<td>Pre-excited QRS complexes</td>
<td>Right ventricular dysplasia</td>
</tr>
<tr>
<td>Prolonged QT interval</td>
<td>Hypertrophic cardiomyopathy</td>
</tr>
<tr>
<td>Right bundle branch block pattern with ST-elevation in leads V1–V3 (Brugada syndrome)</td>
<td>Migraine</td>
</tr>
<tr>
<td>Negative T waves in right precordial leads, epsilon waves and ventricular late potentials suggestive of arrhythmogenic right ventricular dysplasia</td>
<td>Subclavian steal or aortic dissection</td>
</tr>
<tr>
<td>Q waves suggesting myocardial infarction</td>
<td>Seizure</td>
</tr>
</tbody>
</table>

Cardiac ischaemia related syncope is diagnosed when symptoms are present with ECG evidence of acute ischaemia with or without myocardial infarction, independently of its mechanism*.

Arrhythmia related syncope is diagnosed by ECG when there is:
- Sinus bradycardia $<40$ beats $\cdot$ min$^{-1}$ or repetitive sinoatrial blocks or sinus pauses $>3$ s in the absence of negatively chronotropic medications;
- Mobitz II 2nd or 3rd degree atrioventricular block;
- Alternating left and right bundle branch block;
- Rapid paroxysmal supraventricular tachycardia or ventricular tachycardia;
- Pacemaker malfunction with cardiac pauses.

Suspected diagnosis

More commonly, the initial evaluation leads to a suspected diagnosis, which needs to be confirmed by directing testing (Tables 5 and 6).

Unexplained diagnosis

The initial evaluation may lead to no diagnosis (here termed as unexplained syncope). The strategy of

*Note. In the case of ischaemic syncope, the mechanism can be cardiac (low output or arrhythmia) or reflex (Bezold–Jarish reflex), but management is primarily that of ischaemia.
evaluation varies according to the severity and frequency of the episodes and the presence or absence of heart disease (Fig. 2).

**Need for hospitalization**

The admission decision can be considered with two different objectives: for diagnosis or for therapy (Table 7). In patients with syncope in whom the aetiology remains unknown after the initial baseline evaluation a risk stratification can be used for hospitalization decision. The presence of underlying structural heart disease and abnormalities of the baseline ECG are important markers for cardiac syncope. An important, but less frequent, prognostic marker is the family history of sudden death (Table 7).
When is it safe not to hospitalize?

Patients with isolated or rare syncopal episodes, in whom there is no evidence of structural heart disease and who have a normal baseline ECG, have a high probability of having a neurocardiogenic syncope and a low risk of having cardiac syncope. These patients have a good prognosis in terms of survival irrespective of the results of head-up tilt test. The evaluation of these patients generally can be completed entirely on an ambulatory basis. Patients with neurally-mediated syncope, in the absence of structural heart disease and normal ECG, have a good prognosis in terms of survival, and generally do not need specific treatment apart from counselling and general measures already defined.
Part 2. Diagnostic tests and treatment: summary of recommendations


The strength of recommendations has been ranked as follows:

- Class I, when there is evidence for and/or general agreement that the procedure or treatment is useful. Class I recommendations are generally those reported in the sections labelled as ‘Recommendations’ and in the tables.
- Class II, when usefulness of the procedure or treatment is less well established or divergence of opinion exists among the members of the Task Force.
- Class III, when the procedure or treatment is not useful and in some cases may be harmful.

The strength of evidence supporting a particular procedure/treatment option has been ranked as follows:

- Level of Evidence A=Data derived from multiple randomized clinical trials or meta-analyses.
- Level of Evidence B=Data derived from a single randomized trial or multiple non-randomized studies.
- Level of Evidence C=Consensus opinion of experts.

When not expressed otherwise, evidence is of type C.

**Diagnostic tests**

*Carotid sinus massage*

Carotid sinus massage is a tool used to disclose carotid sinus syndrome in patients with syncope. Carotid sinus syndrome is diagnosed in patients who are found to have an abnormal response to carotid sinus massage (carotid sinus hypersensitivity) and an otherwise negative work-up for syncope. The relationship between carotid sinus hypersensitivity and spontaneous, otherwise unexplained, syncope is established (level B).

**Diagnosis**

*Class I*

- The procedure is considered positive if symptoms are reproduced during or immediately after the massage in presence of asystole longer than 3 s and/or a fall in systolic blood pressure of 50 mm Hg or more. A positive response is diagnostic of the cause of syncope in the absence of any other competing diagnosis.

**Tilt testing**

*Recommendations*

**Tilt test protocols**

*Class I*

- Supine pre-tilt phase of at least 5 min when no venous cannulation is performed, and at least 20 min when cannulation is undertaken.
- Tilt angle is 60 to 70 degree.
- Passive phase of a minimum of 20 min and a maximum of 45 min.
- Use of either intravenous isoprenaline/isoproterenol or sublingual nitroglycerin for drug provocation if passive phase has been negative. Drug challenge phase duration of 15–20 min.
For isoprenaline, an incremental infusion rate from 1 up to 3 μg/min in order to increase average heart rate by about 20–25% over baseline, administered without returning the patient to the supine position. For nitroglycerin, a fixed dose of 400 μg nitroglycerin spray sublingually administered in the upright position. The end-point of the test is defined as induction of syncope or completion of the planned duration of tilt including drug provocation. The test is considered positive if syncope occurs (Table 1).

Class II
Divergence of opinion exists in the case of induction of pre-syncope.

Indications
Class I
Tilt testing is indicated for diagnostic purposes:
• In case of unexplained single syncopal episode in high risk settings (e.g. occurrence of, or potential risk for, physical injury or with occupational implications), or recurrent episodes in the absence of organic heart disease, or, in the presence of organic heart disease, after cardiac causes of syncope have been excluded;
• When it will be of clinical value to demonstrate susceptibility to neurally-mediated syncope to the patient.

Class II
Tilt testing is indicated for diagnostic purposes:
• When an understanding of the haemodynamic pattern of syncope may alter the therapeutic approach;
• For differentiating syncope with jerking movements from epilepsy;
• For evaluating patients with recurrent unexplained falls;
• For assessing recurrent presyncope or dizziness.

Class III
• Assessment of treatment.

A single episode without injury and not in a high risk setting.
• Clear-cut clinical vasovagal features leading to a diagnosis when demonstration of a neurally mediated susceptibility would not alter treatment.

Diagnosis
Class I
• In patients without structural heart disease, tilt testing can be considered diagnostic, and no further tests need to be performed when spontaneous syncope is reproduced.
• In patients with structural heart disease, arrhythmias or other cardiac causes should be excluded prior to considering positive tilt test results as evidence suggesting neurally mediated syncope.

Class II
• The clinical meaning of abnormal responses other than induction of syncope is unclear.

Electrocardiographic monitoring
(non-invasive and invasive)

Recommendations

Indications

Class I
• Holter monitoring is indicated in patients with structural heart disease and frequent symptoms or even infrequent when there is a high pre-test probability of identifying an arrhythmia responsible for syncope.
• When the mechanism of syncope remains unclear after full evaluation, External or Implantable Loop
Table 2 Minimal suggested electrophysiological protocol for diagnosis of syncope

- Measurement of sinus node recovery time and corrected sinus node recovery time by repeated sequences of atrial pacing for 30–60 s with at least one low (10–20 beats.min⁻¹ greater than sinus rate) and two higher pacing rates*.
- Assessment of the His–Purkinje system includes measurement of the HV interval at baseline and His–Purkinje conduction with stress by incremental atrial pacing. If the baseline study is inconclusive, pharmacological provocation with slow infusion of ajmaline (1 mg/kg i.v.), procainamide (10 mg/kg i.v.), or disopyramide (2 mg/kg i.v.) is added unless contraindicated.
- Assessment of ventricular arrhythmia inducibility performed by ventricular programmed stimulation at two right ventricular sites (apex and outflow tract), at two basic drive cycle lengths (100 or 120 beats.min⁻¹ and 140 or 150 beats.min⁻¹), with up to two extrastimuli**.
- Assessment of supraventricular arrhythmia inducibility by any atrial stimulation protocol.

Comments:
*When sinus node dysfunction is suspected autonomic blockade may be applied, and measurements repeated.
**A third extrastimulus may be added. This may increase sensitivity, but reduces specificity. Ventricular extrastimulus coupling intervals below 200 ms also reduce specificity.

Recorders are recommended when there is a high pre-test probability of identifying an arrhythmia responsible of for syncope.

**Diagnosis**

**Class I**

- ECG monitoring is diagnostic when a correlation between syncope and an electrocardiographic abnormality (brady- or tachyarrhythmia) is detected.
- ECG monitoring excludes an arrhythmic cause when there is a correlation between syncope and sinus rhythm.
- In the absence of such correlations additional testing is recommended with possible exception of:
  - ventricular pauses longer than 3 s when awake;
  - periods of Mobitz II or 3rd degree atrioventricular block when awake;
  - rapid paroxysmal ventricular tachycardia.

**Electrophysiological testing**

The diagnostic efficiency of the invasive electrophysiological study is highly dependent on the degree of suspicion of the abnormality (pre-test probability), but also on the applied protocol, and the criteria used for diagnosing the presence of clinically significant abnormalities. There are four areas of interest: suspected bradycardia, bundle branch block (impending high degree AV block); suspected supraventricular tachycardia; suspected ventricular tachycardia (Table 2).

**Suspected bradycardia.** The pre-test probability of a transient symptomatic bradycardia as the cause of syncope is relatively high when there is asymptomatic sinus bradycardia (<50 bpm) or sinoatrial block in the absence of negatively chronotropic medications. Sinus node dysfunction can be demonstrated by a prolonged sinus node recovery time. The prognostic value of a prolonged sinus node recovery time is largely unknown. It is opinion of the panel that, in presence of a SNRT>2 s or CSNRT>1 s, sinus node dysfunction may be the cause of syncope.

**Bundle branch block.** In patients with syncope and bifascicular block, an electrophysiological study is highly sensitive in identifying patients with intermittent or impending high degree AV block (level B). This block is the likely cause of syncope in most cases, but not of the high mortality rate observed in these patients that seems mainly related to underlying structural heart disease and ventricular tachyarrhythmias (level B). Unfortunately, ventricular programmed stimulation does not seem to be able correctly to identify these patients and the finding of inducible ventricular arrhythmia should therefore be interpreted with caution.

**Suspected supraventricular tachycardia.** Supraventricular tachycardia presenting as syncope without accompanying palpitations is probably rare. Both non-invasive (transoesophageal) and invasive electrophysiological studies may be used to evaluate the haemodynamic effects of an induced tachycardia.

**Suspected ventricular tachycardia.** Electrophysiological study with programmed electrical stimulation is an effective diagnostic test in patients with coronary artery disease, markedly depressed cardiac function and unexplained syncope (level B). Its utility is more questionable in patients with non-ischaemic dilated cardiomyopathy (level B). Several studies on patients who underwent implantation of an automatic defibrillator showed a high incidence of spontaneous ventricular arrhythmia requiring device therapy, and suppression of syncopal recurrences (level B). However, these results applied to a highly selected, high-risk population that might be not representative of the patients encountered in clinical practice.
**Recommendations**

**Indications**

**Class I**
- An invasive electrophysiological procedure is indicated when the initial evaluation suggests an arrhythmic cause of syncope (in patients with abnormal electrocardiography and/or structural heart disease or syncope associated with palpitations or family history of sudden death).

**Class II**
- Diagnostic reasons: to evaluate the exact nature of an arrhythmia which has already been identified as the cause of the syncope.
- Prognostic reasons: in patients with cardiac disorders, in which arrhythmia induction has a bearing on the selection of therapy; and in patients with high-risk occupations, in whom every effort to exclude a cardiac cause of syncope is warranted.

**Class III**
- In patients with normal electrocardiograms and no heart disease and no palpitations an electrophysiological study is not usually undertaken.

**Diagnosis**

**Class I**
- Normal electrophysiological findings cannot completely exclude an arrhythmic cause of syncope; when an arrhythmia is likely, further evaluations (for example loop recording) are recommended.
- Depending on the clinical context, abnormal electrophysiological findings may not be diagnostic of the cause of syncope.
- An electrophysiological study is diagnostic, and usually no additional tests are required; in the following cases:
  - sinus bradycardia and a very prolonged CSNRT (as discussed in the text);
  - bifascicular block and:
    - a baseline HV interval of ≥100 ms, or
    - 2nd or 3rd degree His–Purkinje block is demonstrated during incremental atrial pacing, or
    - (if the baseline electrophysiological study is inconclusive) high-degree His–Purkinje block is provoked by intravenous administration of ajmaline, procainamide, or disopyramide;
  - previous myocardial infarction and induction of sustained monomorphic ventricular tachycardia;
  - arrhythmogenic right ventricular dysplasia and induction of ventricular tachyarrhythmias;

  — induction of rapid supraventricular arrhythmia which reproduces hypotensive or spontaneous symptoms.

**Class II**
- Divergence of opinion exists on the diagnostic value of electrophysiological study in case of:
  - HV interval of >70 ms but <100 ms;
  - induction of polymorphic ventricular tachycardia or ventricular fibrillation in patients with ischaemic or dilated cardiomyopathy;
  - Brugada syndrome.

**ATP test**

Intravenous injection of adenosine triphosphate (ATP) has recently been proposed as a tool in the investigation of patients with unexplained syncope. In predisposed patients with unexplained syncope, the stimulation of purinergic receptors causes prolonged ventricular pauses due to atrioventricular block, which are considered as possibly responsible for spontaneous attacks (level B).

**Recommendations**

The test requires the rapid injection of a 20 mg bolus of ATP during electrocardiographic monitoring. Asystole lasting more than 6 s, or AV block lasting more than 10 s, is considered abnormal. ATP testing produces an abnormal response in some patients with syncope of unknown origin, but not in controls. The diagnostic and predictive value of the test remains to be confirmed by prospective studies. In the absence of sufficient hard data, the test may be indicated at the end of the diagnostic work-up (Class II).

**Ventricular signal-averaged electrocardiogram**

**Recommendations**

There is general agreement that ventricular signal-averaged electrocardiogram is not diagnostic of the cause of syncope. In patients with syncope and no evidence of structural heart disease, the technique may be useful for guiding the use of electrophysiological studies. Its systematic use is not recommended (Class III).

**Echocardiogram**

Even if echocardiography alone is only seldom diagnostic, this test provides information about the type and severity of underlying heart disease which may be useful for risk stratification. If moderate to severe structural
heart disease is found, evaluation is directed toward a cardiac cause of syncope. On the other hand, in the presence of minor structural abnormalities detected by echocardiography, the probability of cardiac cause of syncope may not be high, and the evaluation may proceed as in patients without structural heart disease.

**Recommendations**

**Indications**

*Class I*

- Echocardiography is recommended in patients with syncope when cardiac disease is suspected.

**Diagnosis**

*Class I*

- Echocardiographic findings may be useful to stratify the risk by assessing the cardiac substrate.
- Echocardiography only makes a diagnosis in severe aortic stenosis and atrial myxoma.

**Exercise testing**

Syncope occurring during exercise may be cardiac (level B), even if some case reports showed that it might be a manifestation of an exaggerated reflex vasodilatation. By contrast, postexertional syncope is almost invariably due to autonomic failure or to a neurally-mediated mechanism (level B).

**Recommendations**

**Indications**

*Class I*

Patients who experience an episode of syncope during or shortly after exertion.

*Class III*

Use of exercise testing is not recommended in patients who do not experience syncope during exercise.

**Diagnosis**

*Class I*

- Exercise testing is diagnostic when ECG and haemodynamic abnormalities are present and syncope is reproduced during or immediately after exercise.
- Exercise testing is diagnostic if Mobitz II second degree or 3rd degree AV block develop during exercise even without syncope.

**Cardiac catheterization and angiography**

**Recommendations**

**Indications**

*Class I*

In patients with syncope suspected to be due, directly or indirectly, to myocardial ischaemia, coronary angiography is recommended in order to confirm the diagnosis and to establish optimal therapy.

*Class III*

Angiography alone is rarely diagnostic for the cause of syncope.

**Neurological and psychiatric evaluation**

**Recommendations**

**Indications**

*Class I*

- Neurological referral is indicated in patients in whom loss of consciousness cannot be attributed to syncope.
- In case of unequivocal syncope, neurological referral is warranted when syncope may be due to autonomic failure or to a cerebrovascular steal syndrome.
- Psychiatric evaluation is recommended when symptoms suggest psychogenic syncope (somatization disorder) or if the patient has a known psychiatric disorder.

*Class III*

- In all other patients with syncope, neurological and psychiatric investigations are not recommended.

**Treatment**

Neurally-mediated reflex syncopeal syndromes

Treatment goals: primarily prevention of symptom recurrence and associated injuries; improved quality of life.

Patients who seek medical advice after having experienced a vasovagal faint principally require reassurance and education regarding the nature of the condition. This assumption is derived from the knowledge of the benign nature of the disease. In particular, based on
review of their medical history, patients should be informed of the likelihood of syncope recurrence. Initial advice should also include review of typical premonitory symptoms which may permit many individuals to recognize an impending episode and thereby avert a frank faint. In general, initial ‘treatment’ of all forms of neurally-mediated reflex syncope comprises education regarding avoidance of triggering events (e.g. hot crowded environments, volume depletion, effects of cough, tight collars, etc.), recognition of premonitory symptoms, and manoeuvres to abort the episode (e.g. supine posture). Additionally, if possible, strategies should address trigger factors directly (for example, suppressing the cause of cough in cough syncope).

When a more aggressive treatment strategy is needed, ‘volume expanders’ (e.g. increased dietary salt/electrolyte intake with fluids [e.g. ‘sport’ drinks, salt tablets]) or moderate exercise training appear to be among the safest initial approaches (level B). Additionally, in highly motivated patients with recurrent vasovagal symptoms, the prescription of progressively prolonged periods of enforced upright posture (so-called ‘tilt-training’) may reduce syncope recurrence (level B).

Many drugs have been used in the treatment of vasovagal syncope (beta-blockers, disopyramide, scopolamine, clonidine, theophylline, fludrocortisone, ephedrine, etilefrine, midodrine, clonidine, serotonin reuptake inhibitors, etc.). In general, while the results have been satisfactory in uncontrolled trials or short-term controlled trials, long-term placebo-controlled prospective trials have been unable to show a benefit of the active drug over placebo. Beta-adrenergic blocking drugs have failed to be effective in several long-term follow-up controlled studies. Thus the evidence fails to support beta-blocker efficacy (level A). Vasconstrictor drugs are potentially more effective in orthostatic hypotension caused by autonomic dysfunction than in the neurally-mediated syncope. Etilerfrine proved to be ineffective (level B). Cardiac pacing has been demonstrated to be effective in highly selected patients affected by cardioinhibitory form (level B).

Cardiac pacing appears to be beneficial in carotid sinus syndrome (level B) and is acknowledged to be the treatment of choice when bradycardia has been documented.

**Recommendations**

It is valuable to assess the relative contribution of cardioinhibition and vasodepression before embarking on specific treatment as there are different therapeutic strategies for the two aspects. Even if evidence of utility of such an assessment exists only for the carotid sinus massage, it is recommended to extend this assessment also by means of tilt testing or implantable loop recorder.

Patients who have syncope in a ‘high risk’ setting (e.g. commercial vehicle driver, machine operator, pilot, commercial painter, competitive athlete) merit specific consideration for treatment. There is no information available regarding the efficacy of treatment in this type of patient, and whether it differs from other patients with neurally-mediated fainting.

Treatment is not necessary in patients who have sustained a single syncope and are not having syncope in a high risk setting.

**Class I**

- Explanation of the risk, and reassurance about the prognosis in vasovagal syncope.
- Avoidance of trigger events as much as possible and reducing magnitude of potential triggers when feasible (e.g. emotional upset) and causal situation in situational syncope.
- Modification or discontinuation of hypotensive drug treatment for concomitant conditions.
- Cardiac pacing in patients with cardioinhibitory or mixed carotid sinus syndrome.

**Class II**

- Volume expansion by salt supplements, an exercise programme or sleeping >10° head-up in posture-related syncope.
- Cardiac pacing in patients with cardioinhibitory vasovagal syncope with a frequency >5 attacks per year or severe physical injury or accident and age >40.
- Tilt training in patients with vasovagal syncope.

**Class III**

- The evidence fails to support the efficacy of beta-adrenergic blocking drugs. Beta-adrenergic blocking drugs may aggravate bradycardia in some cardioinhibitory cases.

**Orthostatic hypotension**

Treatment goals: prevention of symptom recurrence and associated injuries; improved quality of life.

Drug-induced autonomic failure is probably the most frequent cause of orthostatic hypotension. The principal treatment strategy is elimination of the offending agent. It is reasonable for all patients to receive advice and education on factors that influence systemic blood pressure, such as avoiding sudden head-up postural change (especially on waking), standing still for a prolonged period of time, prolonged recumbence during daytime, straining during micturition and defeacation, hyperventilation, high environmental temperature, severe exertion, large meals and alcohol.

Additional treatment principles, used alone or in combination, are appropriate for consideration on an individual patient basis are chronic expansion of intravascular volume by encouraging a higher than normal salt intake and fluid intake; use of fludrocortisone in low dose; and raising the head of the bed on blocks to permit gravitational exposure during sleep (level B). Midodrine
appears to be of particular interest given the rapidly expanding and generally positive experience (level B).

**Recommendations**

**Class I**

- Syncope due to orthostatic hypotension should be treated in all patients. In many instances treatment entails only modification of drug treatment for concomitant conditions.

**Cardiac arrhythmias as primary cause**

Treatment goals: prevention of symptom recurrence, improved quality of life, reduction of mortality risk.

*Sinus node dysfunction (including bradycardia-tachycardia syndrome).* In general, cardiac pacemaker therapy is indicated and has proved highly effective in patients with sinus node dysfunction when bradyarrhythmia has been demonstrated to account for syncope (Class I, level B).

*AV conduction system disease.* Pacing is able to improve survival in patients with heart block as well as prevent syncopal recurrences (Class I, level B). Pacing may also be life-saving in patients with bundle branch block and syncope in whom the mechanism of the faint is suspected to be intermittent AV block. However, it is also critical to consider the possibility that ventricular tachyarrhythmias are responsible for loss of consciousness, since many patients who present with varying degrees of conduction system disease have significant concomitant left ventricular dysfunction.

*Paroxysmal supraventricular and ventricular tachycardias.* Transcatheter ablation has become a very cost-effective treatment option and in paroxysmal supraventricular arrhythmia associated with syncope is probably the treatment of choice (Class I).

In the case of syncope due to ventricular tachycardia, drug therapy may be useful in the setting of normal heart or of heart disease with mild cardiac dysfunction. In patients with depressed cardiac function, the use of implantable pacemaker cardioverter-defibrillators (ICDs) is warranted. Currently, ablation techniques are appropriate first choices in only a few forms of ventricular tachycardia, specifically right ventricular outflow tract tachycardia, bundle-branch reentry tachycardia, and so-called verapamil sensitive left ventricular tachycardias (Table 3).

**Recommendations**

**Class I**

- Syncope due to cardiac arrhythmias must receive treatment appropriate to the cause in all patients in whom it is life-threatening and when there is a high risk of injury.

**Class II**

- Treatment may be employed when the culprit arrhythmia has not been demonstrated and a diagnosis of life-threatening arrhythmia is presumed from surrogate data.
- Treatment may be employed when a culprit arrhythmia has been identified but is not life-threatening or presenting a high risk of injury.

**Structural cardiac or cardiopulmonary disease**

Structural heart disease can cause syncope when circulatory demands outweigh the impaired ability of the heart to increase its output. More than one pathophysiological factor may contribute to the symptoms. Syncope is not solely the result of restricted cardiac output, but may be in part due to inappropriate neurally mediated reflex vasodilation and/or primary cardiac

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**Table 3 Situations in which ICD therapy is likely to be useful**

- Documented syncopal ventricular tachycardia or fibrillation without correctable causes (e.g. drug-induced) (Class I, level A).
- Undocumented syncope likely to be due to ventricular tachycardia or fibrillation:
  - previous myocardial infarction and inducible sustained monomorphic ventricular tachycardia with severe haodynamic compromise, in the absence of another competing diagnosis as a cause of syncope (Class I, level B).
  - unexplained syncope in patients with depressed left ventricular systolic function in the absence of another competing diagnosis as a cause of syncope (Class II, level B).
  - arrhythmogenic right ventricular dysplasia, or hypertrophic obstructive cardiomyopathy, with a family history of sudden death, in the absence of another competing diagnosis for the cause of syncope (Class II).
  - Brugada syndrome or arrhythmogenic right ventricular dysplasia and inducible ventricular tachyarrhythmias with severe haodynamic compromise in the absence of another competing diagnosis for the cause of syncope (Class II).

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arrhythmias. However, the management is primarily that of the underlying disease.

**Recommendations**

**Class I**

Treatment is best directed at amelioration of the specific structural lesion or its consequences.

**Vascular steal syndromes**

Subclavian steal is rare but is the most commonly recognized condition in this group. Direct corrective angioplasty or surgery is usually feasible and effective (Class I).