# 2017 ACC/AHA/HRS Guideline for the Evaluation and **Management of Patients With Syncope**

A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines, and the Heart Rhythm Society

Developed in Collaboration With the American College of Emergency Physicians and Society for Academic Emergency Medicine

Endorsed by the Pediatric and Congenital Electrophysiology Society

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# Circulation

#### **Preamble**

Since 1980, the American College of Cardiology (ACC) and American Heart Association (AHA) have translated scientific evidence into clinical practice guidelines (guidelines) with recommendations to improve cardiovascular health. These guidelines, which are based on systematic methods to evaluate and classify evidence, provide a cornerstone for quality cardiovascular care. The ACC and AHA sponsor the development and publication of guidelines without commercial support, and members of each organization volunteer their time to the writing and review efforts. Guidelines are official policy of the ACC and AHA.

#### **Intended Use**

Practice guidelines provide recommendations applicable to patients with or at risk of developing cardiovascular disease. The focus is on medical practice in the United States, but guidelines developed in collaboration with other organizations may have a global impact. Although guidelines may be used to inform regulatory or payer decisions, their intent is to improve patients' quality of care and align with patients' interests. Guidelines are intended to define practices meeting the needs of patients in most, but not all, circumstances and should not replace clinical judgment.

#### **Clinical Implementation**

Guideline recommended management is effective only when followed by healthcare providers and patients. Adherence to recommendations can be enhanced by shared decision making between healthcare providers and patients, with patient engagement in selecting interventions based on individual values, preferences, and associated conditions and comorbidities.

#### **Methodology and Modernization**

The ACC/AHA Task Force on Clinical Practice Guidelines (Task Force) continuously reviews, updates, and modifies guideline methodology on the basis of published standards from organizations including the Institute of Medicine (1,2) and on the basis of internal reevaluation. Similarly, the presentation and delivery of guidelines are reevaluated and modified on the basis of evolving technologies and other factors to facilitate optimal dissemination of information at the point of care to healthcare professionals. Given time constraints of busy healthcare providers and the need to limit text, the current guideline format delineates that each recommendation be supported by limited text (ideally, <250 words) and hyperlinks to supportive evidence summary tables. Ongoing efforts to further limit text are underway. Recognizing the importance of cost–value considerations in certain guidelines, when appropriate and feasible, an analysis of the value of a drug, device, or intervention may be performed in accordance with the ACC/AHA methodology (3).

To ensure that guideline recommendations remain current, new data are reviewed on an ongoing basis, with full guideline revisions commissioned in approximately 6-year cycles. Publication of new, potentially practice-changing study results that are relevant to an existing or new drug, device, or management strategy will prompt evaluation by the Task Force, in consultation with the relevant guideline writing committee, to determine whether a focused update should be commissioned. For additional information and policies regarding guideline development, we encourage readers to consult the ACC/AHA guideline methodology manual (4) and other methodology articles (5-8).

#### **Selection of Writing Committee Members**

The Task Force strives to avoid bias by selecting experts from a broad array of backgrounds. Writing committee members represent different geographic regions, sexes, ethnicities, races, intellectual perspectives/biases, and

scopes of clinical practice. The Task Force may also invite organizations and professional societies with related interests and expertise to participate as partners, collaborators, or endorsers.

#### **Relationships With Industry and Other Entities**

The ACC and AHA have rigorous policies and methods to ensure that guidelines are developed without bias or improper influence. The complete relationships with industry and other entities (RWI) policy can be found at <a href="http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/relationships-with-industry-policy">http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/relationships-with-industry-policy</a>. Appendix 1 of the current document lists writing committee members' relevant RWI. For the purposes of full transparency, writing committee members' comprehensive disclosure information is available online (<a href="http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIR.00000000000000499/-/DC1">http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIR.0000000000000000499/-/DC1</a>). Comprehensive disclosure information for the Task Force is available at <a href="http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/guidelines-and-documents-task-forces">http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/guidelines-and-documents-task-forces</a>.

#### **Evidence Review and Evidence Review Committees**

When developing recommendations, the writing committee uses evidence-based methodologies that are based on all available data (4-7). Literature searches focus on randomized controlled trials (RCTs) but also include registries, nonrandomized comparative and descriptive studies, case series, cohort studies, systematic reviews, and expert opinion. Only key references are cited.

An independent evidence review committee (ERC) is commissioned when there are 1 or more questions deemed of utmost clinical importance that merit formal systematic review. This systematic review will determine which patients are most likely to benefit from a drug, device, or treatment strategy and to what degree. Criteria for commissioning an ERC and formal systematic review include: a) the absence of a current authoritative systematic review, b) the feasibility of defining the benefit and risk in a time frame consistent with the writing of a guideline, c) the relevance to a substantial number of patients, and d) the likelihood that the findings can be translated into actionable recommendations. ERC members may include methodologists, epidemiologists, healthcare providers, and biostatisticians. The recommendations developed by the writing committee on the basis of the systematic review are marked with "SR".

#### **Guideline-Directed Management and Therapy**

The term *guideline-directed management and therapy* (GDMT) encompasses clinical evaluation, diagnostic testing, and pharmacological and procedural treatments. For these and all recommended drug treatment regimens, the reader should confirm the dosage by reviewing product insert material and evaluate the treatment regimen for contraindications and interactions. The recommendations are limited to drugs, devices, and treatments approved for clinical use in the United States.

#### Class of Recommendation and Level of Evidence

The Class of Recommendation (COR) indicates the strength of the recommendation, encompassing the estimated magnitude and certainty of benefit in proportion to risk. The Level of Evidence (LOE) rates the quality of scientific evidence that supports the intervention on the basis of the type, quantity, and consistency of data from clinical trials and other sources (Table 1) (4-6).

Glenn N. Levine, MD, FACC, FAHA Chair, ACC/AHA Task Force on Clinical Practice Guidelines

# Table 1. Applying Class of Recommendation and Level of Evidence to Clinical Strategies, Interventions, Treatments, or Diagnostic Testing in Patient Care\* (Updated August 2015)

#### **CLASS (STRENGTH) OF RECOMMENDATION** CLASS I (STRONG) Benefit >>> Risk Suggested phrases for writing recommendations: Is recommended Is indicated/useful/effective/beneficial Should be performed/administered/other Comparative-Effectiveness Phrases†: • Treatment/strategy A is recommended/indicated in preference to treatment B Treatment A should be chosen over treatment B Suggested phrases for writing recommendations: Is reasonable Can be useful/effective/beneficial Comparative-Effectiveness Phrases†: • Treatment/strategy A is probably recommended/indicated in preference to treatment B It is reasonable to choose treatment A over treatment B CLASS IIb (WEAK) **Benefit** ≥ **Risk** Suggested phrases for writing recommendations: ■ May/might be reasonable May/might be considered Usefulness/effectiveness is unknown/unclear/uncertain or not well established CLASS III: No Benefit (MODERATE) Benefit = Risk Suggested phrases for writing recommendations: Is not recommended ■ Is not indicated/useful/effective/beneficial Should not be performed/administered/other CLASS III: Harm (STRONG) Risk > Benefit Suggested phrases for writing recommendations: Potentially harmful Causes harm Associated with excess morbidity/mortality

Should not be performed/administered/other

#### LEVEL (QUALITY) OF EVIDENCE‡

#### **LEVEL A**

- High-quality evidence‡ from more than 1 RCT
- Meta-analyses of high-quality RCTs
- One or more RCTs corroborated by high-quality registry studies

#### **LEVEL B-R**

(Randomized)

- Moderate-quality evidence‡ from 1 or more RCTs
- Meta-analyses of moderate-quality RCTs

#### **LEVEL B-NR**

(Nonrandomized)

- Moderate-quality evidence‡ from 1 or more well-designed, well-executed nonrandomized studies, observational studies, or registry studies
- Meta-analyses of such studies

#### **LEVEL C-LD**

(Limited Data)

- Randomized or nonrandomized observational or registry studies with limitations of design or execution
- Meta-analyses of such studies
- Physiological or mechanistic studies in human subjects

#### LEVEL C-EO

(Expert Opinion

Consensus of expert opinion based on clinical experience

COR and LOE are determined independently (any COR may be paired with any LOE).

A recommendation with LOE C does not imply that the recommendation is weak. Many important clinical questions addressed in guidelines do not lend themselves to clinical trials. Although RCTs are unavailable, there may be a very clear clinical consensus that a particular test or therapy is useful or effective.

- \* The outcome or result of the intervention should be specified (an improved clinical outcome or increased diagnostic accuracy or incremental prognostic information).
- † For comparative-effectiveness recommendations (COR I and IIa; LOE A and B only), studies that support the use of comparator verbs should involve direct comparisons of the treatments or strategies being evaluated.
- ‡ The method of assessing quality is evolving, including the application of standardized, widely used, and preferably validated evidence grading tools; and for systematic reviews, the incorporation of an Evidence Review Committee.

COR indicates Class of Recommendation; EO, expert opinion; LD, limited data; LOE, Level of Evidence; NR, nonrandomized; R, randomized; and RCT, randomized controlled trial.

#### 1. Introduction

## 1.1. Methodology and Evidence Review

The recommendations listed in this guideline are, whenever possible, evidence based. An initial extensive evidence review, which included literature derived from research involving human subjects, published in English, and indexed in MEDLINE (through PubMed), EMBASE, the Cochrane Library, the Agency for Healthcare Research and Quality, and other selected databases relevant to this guideline, was conducted from July to October 2015. Key search words included but were not limited to the following: athletes, autonomic neuropathy, bradycardia, carotid sinus hypersensitivity, carotid sinus syndrome, children, death, dehydration, diagnosis, driving, electrocardiogram, electrophysiological study, epidemiology, falls, implantable loop recorder, mortality, older populations, orthostatic hypotension, pediatrics, psychogenic pseudosyncope, recurrent syncope, risk stratification, supraventricular tachycardia, syncope unit, syncope, tilt-table test, vasovagal syncope, and ventricular arrhythmia. Additional relevant studies published through October 2016, during the guideline writing process, were also considered by the writing committee and added to the evidence tables when appropriate. The finalized evidence tables, included in the Online Data Supplement (http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIR.0000000000000499/-/DC2), summarize the evidence used by the writing committee to formulate recommendations. Lastly, the writing committee reviewed documents related to syncope previously published by the ACC and AHA and other organizations and societies. References selected and published in this document are representative and not all inclusive.

An independent ERC was commissioned to perform a systematic review of clinical questions, the results of which were considered by the writing committee for incorporation into this guideline. The systematic review report "Pacing as a Treatment for Reflex-Mediated (Vasovagal, Situational, or Carotid Sinus Hypersensitivity) Syncope" is published in conjunction with this guideline (9).

## 1.2. Organization of the Writing Committee

The writing committee was composed of clinicians with expertise in caring for patients with syncope, including cardiologists, electrophysiologists, a neurologist, an emergency physician, and a pediatric cardiologist. The writing committee included representatives from the ACC, AHA, Heart Rhythm Society (HRS), American Academy of Neurology, American College of Emergency Physicians, and Society for Academic Emergency Medicine.

# 1.3. Document Review and Approval

This document was reviewed by 2 official reviewers each nominated by the ACC, AHA, and HRS; 1 reviewer each from the American Academy of Neurology, American College of Emergency Physicians and Society for © 2017 by the American College of Cardiology Foundation, American Heart Association, Inc., and Heart Rhythm Society

Academic Emergency Medicine, and Pediatric and Congenital Electrophysiology Society; a lay/patient representative; and 25 individual content reviewers. Reviewers' RWI information was distributed to the writing committee and is published in this document (Appendix 2).

This document was approved for publication by the governing bodies of the ACC, AHA, and HRS and was endorsed by the Pediatric and Congenital Electrophysiology Society.

## 1.4. Scope of the Guideline

The purpose of this ACC/AHA/HRS guideline is to provide contemporary, accessible, and succinct guidance on the management of adult and pediatric patients with suspected syncope. This guideline is intended to be a practical document for cardiologists, arrhythmia specialists, neurologists, emergency physicians, general internists, geriatric specialists, sports medicine specialists, and other healthcare professionals involved in the care of this very large and heterogeneous population. It is not a review of physiology, pathophysiology, or mechanisms of underlying conditions associated with syncope. The nature of syncope as a symptom required that the writing committee consider numerous conditions for which it can be a symptom, and as much as possible, we have addressed the involvement of syncope only as a presenting symptom. Because of the plausible association of syncope and sudden cardiac death (SCD) in selected populations, this document discusses risk stratification and prevention of SCD when appropriate. The use of the terms selected populations and selected patients in this document is intended to direct healthcare providers to exercise clinical judgment, which is often required during the evaluation and management of patients with syncope. When a recommendation is made to refer a patient to a specialist with expertise for further evaluation, such as in the case of autonomic neurology, adult congenital heart disease (ACHD), older populations, or athletes, the writing committee agreed to make Class IIa recommendations because of the paucity of outcome data. The definition of older populations has been evolving. Age >75 years is used to define older populations or older adults in this document, unless otherwise specified. If a study has defined older adults by a different age cutoff, the relevant age is noted in those specific cases. Finally, the guideline addresses the management of syncope with the patient as a focus, rather than larger aspects of health services, such as syncope management units. The goals of the present guideline are:

- To define syncope as a symptom, with different causes, in different populations and circumstances.
- To provide guidance and recommendations on the evaluation and management of patients with suspected syncope in the context of different clinical settings, specific causes, or selected circumstances.
- To identify key areas in which knowledge is lacking, to foster future collaborative research opportunities and efforts.

In developing this guideline, the writing committee reviewed the evidence to support recommendations in the relevant ACC/AHA guidelines noted in Table 2 and affirms the ongoing validity of the related recommendations in the context of syncope, thus obviating the need to repeat existing guideline

recommendations in the present guideline when applicable or when appropriate. Table 2 also contains a list of other statements that may be of interest to the reader.

Table 2. Relevant ACC/AHA Guidelines

Table 2. Relevant ACC/AHA Guidelines  Title	Organization	Publication Year (Reference)	
ACC/AHA guideline policy relevant to the management of	of syncope	, , , , , , , , ,	
Supraventricular tachycardia	ACC/AHA/HRS	2015 (10)	
Valvular heart disease	AHA/ACC	2014 (11)	
Device-based therapies for cardiac rhythm abnormalities	ACCF/AHA/HRS	2012 (12)	
Ventricular arrhythmias and sudden cardiac death	ACC/AHA/ESC	2006 (13)*	
Other ACC/AHA guidelines of interest			
Hypertension*	ACC/AHA		
Stable ischemic heart disease	ACC/AHA/ACP/	2012 and 2014	
	AATS/PCNA/SCAI/STS	(14,15)	
Atrial fibrillation	AHA/ACC/HRS	2014 (16)	
Non–ST-elevation acute coronary syndromes	AHA/ACC	2014 (17)	
Assessment of cardiovascular risk	ACC/AHA	2013 (18)	
Heart failure	ACC/AHA	2013 (19)*	
Hypertrophic cardiomyopathy	ACC/AHA	2011 (20)	
Assessment of cardiovascular risk in asymptomatic adults	ACC/AHA	2010 (21) ation.	
Adult congenital heart disease	ACC/AHA	2008 (22)*	
Other related references			
Scientific statement on electrocardiographic early repolarization	АНА	2016 (23)	
Expert consensus statement on the diagnosis and treatment of postural tachycardia syndrome, inappropriate sinus tachycardia, and vasovagal syncope	HRS	2015 (24)	
Guidelines for the management of patients with ventricular arrhythmias and the prevention of sudden cardiac death	ESC	2015 and 2013 (25,26)	
Expert consensus statement on the recognition and management of arrhythmias in adult congenital heart disease	PACES/HRS	2014 (27)	
Expert consensus statement on the use of implantable cardioverter-defibrillator therapy in patients who are not included or not well represented in clinical trials	HRS/ACC/AHA	2014 (28)	
Expert consensus statement on ventricular arrhythmias	EHRA/HRS/APHRS	2014 (29)	
Expert consensus statement on the diagnosis and management of patients with inherited primary arrhythmia syndromes	HRS/EHRA/APHRS	2013 (25)	
Guidelines for the diagnosis and management of syncope	ESC	2009 (30)	

<sup>\*</sup>Revisions to the current documents are being prepared, with publication expected in 2017.

AATS indicates American Association for Thoracic Surgery; ACC, American College of Cardiology; ACCF, American College of Cardiology Foundation; ACP, American College of Physicians; AHA, American Heart Association; APHRS, Asia Pacific Heart Rhythm Society; EHRA, European Heart Rhythm Association; ESC, European Society of Cardiology; HRS, Heart Rhythm Society; PACES, Pediatric and Congenital Electrophysiology Society; PCNA, Preventive Cardiovascular Nurses Association; SCAI, Society for Cardiovascular Angiography and Interventions; and STS, Society of Thoracic Surgery.

# 2. General Principles

## 2.1. Definitions: Terms and Classification

For the purpose of this guideline, definitions of syncope and relevant terms are provided in Table 3.

**Table 3. Relevant Terms and Definitions\*** 

Term	Definition/Comments and References
Syncope	A symptom that presents with an abrupt, transient, complete loss of consciousness, associated with inability to maintain postural tone, with rapid and spontaneous recovery. The presumed mechanism is cerebral hypoperfusion (24,30). There should not be clinical features of other nonsyncope causes of loss of consciousness, such as seizure, antecedent head trauma, or apparent loss of consciousness (i.e., pseudosyncope) (24,30).
Loss of consciousness	A cognitive state in which one lacks awareness of oneself and one's situation, with an inability to respond to stimuli.
Transient loss of consciousness	Self-limited loss of consciousness (30) can be divided into syncope and nonsyncope conditions. Nonsyncope conditions include but are not limited to seizures, hypoglycemia, metabolic conditions, drug or alcohol intoxication, and concussion due to head trauma. The underlying mechanism of syncope is presumed to be cerebral hypoperfusion, whereas nonsyncope conditions are attributed to different mechanisms.
Presyncope (near-syncope)	The symptoms before syncope. These symptoms could include extreme lightheadedness; visual sensations, such as "tunnel vision" or "graying out"; and variable degrees of altered consciousness without complete loss of consciousness. Presyncope could progress to syncope, or it could abort without syncope.
Unexplained syncope (syncope of undetermined etiology)	Syncope for which a cause is undetermined after an initial evaluation that is deemed appropriate by the experienced healthcare provider. The initial evaluation includes but is not limited to a thorough history, physical examination, and ECG.
Orthostatic intolerance	A syndrome consisting of a constellation of symptoms that include frequent, recurrent, or persistent lightheadedness, palpitations, tremulousness, generalized weakness, blurred vision, exercise intolerance, and fatigue upon standing. These symptoms can occur with or without orthostatic tachycardia, OH, or syncope (24). Individuals with orthostatic intolerance have ≥1 of these symptoms associated with reduced ability to maintain upright posture.
Orthostatic tachycardia	A sustained increase in heart rate of $\geq 30$ bpm within 10 min of moving from a recumbent to a quiet (nonexertional) standing position (or $\geq 40$ bpm in individuals 12–19 y of age) (24,30,31).
Orthostatic hypotension (OH)	A drop in systolic BP of $\geq$ 20 mm Hg or diastolic BP of $\geq$ 10 mm Hg with assumption of an upright posture (31).
• Initial (immediate) OH	A transient BP decrease within 15 s after standing, with presyncope or syncope (31,32).
• Classic OH	A sustained reduction of systolic BP of $\geq$ 20 mm Hg or diastolic BP of $\geq$ 10 mm Hg within 3 min of assuming upright posture (31).
• Delayed OH	A sustained reduction of systolic BP of $\geq$ 20 mm Hg (or 30 mm Hg in patients with supine hypertension) or diastolic BP of $\geq$ 10 mm Hg that takes $>$ 3 min of upright posture to develop. The fall in BP is usually gradual until reaching the threshold (31).
• Neurogenic OH	A subtype of OH that is due to dysfunction of the autonomic nervous system and not solely due to environmental triggers (e.g., dehydration or drugs) (33,34). Neurogenic OH is due to lesions involving the central or peripheral autonomic nerves.
Cardiac (cardiovascular) syncope	Syncope caused by bradycardia, tachycardia, or hypotension due to low cardiac index, blood flow obstruction, vasodilatation, or acute vascular dissection (35,36).
Noncardiac syncope	Syncope due to noncardiac causes which include reflex syncope, OH, volume depletion, dehydration, and blood loss (35).

Reflex (neurally	Syncope due to a reflex that causes vasodilation, bradycardia, or both (24,30,31).
mediated) syncope  • Vasovagal syncope (VVS)	The most common form of reflex syncope mediated by the vasovagal reflex. VVS 1) may occur with upright posture (standing or seated or with exposure to emotional stress, pain, or medical settings; 2) typically is characterized by diaphoresis, warmth, nausea, and pallor; 3) is associated with vasodepressor hypotension and/or inappropriate bradycardia; and 4) is often followed by fatigue. Typical features may be absent in older patients (24). VVS is often preceded by identifiable triggers and/or by a characteristic prodrome. The diagnosis is made primarily on the basis of a thorough history, physical examination, and eyewitness observation, if available.
Carotid sinus syndrome	Reflex syncope associated with carotid sinus hypersensitivity (30). Carotid sinus hypersensitivity is present when a pause ≥3 s and/or a decrease of systolic pressure ≥50 mm Hg occurs upon stimulation of the carotid sinus. It occurs more frequently in older patients. Carotid sinus hypersensitivity can be associated with varying degrees of symptoms. Carotid sinus syndrome is defined when syncope occurs in the presence of carotid sinus hypersensitivity.
Situational syncope	Reflex syncope associated with a specific action, such as coughing, laughing, swallowing, micturition, or defecation. These syncope events are closely associated with specific physical functions.
Postural (orthostatic) tachycardia syndrome (POTS)	A clinical syndrome usually characterized by all of the following: 1) frequent symptoms that occur with standing (e.g., lightheadedness, palpitations, tremulousness, generalized weakness, blurred vision, exercise intolerance, and fatigue); and 2) an increase in heart rate of ≥30 bpm during a positional change from supine to standing (or ≥40 bpm in those 12–19 y of age); and 3) the absence of OH (>20 mm Hg reduction in systolic BP). Symptoms associated with POTS include those that occur with standing (e.g., lightheadedness, palpitations); those not associated with particular postures (e.g., bloating, nausea, diarrhea, abdominal pain); and those that are systemic (e.g., fatigue, sleep disturbance, migraine headaches) (37). The standing heart rate is often >120 bpm (31,38-42).
Psychogenic pseudosyncope	A syndrome of <i>apparent</i> but not true loss of consciousness that may occur in the absence of identifiable cardiac, reflex, neurological, or metabolic causes (30).

<sup>\*</sup>These definitions are derived from previously published definitions from scientific investigations, guidelines, expert consensus statements, and Webster dictionary after obtaining consensus from the WC

BP indicates blood pressure; ECG, electrocardiogram; OH, orthostatic hypotension; POTS, postural tachycardia syndrome; and VVS, vasovagal syncope.

# 2.2. Epidemiology and Demographics

Syncope has many causes and clinical presentations; the incidence depends on the population being evaluated. Estimates of isolated or recurrent syncope may be inaccurate and underestimated because epidemiological data have not been collected in a consistent fashion or because a consistent definition has not been used. Interpretation of the symptoms varies among the patients, observers, and healthcare providers. The evaluation is further obscured by inaccuracy of data collection and by improper diagnosis.

Studies of syncope report prevalence rates as high as 41%, with recurrent syncope occurring in 13.5% (43). In a cross section of 1,925 randomly selected residents of Olmsted County, MN, with a median age of 62 years (all age >45 years), 364 reported an episode of syncope in their lifetime; the estimated prevalence of syncope was 19%. Females reported a higher prevalence of syncope (22% versus 15%, p<0.001) (44). The incidence follows a trimodal distribution in both sexes, with the first episode common around 20, 60, or 80 years of age and the third peak occurring 5 to 7 years earlier in males (45). Predictors of recurrent syncope in older adults are aortic stenosis, impaired renal function, atrioventricular (AV) or left bundle-branch block, male sex, © 2017 by the American College of Cardiology Foundation, American Heart Association, Inc., and Heart Rhythm Society

chronic obstructive pulmonary disorder, heart failure (HF), atrial fibrillation (AF), advancing age, and orthostatic medications (45), with a sharp increase in incidence after 70 years of age (35). Reflex syncope was most common (21%), followed by cardiac syncope (9%) and orthostatic hypotension (OH) (9%), with the cause of syncope unknown in 37% (35). In patients with New York Heart Association class III–IV HF, syncope is present in 12% to 14% of patients (46,47).

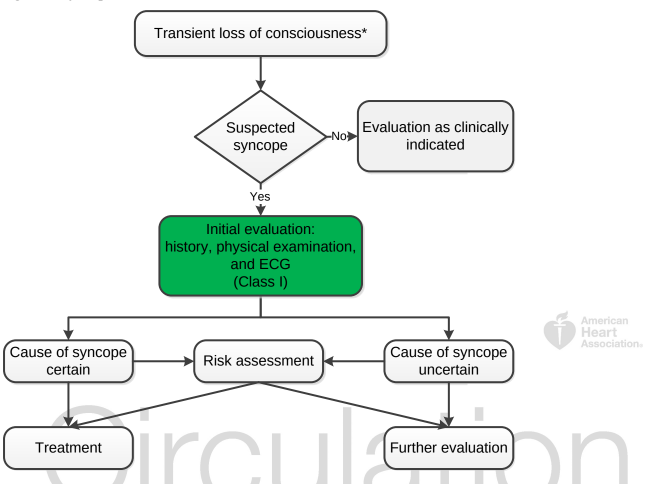
In older adults, there is a greater risk of hospitalization and death related to syncope. The National Hospital Ambulatory Medical Care Survey reported 6.7 million episodes of syncope in the emergency department (ED), or 0.77% of all ED patients. Among patients >80 years of age, 58% were admitted to hospital (48). The prevalence of syncope as a presenting symptom to the ED ranged from 0.8% to 2.4% in multiple studies in both academic and community settings (49-55).

Older institutionalized patients have a 7% annual incidence of syncope, a 23% overall prevalence, and a 30% 2-year recurrence rate (56). The incidence of syncope in older adults may overlap with falls, so it may be difficult to distinguish one from the other. Older adults are predisposed to falls when syncope occurs, with a 1-year fall rate of 38% among fainters versus 18.3% among nonfainters (57).

### 2.3. Initial Evaluation of Patients With Syncope

The time interval between the index syncopal event and the initial evaluation can vary significantly according to the medical necessity for evaluation and the patient's effort in seeking evaluation. The clinical setting in which the initial evaluation takes place also varies. The patient could seek evaluation in an outpatient setting with a generalist or a specialist or in the ED at a hospital. The recommendations in the present section are intended for consideration under the general principles of what constitutes GDMT during initial evaluation, regardless of the clinical setting. These general principles for the initial evaluation are shown in Figure 1. Additional evaluation is discussed in subsequent sections according to the outcomes of initial evaluation or in the presence of specific disease conditions.

Figure 1. Syncope Initial Evaluation



<sup>\*</sup>See relevant terms and definitions in Table 3.

Colors correspond to Class of Recommendation in Table 1. This figure shows the general principles for initial evaluation of all patients after an episode of syncope.

ECG indicates electrocardiogram.

## 2.3.1. History and Physical Examination: Recommendation

Recommen	Recommendation for History and Physical Examination				
COR	LOE	Recommendation			
I	B-NR	A detailed history and physical examination should be performed in patients with syncope (58-66).			
See Online Data Supplement 1.		The history should aim to identify the prognosis, diagnosis, reversible or ameliorable factors, comorbidities, medication use, and patient and family needs. Cardiac syncope carries a significantly worse prognosis than does neurally mediated syncope. Prognostic factors generally separate neurally mediated from cardiac syncope and are described in Section 2.3.3. The diagnostic history focuses on the situations in which syncope occurs, prodromal symptoms that provide physiological insight, patient's self-report, bystander observations of the event and vital signs, and post-event symptoms. Video recordings are helpful when available. Time relationship to meals and physical activities and duration of the prodrome are helpful in differentiating neurally mediated syncope from			
		cardiac syncope. Comorbidities and medication use are particularly important			

factors in older patients. A history of past medical conditions should be obtained, particularly with regard to the existence of preexisting cardiovascular disease (58-66). A family history should be obtained, with particular emphasis on histories of syncope or sudden unexplained death (or drowning). Historical characteristics associated with, though not diagnostic of, cardiac and noncardiac syncope are summarized in Table 4.

The physical examination should include determination of orthostatic blood pressure and heart rate changes in lying and sitting positions, on immediate standing, and after 3 minutes of upright posture (31). Careful attention should be paid to heart rate and rhythm, as well the presence of murmurs, gallops, or rubs that would indicate the presence of structural heart disease. A basic neurological examination should be performed, looking for focal defects or other abnormalities that would suggest need for further neurological evaluation or referral.

Table 4. Historical Characteristics Associated With Increased Probability of Cardiac and Noncardiac Causes of Syncope (60.67-75)

More Often Associated With Cardiac Causes of Syncope	
• Older age (>60 y)	Heart
• Male sex	Association
• Presence of known ischemic heart disease, structural heart disease, previous arrhythmias, or function	r reduced ventricular
• Brief prodrome, such as palpitations, or sudden loss of consciousness without prodrome	
Syncope during exertion	
Syncope in the supine position	
• Low number of syncope episodes (1 or 2)	
Abnormal cardiac examination	
• Family history of inheritable conditions or premature SCD (<50 y of age)	
Presence of known congenital heart disease	
More Often Associated With Noncardiac Causes of Syncope	
• Younger age	
No known cardiac disease	
Syncope only in the standing position	
Positional change from supine or sitting to standing	
Presence of prodrome: nausea, vomiting, feeling warmth	
• Presence of specific triggers: dehydration, pain, distressful stimulus, medical environment	
Situational triggers: cough, laugh, micturition, defecation, deglutition	
Frequent recurrence and prolonged history of syncope with similar characteristics	

SCD indicates sudden cardiac death.

# 2.3.2. Electrocardiography: Recommendation

Recommer	ndation for	Electrocardiography	
COR	LOE	Recommendation	
<b>T</b>	B-NR	In the initial evaluation of patients with syncope, a resting 12-lead	
1	B-NK	electrocardiogram (ECG) is useful (76).	
See Online Data Supplement 2.		ECG is widely available and inexpensive and can provide information about the	
		I potential and specific callse of the syncope episode (e.g., bradyarrhythmia with	
		sinus pauses or high-grade conduction block; ventricular tachyarrhythmia). It may	

demonstrate an underlying arrhythmogenic substrate for syncope or SCD. Subsets of patients with Wolff-Parkinson-White syndrome, Brugada syndrome, long-QT syndrome (LQTS), hypertrophic cardiomyopathy (HCM), or arrhythmogenic right ventricular cardiomyopathy (ARVC) have characteristic ECG features, which can prompt the decision to pursue further evaluation.

Despite the benefit of identifying a likely cause or potential clue about the cause of syncope from the ECG, prospective studies did not conclude that ECG findings significantly affected subsequent management (73,77-80). The prognostic value of an abnormal ECG in patients with syncope has been questioned, as well (69,81). However, a multicenter, prospective, observational study (76) concluded that the presence of AF, intraventricular conduction disturbances, voltage criteria for left ventricular (LV) hypertrophy, and ventricular pacing were associated with increased risk of death from all causes at 1 year.

#### 2.3.3. Risk Assessment: Recommendations

Syncope is a symptom that can be due to various causes, ranging from benign to life-threatening conditions. Risk stratification during initial evaluation is important for guiding the treatment and preventing long-term morbidity and mortality. However, risk stratification schemes for short- and long-term clinical outcomes are limited by the inclusion of all patients with syncope, without regard to the presence or absence of underlying medical conditions associated with syncope. For example, outcomes would not be expected to be similar for patients with vasovagal syncope (VVS), heart block with preserved ejection fraction, advanced cardiomyopathy and HF, acute gastric bleeding, or aortic dissection. The short-term prognosis of patients presenting with syncope is mainly related to the cause of syncope and the acute reversibility of the underlying condition; long-term prognosis is related to the effectiveness of therapy and the severity and progression of underlying diseases, especially cardiac or terminal illnesses.

Although having precise definitions for high-, intermediate-, and low-risk patient groups after an episode of syncope would be useful for managing these patients, evidence from current clinical studies renders this proposal challenging because of a large number of confounders. Risk markers from history, physical examination, laboratory investigations, study endpoints, adverse event rates, and time intervals between these events are variable from study to study. Current data are best grouped into short-term risk (associated with outcomes in the ED and up to 30 days after syncope) and long-term risk (up to 12 months of follow-up). Risk markers are summarized in Table 5 (64,67-70,72-75,82-98). The types of events, event rates, and study durations from investigations that estimated risk scores are summarized in Table 6 (64,65,76,81,87,89,92,97,99).

Recommendations for Risk Assessment			
COR	LOE	Recommendations	
I	B-NR	Evaluation of the cause and assessment for the short- and long-term morbidity and mortality risk of syncope are recommended (Table 5) (68,82,83,100).	

		Syncope may be an acute result of major hemodynamic abnormalities or a			
		manifestation of serious underlying disease. Thus, assessment of the cause of			
		syncope and underlying comorbidities is necessary.			
		Short-term adverse events and deaths are determined largely by the			
		cause of syncope and the effectiveness of the treatment. In patients without a			
See On	line Data	presumptive cause of syncope, risk stratification for potential short-term			
Supplem	nents 3 and	outcomes is necessary for immediate decision making in the acute setting.			
	4.	Potential predictors of increased short-term risk of death and serious outcomes			
		are listed in Table 5. Long-term adverse events and deaths are more likely			
		determined by the underlying medical comorbidities, many of which are			
		cardiac. The evaluation of patients with syncope should include a full			
		assessment of the long-term risk factors, including those listed in Table 5			
		(69,70,72-74,84-93,95,97).			
TTL	B-NR	Use of risk stratification scores may be reasonable in the management of			
IIb	B-NK	patients with syncope (67,68,72,73,75,87,89,100,101).			
		Investigators have reported numerous risk scores to predict adverse outcomes			
	after syncope (examples in Table 6). This literature has important limitations,				
	including inconsistent definitions of syncope, outcomes, outcome time frames,				
See On	See Online Data and predictors; inclusion of patients with serious outcomes already identified in				
Supplem	applements 3 and the ED, which biases risk scores toward identifying "obvious" events; the use of				
	4. composite outcomes that combine events with different pathophysiologies;				
		small samples that limited model reliability; and limited external validation.			
		Risk scores have not performed better than unstructured clinical judgment			
		(64,67-75,96,98).			

Table 5. Short- and Long-Term Risk Factors\*

Short-Term Risk Factors (≤30 d)	Long-Term Risk Factors (>30 d)				
History: Outpatient Clinic or ED Evaluation					
Male sex (74,85,101,102)	Male sex (68,90)				
Older age (>60 y) (88)	Older age (90)				
No prodrome (68)	Absence of nausea/vomiting preceding syncopal event (93)				
Palpitations preceding loss of consciousness (83)	VA (68,90)				
Exertional syncope (83)	Cancer (68)				
Structural heart disease (70,83,88,101,103)	Structural heart disease (68,103)				
HF (74,83,85,88)	HF (90)				
Cerebrovascular disease (70)	Cerebrovascular disease (68)				
Family history of SCD (70)	Diabetes mellitus (104)				
Trauma (68,101)	High CHADS-2 score (95)				
Physical Examination or Laboratory Investigat	ion				
	Abnormal ECG (84,90,93)				
Evidence of bleeding (83)	Lower GFR				
Persistent abnormal vital signs (70)					
Abnormal ECG (68,72,74,75,105)					
Positive troponin (75)					

<sup>\*</sup>Definitions for clinical endpoints or serious outcomes vary by study. The specific endpoints for the individual studies in this table are defined in Data Supplements 3 and 4 and summarized in Table 6 for selected studies. This table includes individual risk predictors from history, physical examination, and laboratory studies associated with adverse outcomes from selected studies.

CHADS-2 indicates congestive heart failure, hypertension, age ≥75 years, diabetes mellitus, and stroke or transient ischemic attack; ECG, electrocardiogram; ED, emergency department; GFR, glomerular filtration rate; HF, heart failure; SCD, sudden cardiac death; and VA, ventricular arrhythmias.

Table 6. Examples of Syncope Risk Scores

		Sample	Events	Outcome	ED		NPV
Study/Reference	Year	N	N	Definition	Events*	Predictors	(%)†
Martin (90)	1997	252	104 (41%)	1-y death/ arrhythmia	Yes	Abnormal ECG#; >45 y of age; VA; HF	93
Sarasin (74)	2003	175	30 (17%)	Inpatient arrhythmia	Yes	Abnormal ECG#; >65 y of age; HF	98
OESIL (67)	2003	270	31 (11%)	1-y death	N/A	Abnormal ECG#; >65 y of age; no prodrome; cardiac history	100
SFSR (72)	2004	684	79 (12%)	7-d serious events§	Yes	Abnormal ECG#; dyspnea; hematocrit; systolic BP <90 mm Hg; HF	99
Boston Syncope Rule (70)	2007	293	68 (23%)	30-d serious events‡	Yes	Symptoms of acute coronary syndrome; worrisome cardiac history; family history of SCD; VHD; signs of conduction earlies disease; volume depletion; persistent abnormal vital signs; primary central nervous event	100
Del Rosso (69)	2008	260	44 (17%)	Cardiac etiology	N/A	Abnormal ECG#/cardiac history; palpitations; exertional; supine; precipitant (low-risk factor); autonomic prodrome (low-risk factors)	99
STePS (68)	2008	676	41 (6%)	10-d serious events	Yes	Abnormal ECG#; trauma; no prodrome; male sex	
Syncope Risk Score (75)	2009	2,584	173 (7%)	30-d serious events¶	No	Abnormal ECG#; >90 y of age; male sex; positive troponin; history of arrhythmia; systolic BP >160 mm Hg; near-syncope (a low-risk factor)	97
ROSE (73)	2010	550	40 (7%)	30-d serious events¶	Yes	Abnormal ECG#; B-natriuretic peptide; hemoglobin; O <sub>2</sub> Sat; fecal occult blood	98

<sup>\*</sup>Did the study include events diagnosed during the ED evaluation?

AVB indicates atrioventricular block; BBB, bundle-branch block; BP, blood pressure; ECG, electrocardiogram; ED, emergency department; HF, heart failure; MI, myocardial infarction; N/A, not available; NPV, negative predictive value; O<sub>2</sub>Sat, oxygen saturation; OESIL, Osservatorio Epidemiologico sulla Sincope nel Lazio; ROSE, Risk Stratification of

<sup>†</sup>NPV: negative predictive value for lowest risk group for the specific events defined by the study.

<sup>‡</sup>Events: death, major therapeutic procedure, MI, arrhythmia, pulmonary embolism, stroke, sepsis, hemorrhage, or life-threatening sequelae of syncope.

<sup>§</sup>Events: death, MI, arrhythmia, pulmonary embolism, stroke, hemorrhage, or readmission.

Events: death, major therapeutic procedure, or readmission.

Events: death, arrhythmia, MI, new diagnosis of severe structural heart disease, pulmonary embolism, aortic dissection, stroke/TIA, cerebral hemorrhage, or significant anemia requiring blood transfusion.

<sup>#</sup>Abnormal ECG is defined variably in these studies. In the context of syncope evaluation, an abnormal ECG is any rhythm other than normal sinus rhythm, conduction delays (BBB, type-2 second-degree AVB or third-degree AVB), presence of Q waves, ST abnormalities, or prolonged OT interval.

Syncope in the ED; SCD, sudden cardiac death; SFSR, San Francisco Syncope Rule; STePS, Short-Term Prognosis of Syncope Study; TIA, transient ischemic attack; VA, ventricular arrhythmias; and VHD, valvular heart disease.

#### 2.3.4. Disposition After Initial Evaluation: Recommendations

The evaluating provider must decide whether further workup can continue in an outpatient setting or whether hospital-based evaluation is required. The purpose of hospital-based evaluation is to expedite the treatment of identified serious conditions or to continue the diagnostic evaluation in the absence of a presumptive cause of syncope (105,106).

The disposition decision is complicated by varying resources available for immediate testing, a lack of consensus on acceptable short-term risk of serious outcomes, varying availability and expertise of outpatient diagnostic clinics, and the lack of data demonstrating that hospital-based evaluation improves outcomes. In patients with a presumptive cause of reflex-mediated syncope and no other dangerous medical conditions identified, hospital-based evaluation is unlikely to provide benefit (35). In patients with perceived higher risk, the healthcare provider may recommend a hospital-based evaluation. In this setting, a structured ED protocol can be effective as an alternative to inpatient admission (107-110).

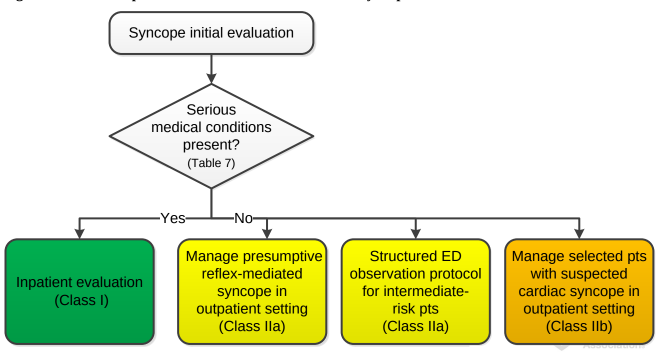
Decision support algorithms may reduce health service use in the evaluation of syncope (Figures 1 and 2) (105,111-113), although there are currently insufficient data to advocate the use of specific decision support algorithms for making disposition decisions.

Specialized syncope evaluation units may lead to reduced health service use and increased diagnostic rates (114-119). However, the logistical and financial feasibility of specialized syncope units in North American settings is unknown. A wider acceptance of syncope units requires further evidence of improvement in clinical outcomes. Individual risk factors (Table 5) and risk scores (Table 6) are correlated with short- and long-term clinical outcomes, but they are not primary determinants for admission to hospital. Presence of  $\geq 1$  serious medical condition, summarized in Table 7, is the key determinant for further in-hospital management of patients after syncope (90,98).

Recommendations for Disposition After Initial Evaluation			
COR	LOE	Recommendations	
I	B-NR Hospital evaluation and treatment are recommended for patients present with syncope who have a serious medical condition potentially relevant to the cause of syncope identified during initial evaluation (105,106,120).		
	ine Data nts 5 and 6.	Table 7 provides examples of serious conditions associated with syncope that may require inpatient evaluation and "treatment."  Arrhythmic causes may require consideration of pacemaker / implantable cardioverter-defibrillator (ICD) placement or revision and/or medication modification. Cardiac causes require treatment of the underlying condition (e.g., medication management and consideration of surgical intervention for critical aortic stenosis). Finally, a large spectrum of noncardiac serious conditions may be associated with syncope and require management of the underlying problem (e.g., severe anemia from a gastrointestinal bleed).	

IIa	C-LD	It is reasonable to manage patients with presumptive reflex-mediated syncope in the outpatient setting in the absence of serious medical conditions (35).
See Online Data Supplements 5 and 6.		Patients with presumptive VVS have a long-term risk of death similar to that of risk-matched patients without syncope (35). Hospital-based evaluation for presumptive VVS is unlikely to improve long-term outcomes. Possible exceptions that might require hospital-based evaluation include frequent recurrent syncope with risk of injury or identified injury related to syncope.
In intermediate-risk patients with an unclear cause of syncope, use of a structured ED observation protocol can be effective in reducing hospital		In intermediate-risk patients with an unclear cause of syncope, use of a structured ED observation protocol can be effective in reducing hospital admission (107-110).
See Online Data Supplements 5 and 6.		Two small RCTs suggest that structured ED-based protocols, consisting of time-limited observation and expedited access to cardiac testing/consultation, result in reduced health service use without adverse impact on clinical outcomes when compared with unstructured hospital admission. "Intermediate" risk factors included the following: ≥50 years of age; prior history of cardiac disease, cardiac device without evidence of dysfunction, concerning ECG findings, or family history of early SCD; and symptoms not consistent with reflex-mediated syncope. Both trials also allowed unstructured physician judgment to identify intermediate-risk patients (107-110).
IIb	It may be reasonable to manage selected patients with suspected cardiac	
See Online Data Supplements 5 and 6.		Hospital-based evaluation of syncope of unclear cause, in the absence of other serious identified medical conditions, has not demonstrated an improvement in patient-relevant outcomes. Several observational studies suggest modest diagnostic yield of hospital admission (121-123). Patients evaluated for suspected cardiac syncope in outpatient settings are seldom admitted for diagnostic purposes, and it may be reasonable to extend a similar approach to EDs after initial evaluation is completed in the ED. Primary providers can consider expedited referral to specialists with expertise in syncope, as indicated by availability of resources and provider's assessment of short-term risk of serious outcomes, as an alternative to extended hospital-based evaluation.

Figure 2. Patient Disposition After Initial Evaluation for Syncope



Colors correspond to Class of Recommendation in Table 1. ED indicates emergency department; pts, patients.

Table 7. Examples of Serious Medical Conditions That Might Warrant Consideration of Further Evaluation and Therapy in a Hospital Setting

Cardiac Arrhythmic Conditions	Cardiac or Vascular Nonarrhythmic Conditions	Noncardiac Conditions
<ul><li>Sustained or symptomatic VT</li><li>Symptomatic conduction system</li></ul>	Cardiac ischemia     Severe aortic stenosis	• Severe anemia/gastrointestinal
disease or Mobitz II or third-degree heart block  Symptomatic bradycardia or sinus pauses not related to neurally mediated syncope  Symptomatic SVT  Pacemaker/ICD malfunction  Inheritable cardiovascular conditions predisposing to arrhythmias	<ul> <li>Cardiac tamponade</li> <li>HCM</li> <li>Severe prosthetic valve dysfunction</li> <li>Pulmonary embolism</li> <li>Aortic dissection</li> <li>Acute HF</li> <li>Moderate-to-severe LV dysfunction</li> </ul>	bleeding  • Major traumatic injury due to syncope  • Persistent vital sign abnormalities

HCM indicates hypertrophic cardiomyopathy; HF, heart failure; ICD, implantable cardioverter-defibrillator; LV, left ventricular; SVT, supraventricular tachycardia; and VT, ventricular tachycardia.

## 3. Additional Evaluation and Diagnosis

The selection of a given diagnostic test, after the initial history, physical examination, and baseline ECG, is a clinical decision based on the patient's clinical presentation, risk stratification, and a clear understanding of diagnostic and prognostic value of any further testing. A broad-based use of additional testing is costly and often ineffective. This section provides recommendations for the most appropriate use of additional testing for syncope evaluation. See Figure 3 for the algorithm for additional evaluation and diagnosis for syncope.

Syncope additional evaluation and diagnosis Initial evaluation: history, physical exam, ECG (Class I) Initial evaluation Initial evaluation Stress testing clear unclear (Class IIa)† TTE Initial Initial Initial (Class IIa)† No additional Targeted blood evaluation evaluation evaluation evaluation testing Options suggests suggests reflex suggests CV needed\* (Class IIa)† **EPS** neurogenic OH syncope abnormalities (Class IIa)† Referral for Cardiac monitor MRI or CT Tilt-table autonomic selected based (Class IIb)† testing evaluation on frequency (Class IIa)† (Class IIa)† and nature (Class I) Options **Ambulatory Implantable** external cardiac cardiac monitor monitor (Class IIa)† (Class IIa)†

Figure 3. Additional Evaluation and Diagnosis for Syncope

Colors correspond to Class of Recommendation in Table 1.

CT indicates computed tomography; CV, cardiovascular; ECG, electrocardiogram; EPS, electrophysiological study; MRI, magnetic resonance imaging; OH, orthostatic hypotension; and TTE, transthoracic echocardiography.

## 3.1. Blood Testing: Recommendations

The availability of simple and accurate biomarkers might streamline risk stratification and diagnosis of the cause of syncope. This section reviews circulating biomarkers, which are being evaluated as markers either of hypotension or underlying disease processes. None have met with strong success.

Recommendations for Blood Testing			
COR	LOE	Recommendations	
IIa	B-NR	Targeted blood tests are reasonable in the evaluation of selected patients with syncope identified on the basis of clinical assessment from history, physical examination, and ECG (124).	
See Online Data Supplements 7 and 8.		Although broad-panel testing is common in clinical practice at the point of triage, there are no data on the utility of this approach. Data to support specific blood testing are largely descriptive data from case series and registries.  Complete blood count and electrolyte panel are frequently obtained during syncope evaluation. The diagnostic yield is low when these are used routinely;	

<sup>\*</sup>Applies to patients after a normal initial evaluation without significant injury or cardiovascular morbidities; patients followed up by primary care physician as needed.

<sup>†</sup>In selected patients (see Section 1.4).

however, when these blood tests are conducted in patients with a suspected
related diagnosis (e.g., history of peptic ulcer disease, or tarry stools associated
with OH on physical examination), test results can be diagnostic and useful for
guiding therapy. Thus, specific testing should stem from the assessment by
history and physical examination when the nature of the syncope presentation
or associated comorbidities suggests a diagnostic or more likely prognostic
role for laboratory testing. Results have not been linked to clinical decision
making or outcomes (125-128).
Usefulness of brain natriuretic peptide and high-sensitivity troponin
measurement is uncertain in patients for whom a cardiac cause of syncope
is suspected (125,127,129,130).
Although data to support biomarker testing are in general relatively weak,
there are sufficient data to suggest that natriuretic peptide is elevated in
patients whose subsequent cause for syncope is determined to be cardiac. A
systematic review of biomarkers found little value in contemporary troponin
measurement unless acute myocardial infarction is suspected, and there is
modest predictive value for high-sensitivity troponin and natriuretic peptides
for major adverse cardiovascular events. The ability of troponin and natriuretic
peptide measurement to influence clinical decision making or patient outcome
is unknown (129).
Routine and comprehensive laboratory testing is not useful in the
evaluation of patients with syncope (126,131).
There are no data on the utility of a standardized broad panel of laboratory
testing in patients with syncope. Specific cardiac biomarkers may play a
limited role when directed by clinical suspicion from the baseline assessment.
There is little biological plausibility linking the remaining elements of broad-
panel laboratory testing to the presentation or mechanism of syncope.

## 3.2. Cardiovascular Testing

Cardiovascular causes of syncope are common. The presence of significant cardiovascular diseases, often associated with the cardiovascular causes of syncope, portends a poor prognosis (35,132). As such, cardiovascular testing can be a critical element in the evaluation and management of selected patients with syncope. It is important also to recognize that the abnormalities found during cardiovascular testing may not have a causal relationship to syncope itself. Determining the significance of such abnormalities, their causality, and whether subsequent treatment is merited requires clinical judgment and appropriate selection of cardiovascular testing.

# 3.2.1. Cardiac Imaging: Recommendations

Recommendations for COR LOE		Recommendations	
COR	LUE		
IIa	B-NR	Transthoracic echocardiography can be useful in selected patients presenting with syncope if structural heart disease is suspected (80,99,124).	
See Online Data Supplement 9.		Cardiac imaging is often used to identify a structural cardiac abnormality, and imaging with transthoracic echocardiography is widely used for this purpose because it is noninvasive and low risk. Transthoracic echocardiography can be useful when healthcare providers are concerned about the presence of valvular disease (e.g., aortic stenosis), HCM, or LV dysfunction (124,133). In a retrospective study of patients presenting with syncope and suspected cardiac disease after history, physical examination, or ECG, the echocardiogram suggested a diagnosis of cardiac syncope in 48% of the study cohort (99). In a prospective evaluation of 650 patients referred for syncope of unknown origin, 88 patients had an abnormal history or ECG; an echocardiogram showed systolic dysfunction (LV ejection fraction ≤40%) in 24 patients (80); and 50% of patient with LV systolic dysfunction had manifest arrhythmias, compared with 9% with minor, incidental abnormalities (p<0.01). Although an echocardiogram may not be able to establish the immediate cause of syncope, it provides information for a potential disease substrate related to prognosis.	
IIb	B-NR	Computed tomography (CT) or magnetic resonance imaging (MRI) may be useful in selected patients presenting with syncope of suspected cardiac etiology (134).	
See Online Data Supplement 9.		Imaging modalities, including CT and MRI, are usually reserved for selected patients presenting with syncope, especially when other noninvasive means are inadequate or inconclusive. These modalities offer superior spatial resolution in delineating cardiovascular anatomy (e.g., in patients with structural, infiltrative, or congenital heart disease [CHD]) (135,136). The use of CT and MRI in contemporary cardiology is increasing (137,138). Their role in the evaluation of syncope has been investigated (139). The use of CT or MRI increased from 21% in 2001 to 45% in 2010, as reported in a series of patients evaluated for syncope in the ED (134). MRI is useful when there is a suspicion of ARVC or cardiac sarcoidosis (140,141). When pulmonary embolism is suspected in patients presenting with syncope to the hospital, CT can confirm the diagnosis in selected patients (128). CT or MRI may not provide answers about the cause of syncope. They provide information on the structural disease substrate relevant to the overall diagnosis and subsequent evaluation and follow-up in selected patients presenting with syncope.	
III: No Benefit	B-NR	Routine cardiac imaging is not useful in the evaluation of patients with syncope unless cardiac etiology is suspected on the basis of an initial evaluation, including history, physical examination, or ECG (77,99).	
See Online Data Supplement 9.		Although some investigators have advocated for cardiac imaging—particularly transthoracic echocardiography—as a routine screening examination for patients with syncope who lack clear signs or symptoms of cardiovascular disease (133), clinical evidence does not support such practice. Unexpected findings on	

echocardiograms to explain syncope are uncommon; a "screening" echocardiogram is of low utility (142). In 1 evaluation of 2,106 inpatients with syncope, a battery of testing, including cardiac enzymes, CT scans, echocardiography, carotid ultrasonography, and electroencephalography, contributed to the diagnosis or management in <5% of cases and helped determine the etiology of syncope <2% of the time (77). Similarly, in another retrospective series of 128 inpatients with syncope, it was found that echocardiograms in patients with no clinical evidence of heart disease according to history, physical examination, or ECG either were normal (63%) or provided no useful additional information for arriving at a diagnosis (37%) (99). Finally, radionuclide imaging and cardiac catheterization have little role in the evaluation of syncope.

#### 3.2.2. Stress Testing: Recommendation

Recommendation for Stress Testing			
COR	LOE	Recommendation	
IIa	IIa C-LD Exercise stress testing can be useful to establish the cause of syncope in selected patients who experience syncope or presyncope during exertion (132,143).		
	ine Data nent 10.	Exertion can result in syncope in a variety of conditions, including structural lesions, such as hypertrophic obstructive cardiomyopathy and aortic stenosis; interarterial anomalous coronary artery and pulmonary arterial hypertension; and channelopathies, such as LQTS (type 1) and catecholaminergic polymorphic ventricular tachycardia (CPVT). Subjecting a patient to a treadmill exercise test to reproduce the symptoms or evaluate the hemodynamic response to exertion (e.g., hypotension) must be done with extreme caution and in an environment with proper advanced life support.  In a prospective evaluation of 433 patients in which tachyarrhythmia was studied as the etiology for exertional syncope (132), an ECG stress evaluation was felt to be the sole test useful in identifying a presumptive cause of syncope in only 2 patients. However, bradyarrhythmia may ultimately be responsible for exertional syncope as well, and may only be elicited during stress testing. In rare instances, exercise-induced ischemia (143-146) or coronary vasospasm (147) may lead to high grade/infranodal AV block in patients with underlying coronary disease.	

## 3.2.3. Cardiac Monitoring: Recommendations

Although cardiac monitoring is often used in the evaluation of palpitations or intermittent arrhythmias, the following recommendations and discussion are focused primarily on the use of monitoring for the evaluation of patients with syncope. The choice of monitoring system and duration should be appropriate to the likelihood that a spontaneous event will be detected and the patient may be incapacitated and unable to voluntarily trigger the recording system.

Recommendations for Cardiac Monitoring		
COR	LOE	Recommendation
I	С-ЕО	The choice of a specific cardiac monitor should be determined on the basis of the frequency and nature of syncope events.
N/A		The technology of cardiac rhythm monitoring is dynamic and advancing at rapid speed. Several types of ambulatory cardiac rhythm monitoring are summarized in Table 8. Their selection and usefulness are highly dependent on patient characteristics with regard to the frequency of syncope and the likelihood of an
IIa B-NR		arrhythmic cause of syncope (148).  To evaluate selected ambulatory patients with syncope of suspected arrhythmic etiology, the following external cardiac monitoring approaches can be useful:  1. Holter monitor (149-153) 2. Transtelephonic monitor (150,154,155) 3. External loop recorder (150,154-156) 4. Patch recorder (157-159) 5. Mobile cardiac outpatient telemetry (160,161).
See Online Data Supplements 11 and 12.		The types of external monitoring devices are summarized in Table 8. The effectiveness of any external cardiac monitoring device for syncope evaluation is related to the duration of monitoring, continuous versus intermittent monitoring, frequency of syncope, duration of prodrome, and suddenness of incapacitation. The patient activation, before or after an event, allows for symptom rhythm correlation; however, some external loop recorders are of limited use in patients who are temporarily incapacitated around the time of syncope. External loop recorders are also limited by infrequent syncopal events. The advantage of an external loop recorder over Holter monitoring stems from a longer monitoring period, which confers a higher yield than Holter monitoring (149,153) and may offer a diagnosis after a negative Holter evaluation (150). Although the diagnostic yield of an external loop recorder may be lower than that of an implantable cardiac monitor (ICM), the noninvasive strategy is reasonable as a first approach. One prospective, multicenter study of 392 patients (28% with syncope) reported a 4-week diagnostic yield of 24.5%, with recurrent events and previous history of supraventricular arrhythmias being strong predictors of diagnostic events (156).  The advances of new patch-based devices offer another and often less cumbersome means of identifying an arrhythmic cause for syncope (157-159). The duration of monitoring (2 to 14 days) is often shorter than for the external loop recorder or mobile continuous outpatient telemetry.  Some practices offer mobile continuous outpatient telemetry devices, which provide real-time arrhythmia monitoring and analysis. An RCT (161) of 266 patients with suspected intermittent arrhythmias demonstrated that an arrhythmia was diagnosed in 88% of mobile continuous outpatient telemetry patients versus 75% of external loop recorder patients (p=0.008). Importantly, there was a similar result in the subgroup of patients presenting with syncope or presyncope, with a significantly higher di
IIa	B-R	outpatient telemetry group (89% versus 69%; p=0.008).  To evaluate selected ambulatory patients with syncope of suspected arrhythmic etiology, an ICM can be useful (149,150,153,161-175).

See Online Data Supplements 11 and 12. Several RCTs and observational studies have demonstrated a benefit of the ICM in establishing a diagnosis in syncope of unclear etiology. In a prospective study of 60 patients with syncope of unknown origin, the diagnosis (primarily bradyarrhythmia) was made in 55% with ICM, compared with a 19% diagnostic yield with conventional testing (external loop recorder, followed by tilt-table testing and electrophysiological study [EPS]) (p=0.0014) (162). These findings are consistent with other studies, which generally have shown that patients who underwent the ICM approach experienced higher rates of diagnosis than those of patients who underwent the conventional approach (164,176,177). A study on cost-effectiveness of the ICM strategy reported that the mean cost per participant was higher but the cost per diagnosis was lower in patients who received ICM than in patients who underwent conventional approaches (162,164,178). Key confounders in cost assessment include differences in healthcare settings, heterogeneity of patient populations, pricing of devices and healthcare delivery, and changing technology.

**Table 8. Cardiac Rhythm Monitors** 

Types of Monitor	Device Description	Patient Selection
Holter monitor (151-153)	<ul> <li>A portable, battery-operated device,</li> <li>Continuous recording for 24–72 h; up to 2 wk with newer models</li> <li>Symptom rhythm correlation can be achieved through a patient event diary and patient-activated annotations</li> </ul>	Symptoms frequent enough to be detected within a short period (24–72 h) of monitoring*
Patient-activated, transtelephonic monitor (event monitor) (150,154,155)	A recording device that transmits patient- activated data (live or stored) via an analog phone line to a central remote monitoring station (e.g., physician office)	<ul> <li>Frequent, spontaneous symptoms likely to recur within 2–6 wk</li> <li>Limited use in patients with frank syncope associated with sudden incapacitation</li> </ul>
External loop recorder (patient or auto triggered)† (150,154,155)	<ul> <li>A device that continuously records and stores rhythm data over weeks to months</li> <li>Patient activated, or auto triggered (e.g., to record asymptomatic arrhythmias) to provide a recording of events antecedent to (3–14 min), during, and after (1–4 min) the triggered event</li> <li>Newer models are equipped with a cellular phone, which transmits triggered data automatically over a wireless network to a remote monitoring system</li> </ul>	• Frequent, spontaneous symptoms related to syncope, likely to recur within 2–6 wk
External patch recorders (157-159)	<ul> <li>Patch device that continuously records and stores rhythm data, with patient-trigger capability to allow for symptom-rhythm correlation</li> <li>No leads or wires, and adhesive to chest wall/sternum</li> <li>Various models record from 2–14 d</li> <li>Offers accurate means of assessing burden of atrial fibrillation</li> <li>Patient activated, or auto triggered (e.g., to record asymptomatic arrhythmias) to provide a recording of events antecedent to, during, and after the triggered event</li> </ul>	<ul> <li>Can be considered as an alternative to external loop recorder</li> <li>Given that it is leadless, can be accurately self-applied, and is largely water resistant, it may be more comfortable and less cumbersome than an external loop recorder, potentially improving compliance</li> <li>Unlike Holter monitors and other external monitors, it offers only 1-lead recording</li> </ul>

Mobile cardiac outpatient telemetry (160,161)	<ul> <li>Device that records and transmits data (up to 30 d) from preprogrammed arrhythmias or patient activation to a communication hub at the patient's home</li> <li>Significant arrhythmias are detected; the monitor automatically transmits the patient's ECG data through a wireless network to the central monitoring station, which is attended by trained technicians 24 h/d</li> <li>This offers the potential for real-time, immediate feedback to a healthcare provider for evaluation</li> </ul>	<ul> <li>Spontaneous symptoms related to syncope and rhythm correlation</li> <li>In high-risk patients whose rhythm requires real-time monitoring</li> </ul>
Implantable cardiac monitor (162,167,179- 181)	<ul> <li>Subcutaneously implanted device, with a battery life of 2–3 y</li> <li>Triggered by the patient (or often family member witness) to store the event</li> <li>Models allow for transtelephonic transmission, as well as automatic detection of significant arrhythmias with remote monitoring</li> </ul>	Recurrent, infrequent, unexplained syncope (or suspected atypical reflex syncope) of suspected arrhythmic cause after a nondiagnostic initial workup, with or without structural heart disease

<sup>\*</sup>Includes history, physical examination, and 12-lead ECG; may include nondiagnostic tilt-table test or electrophysiological study.

## 3.2.4. In-Hospital Telemetry: Recommendation

Recommendation for In-Hospital Telemetry			
COR			
I	B-NR	Continuous ECG monitoring is useful for hospitalized patients admitted	
		Continuous ECG monitoring is useful for hospitalized patients admitted for syncope evaluation with suspected cardiac etiology (77,182,183).  Given that patients with syncope and structural heart disease are at high risk of death or significant arrhythmia (184), inpatient telemetry could be a valuable diagnostic modality. However, the diagnostic yield of inpatient telemetry is low in the absence of high suspicion about an arrhythmic cause (183). One study of 172 patients with syncope presenting to the ED and admitted to a telemetry unit revealed a diagnostic yield in 18% of patients, with 15% demonstrating bradyarrhythmias (182). The yield was highest in older patients with HF. No deaths occurred within an average monitoring time of 4.8±2.7 days. In 1 prospective study of 2,240 patients admitted to a telemetry unit, patients admitted for syncope (10%) had low rates of unexpected intensive care transfer, and most were unrelated to arrhythmic conditions (185). Furthermore, in another prospective evaluation of 205	
		patients admitted to telemetry, significant arrhythmias were seen in only 12 patients with known or suspected coronary artery disease or in those with previously documented arrhythmias (183). No arrhythmias or interventions occurred in the 7% of patients who were assigned to telemetry because of syncope. A large, prospective evaluation of 2,106 patients admitted with syncope demonstrated high telemetry use (95%) but a diagnostic yield of only 5% (77). Continuous telemetry in the hospital for patients presenting with syncope not suspected of a cardiac etiology is not cost-effective (186,187).	

<sup>†</sup>Higher yield in patients who are able to record a diary to correlate with possible arrhythmia. ECG indicates electrocardiogram.

#### 3.2.5. Electrophysiological Study: Recommendations

The EPS can identify a substrate for clinical bradyarrhythmia or tachyarrhythmia as a potential cause of syncope after a nondiagnostic initial evaluation. Despite these purported benefits, EPS has a limited role in the evaluation of syncope, especially in patients without known heart disease or with low suspicion of an arrhythmic etiology (117,187,188). The sensitivity and specificity of EPS to assess sinus node dysfunction and AV conduction disease in patients with syncope are variable, depending on patient selection and pretest probability of a bradycardia substrate (189-191).

Inducible ventricular tachycardia (VT) in patients with syncope, ischemic heart disease, and a prior history of myocardial infarction is predictive of spontaneous VT and prognosis. The causal relationship between the inducible VT during EPS and syncope requires clinical correlation. The lack of an inducible sustained monomorphic VT predicts lower risk of spontaneous VT and better prognosis (192). The overall role of EPS in the evaluation of ventricular arrhythmias (VA) in patients with syncope has diminished in the past 2 decades. This is primarily due to the use of ICD as a Class I indication for the primary prevention of SCD in patients with ischemic or nonischemic cardiomyopathy and significant LV dysfunction (ejection fraction ≤35%). An EPS is no longer required in patients with syncope before consideration of ICD therapy. However, although ICDs may reduce risk of death, they may not prevent syncope. The role of EPS in patients with syncope suspected to be due to VA and acquired nonischemic heart disease is unproven (193-198).

<b>Recommendations for</b>		·EPS	
COR	OR LOE Recommendations		
IIa	B-NR	EPS can be useful for evaluation of selected patients with syncope of suspected arrhythmic etiology (91,151,199-205).	
See Onli Supplen		Diagnostic results detected during EPS occur predominantly in patients who have cardiac disease (e.g., conduction system delay, coronary artery disease, cardiomyopathy, and valvular heart disease). Most of the literature evaluating EPS as a means to diagnose syncope is relatively old, and the data were obtained in referral centers where there was a high pretest probability of an arrhythmia. Eight of these small retrospective studies (91,199-205) (total n=625) found that, of the 406 patients with cardiac disease or an abnormal ECG, 41% had a positive result (of these, 21% had VT and 34% had a bradycardia) (151). Of 219 patients without evidence of heart disease, only 5% had a positive result (1% with VT and 10% with evidence of substrate for symptomatic bradycardia). Overall, the diagnostic yield of EPS was approximately 50% and 10% in patients with and without structural heart disease, respectively.	
B-NR EPS is not recommended for syncope evaluation in patients with a normal ECG and normal cardiac structure and function, unless an arrhythmic etiology is suspected (205-207).		· · · · · · · · · · · · · · · · · · ·	
See Online Data Supplement 14.		One prospective evaluation of 247 patients with syncope of undetermined etiology who underwent EPS found that the diagnostic yield was significantly higher in patients with an abnormal ECG than in those with a normal ECG (22% versus 3.7%) and that the diagnostic yield was low in patients with a normal	

ECG and without cardiac disease (2.6%) (206). In another small series of 34 patients with unexplained syncope who had normal ECGs and normal testing otherwise and who underwent EPS (205), the results were diagnostic in only 4 patients; the results were abnormal but not diagnostic in 2 patients and were normal in the remaining 28 patients. In another evaluation of 421 patients with undiagnosed syncope who underwent noninvasive testing as a means of predicting abnormal EPS findings, a normal ECG and ambulatory monitor were associated with a lower risk of EPS abnormalities than were an abnormal ECG and ambulatory monitor (9% versus 82%) (207).

## 3.2.6. Tilt-Table Testing: Recommendations

Recommendations for Tilt-Table Testing		
LOE	Recommendations	
B-R	If the diagnosis is unclear after initial evaluation, tilt-table testing can be useful for patients with suspected VVS (208-213).	
	Tilt-table testing has been used to evaluate patients with syncope for nearly 3 decades (208). It is an orthostatic stress test to assess the susceptibility of a vasovagal response to a postural change from a supine to an upright position. A positive response is defined as inducible presyncope or syncope associated with hypotension, with or without bradycardia (less commonly asystole). The hemodynamic response to the tilt maneuver determines whether there is a cardioinhibitory, vasodepressor, or mixed response (214). There is general consensus that a tilt-table angle of 70 degrees for 30 to 40 minutes would provide optimal yield (211,213,215). Adjunctive agents, such as a low dose of isoproterenol infusion or sublingual nitrates, may improve sensitivity but decrease specificity (210,212,216,217). A positive tilt-table test suggests a tendency or predisposition to VVS induced in the laboratory. This observation during tilt-table testing cannot necessarily define a causal etiology or be entirely conclusive of a reflex mechanism for syncope in the clinical setting. Correlation of tilt-table—induced findings to patients' clinical presentation is critically important to prevent consequences of false-positive results from tilt-table testing.  The utility of tilt-table testing is highest in patients with a suspected VVS when syncope is recurrent. Several factors have reduced the role of tilt-table testing in the evaluation of syncope: the overall moderate sensitivity, specificity, and reproducibility of tilt-table testing; the presence of false-positive response in controls; the increasing recognition of VVS from a structured history taking; and the availability of long-term cardiac monitoring (24,211,213).  Tilt-table testing can be useful for patients with syncope and suspected	
B-NR	delayed OH when initial evaluation is not diagnostic (218,219).	
	OH with standing, or a similar fall in blood pressure within 3 minutes of upright tilt-table testing to 60 degrees (220), is distinct from delayed OH, characterized by a sustained decrease in blood pressure occurring beyond 3 minutes of standing or upright tilt-table testing (220,221). Delayed OH may be responsible for syncopal episodes or symptoms of orthostatic intolerance only after prolonged standing. In 1 retrospective study of 230 patients with OH, only 46% had OH within 3 minutes of head-up tilt; 15% had OH between 3 and 10	
	B-R ine Data nent 15.	

		minutes; and 39% had OH only after 10 minutes of tilt-table testing (218). In
		10-year follow-up data from 165 of these patients, 54% of individuals with
		delayed OH progressed to classic OH (219). The 10-year death rate in
		individuals with delayed OH was 29%, compared with 64% and 9% in
		individuals with baseline OH and controls, respectively.
		Tilt-table testing is reasonable to distinguish convulsive syncope from
IIa	B-NR	epilepsy in selected patients (222-225).
See Online Data Supplement 15.		Convulsive syncope is a term that can be used to describe any form of syncope manifesting with convulsive movements (e.g., myoclonus). Prolonged convulsions and marked postictal confusion are uncommon in patients with syncope associated with convulsive movements (226), and fatigue is frequent after reflex syncope and may be confused with a postictal state (226). Tilt-table testing has been shown to be of value in this clinical setting when a detailed history cannot clearly determine whether the convulsive movements were secondary to syncope, given the need for objective evidence to help distinguish this entity from true epileptic seizures. In a prospective study of 15 patients with recurrent unexplained seizure-like episodes who were unresponsive to antiepileptic therapy (223), 67% had convulsive movements associated with can hypotension and bradycardia during tilt-table testing. In another study of 74 arr hypotension and bradycardia during tilt-table testing. In another study of 74 contain patients with a questionable diagnosis of epilepsy (because of drug-refractory seizures or clinically suspected not to be true epilepsy), a cardiac diagnosis was established in 42% of patients, with >25% developing profound hypotension or bradycardia during the head-up tilt-table test, confirming the diagnosis of VVS (225). Taken together, it can be estimated from these studies that approximately 50% of patients with either questionable or drug-refractory epilepsy have positive tilt-table tests suggestive of a vasovagal etiology (226).
IIa	B-NR	Tilt-table testing is reasonable to establish a diagnosis of pseudosyncope (227-229).
See Online Data Supplement 15.		Psychogenic pseudosyncope should be suspected when patients present with frequent (even daily) symptoms that mimic VVS (and, in some cases, with a history of true VVS). It is often challenging to differentiate psychogenic syncope from true syncope. However, tilt-table testing may help to elucidate the diagnosis. During tilt-table testing, the apparent unconsciousness with loss of motor control, combined with normal blood pressure and heart rate (and a normal electroencephalogram [EEG] if such a recording is obtained), rules out true syncope and most forms of epilepsy (227-229). In 1 study of 800 patients who underwent tilt-table testing, approximately 5% were diagnosed with pseudosyncope. Compared with patients with VVS, eye closure during the event, long periods of apparent transient loss of consciousness, and increased heart rate and blood pressure are highly specific for pseudosyncope. One study of 21 patients with suspected pseudosyncope who were subjected to tilt-table testing with continuous monitoring of the ECG, EEG, and blood pressure revealed 17 patients with non-epileptiform limb shaking without significant changes on an EEG or hemodynamic changes (227).
III: No Benefit	B-R	Tilt-table testing is not recommended to predict a response to medical treatments for VVS (230,231).

See Online Data	One of the purported advantages of tilt-table testing, in addition to suggesting a
	diagnosis of VVS, is the ability to assess the efficacy of pharmacological
	therapeutics in suppressing a vasovagal response to postural stress by evaluating
Supplement 15.	the effectiveness of a therapy during repeated testing (230,231). Several small
	studies suggested a possible benefit, but these data were limited by the lack of
	reproducibility of tilt-table testing (232-235).

# 3.3. Neurological Testing

#### 3.3.1. Autonomic Evaluation: Recommendation

Syncope due to neurogenic OH is common in patients with central or peripheral autonomic nervous system damage or dysfunction. Its causes should be sought so as to provide efficient, accurate, and effective management. Some symptoms of neurogenic OH may differ from those due to dehydration, drugs, and cardiac and reflex syncope; these include persistent and often progressive generalized weakness, fatigue, visual blurring, cognitive slowing, leg buckling, and the "coat hanger" headache (a triangular headache at the base of the neck due to trapezius ischemia). These symptoms may be provoked or exacerbated by exertion, prolonged standing, meals, or increased ambient temperature. Confirmation of specific neurogenic OH conditions causing syncope often requires additional autonomic evaluation.

Recommendation for Autonomic Evaluation			
COR	LOE	Recommendation	
IIa	C-LD	Referral for autonomic evaluation can be useful to improve diagnostic and prognostic accuracy in selected patients with syncope and known or suspected neurodegenerative disease (219,236-239).	
See On	line Data	The care of patients with neurogenic OH is complex, especially in individuals with neurodegenerative disease. Care providers must be knowledgeable in the pathophysiology of the autonomic nervous system and the pharmacology of treatments for neurodegenerative disease (33,240). Many symptomatic treatments for neurodegenerative disease will increase the risk of syncope due to worsening OH; selection of these treatments needs to be balanced against the increased morbidity of not treating the symptoms of the neurodegenerative disease. Such care may be provided by a neurologist, cardiologist, internist, or other physician who has sufficient training to treat these complicated patients.	
Supplement 16.		Syncope due to neurogenic OH is caused by either central or peripheral autonomic nervous system damage or dysfunction. Central autonomic degenerative disorders include multiple system atrophy (241), Parkinson's disease (242), and Lewy Body dementia (238). Peripheral autonomic dysfunction may be due to a selective degeneration of peripheral autonomic neurons, known as pure autonomic failure (243), or may accompany autonomic peripheral neuropathies, such as neuropathies due to diabetes amyloidosis, immunemediated neuropathies, hereditary sensory and autonomic neuropathies, and inflammatory neuropathies. Peripheral neuropathies due to vitamin B <sub>12</sub>	

deficiency, neurotoxic exposure, HIV and other infections, and porphyria are less common causes of neurogenic OH (240).

It can be useful to consider referring patients with the following characteristics for autonomic evaluation: Parkinsonism (241,244-246) or other central nervous system features (247,248), peripheral neuropathies (240), underlying diseases known to be associated with a peripheral neuropathy (240,248), progressive autonomic dysfunction without central or peripheral nervous system features (243,248), postprandial hypotension (248,249), and known or suspected neuropathic postural tachycardia syndrome (POTS)

(37,248,250). Autonomic evaluation may 1) determine the underlying cause of neurogenic OH, 2) provide prognostic information, and 3) have therapeutic

### 3.3.2. Neurological and Imaging Diagnostics: Recommendations

implications.

Many patients undergo extensive neurological investigation after an uncomplicated syncope event, despite the absence of neurological features on history or examination. A systematic review found that EEG, CT, MRI, and carotid ultrasound were ordered in 11% to 58% of patients with a presentation of syncope (78). The evidence suggests that routine neurological testing is of very limited value in the context of syncope evaluation and management; the diagnostic yield is low, with very high cost per diagnosis (36,77,78,251-260). The recommendations pertain to the use of these investigations in patients with syncope and not in patients in the wider category of transient loss of consciousness.

Recommendations for Neurological Diagnostics		
COR	LOE	Recommendations
IIa	C-LD	Simultaneous monitoring of an EEG and hemodynamic parameters during tilt-table testing can be useful to distinguish among syncope, pseudosyncope, and epilepsy (229,261-263).
See Online Data Supplement 16.		Although a thoughtful and detailed history usually suffices to distinguish among convulsive syncope, epileptic convulsions, and pseudosyncope, an EEG is particularly important when a diagnosis cannot be established after a thorough initial evaluation. ECG findings are characteristic if an episode can be induced during the tilt-table testing (261-263). Epileptiform discharges are recorded during epileptic convulsions whereas, during syncope, an EEG generally shows diffuse brainwave slowing with delta waves and a flat line pattern (263). Pseudosyncope and psychogenic nonepileptic seizures are associated with a normal EEG (229).
III: No Benefit	B-NR	MRI and CT of the head are not recommended in the routine evaluation of patients with syncope in the absence of focal neurological findings or head injury that support further evaluation (78,260).
See Online Data Supplement 16.		Syncope is due to global cerebral hypoperfusion, and brain structural abnormalities are rare. Nonetheless, MRI and CT are frequently used and infrequently helpful. In 5 studies investigating patients with syncope, MRI was used in 11% of 397 patients and established a diagnosis in only 0.24%. Similarly,

		in 10 studies of investigation of syncope, CT was used in 57% of 2,728 patients
		and established a diagnosis in only 1% (77,78,256,257,260). Given the cost and
		impact on health service facilities, MRI and CT should not be routinely used in
		the assessment of syncope. Neurological imaging may be indicated if significant
		head injury as a result of syncope is suspected. Although there is general concern
		about potential radiation-mediated harm from CT, there are very limited data on
		the actual harm from CT for syncope evaluation.
III: No		Carotid artery imaging is not recommended in the routine evaluation of
Benefit	B-NR	patients with syncope in the absence of focal neurological findings that support further evaluation (77,78,256,257,260).
		Syncope is due to global cerebral hypoperfusion and therefore not to unilateral
C O - 1'	D	ischemia. A review of 5 studies of carotid artery ultrasound and Doppler use in
See Onli		patients with syncope found that these modalities were used in 58% of 551
Supplen	ient 16.	patients and established a diagnosis in 0.5% (77,78,256,257,260). Carotid artery
		ultrasound should not be routinely used in the assessment of syncope.
III: No		Routine recording of an EEG is not recommended in the evaluation of
Benefit	B-NR	patients with syncope in the absence of specific neurological features suggestive of a seizure (36,77,254-258).
		EEGs are ordered frequently for the evaluation of syncope. A review of 7 studies
See Online Data Supplement 16.		of use of an EEG in patients with syncope found that it was used in 52% of 2,084
		patients and established a diagnosis in 0.7% (36,77,254-258). EEGs should not
		be routinely used in the assessment of syncope.

# 4. Management of Cardiovascular Conditions

The writing committee reviewed the evidence to support recommendations in the relevant ACC/AHA guidelines and affirms the ongoing validity of the related recommendations in the context of syncope, thus obviating the need to repeat existing guideline recommendations in the present guideline, except for the specific cardiac conditions in Sections 4.2.4, 4.2.5, and 4.3 for which ACC/AHA guidelines are not available. The relevant guidelines are noted in Table 2.

It is pertinent to note that the principles of evaluation and management of syncope in patients with various cardiac conditions are the same as for other noncardiac conditions. A thorough history, physical examination, and baseline ECG are recommended in all patients. The determination of the immediate cause of syncope may be related, indirectly related, or unrelated to the underlying cardiac condition. Management of patients with syncope and heart disease would include treating the immediate cause of syncope and further assessing long-term management strategies to improve prognosis. The recommendations stated in this section focus on syncope relevant to and within the context of the specific stated cardiac condition.

# 4.1. Arrhythmic Conditions

Cardiac arrhythmia is a common cause of syncope, and the prompt identification of an arrhythmic etiology has diagnostic and prognostic implications. When bradyarrhythmias and tachyarrhythmias are discovered in patients

with syncope, determining their causal relationship to syncope often poses challenges for the practitioner. The baseline presence of an arrhythmia does not necessarily represent the etiology of syncope (e.g., marked resting bradycardia in a young patient with syncope). Furthermore, determining the significance of atrial tachyarrhythmias and VT—which are often paroxysmal and occult on initial evaluation—poses additional challenges and may warrant a more extensive evaluation (Section 3.2). Section 4.1 broadly outlines strategies to guide the practitioner when evaluating patients with bradycardia, supraventricular arrhythmias (including AF), and VT.

## 4.1.1. Bradycardia: Recommendation

Recommendation for Bradycardia			
COR	LOE	Recommendation	
I	С-ЕО	In patients with syncope associated with bradycardia, GDMT is	
_	C-EO	recommended (12,264).	
		A search and review of papers on syncope and bradycardia has been performed	
		since the last guidelines were published in 2008 and 2012 (12,264). The writing	
		committee supports the previous recommendations pertaining to syncope in	
		patients with sinus node dysfunction and AV conduction diseases. In adult patients	
		presenting with syncope and chronic bifascicular block but without documented	
		high-degree AV block, for whom other causes have been excluded, an RCT (265)	
		showed that a dual-chamber pacemaker reduced recurrent syncope. The evidence	
		continues to support, without change from the previous recommendation, the	
N	/ <b>A</b>	notion that permanent pacemaker implantation is reasonable for syncope in	
IN IN	/A	patients with chronic bifascicular block when other causes have been excluded.	
		The use of adenosine triphosphate in the evaluation of syncope in older	
		patients continues to evolve. In a small, single-blind trial of older patients (mean	
		age 75 years) randomized to active pacing or back-up pacing with documented	
		adenosine triphosphate–sensitive sinoatrial or AV block, there was a 75% risk	
		reduction in syncope recurrence with dual-chamber pacing (266). Adenosine	
		triphosphate is not available in the United States. The writing committee has	
		reached a consensus not to make a new recommendation on its use for syncope	
		evaluation because of the limited data at this time.	

# 4.1.2. Supraventricular Tachycardia: Recommendation

Recommendations for Supraventricular Tachycardia (SVT)		
COR	LOE	Recommendation
I	C-EO	In patients with syncope and SVT, GDMT is recommended (10).
N	/A	Although patients with SVT frequently manifest palpitations and lightheadedness, syncope is uncommon. Of note, older patients with paroxysmal SVT are more prone to syncope or near-syncope than are younger patients; these symptoms appear to be independent of the rate of tachycardia, which is generally slower in older adult patients than in younger patients (267,268). Younger patients with SVT causing syncope generally have a very rapid tachycardia. Evaluation of syncope in patients with Wolff-Parkinson-White syndrome with preexcitation on ECG

		requires a thorough history to differentiate an arrhythmic syncope from a
		nonarrhythmic syncope, such as VVS, in younger patients (269). When a patient
!		
		with syncope reports antecedent palpitations and lightheadedness, VT should be
		more strongly suspected than SVT. EPS may be useful to distinguish a VT from an
		SVT responsible for syncope associated with these antecedent symptoms. It should
		be noted that palpitations can also precede vasovagal faints due to sinus
		tachycardia, so not all palpitations are necessarily due to paroxysmal SVT or VT.
Ι	С-ЕО	In patients with AF, GDMT is recommended (16).
		AF can be associated with syncope. As with other forms of SVT, syncope from a
		rapid ventricular response (in the absence of preexcitation) is relatively unusual.
		Patients with chronic AF merit control of the ventricular response or maintenance
	/ <b>A</b>	of sinus rhythm with appropriate antiarrhythmic therapy (in carefully selected
N/A		patients) (16). Patients with paroxysmal AF are predisposed to an abnormal neural
		response during both sinus rhythm and arrhythmia, and the onset of AF may
		trigger VVS (270). In patients with sinus node dysfunction, syncope could occur
		upon termination of AF when prolonged pauses are present.

# 4.1.3. Ventricular Arrhythmia: Recommendation



Recommendation for VA			
COR	LOE	Recommendation	
I	С-ЕО	In patients with syncope and VA, GDMT is recommended	
1	C-EO	(12,13,220,264,271).	
		Patients with VA (monomorphic or polymorphic) can present with syncope,	
		whether it is nonsustained or sustained. The mechanism of syncope from VA is	
		multifactorial, including: rapid rate, abrupt change in rate, abnormal atrial and	
		ventricular activation relationships, dyssynchrony of ventricular activation,	
		changes in autonomic tone, and body position during the VA (272). One study of	
N/A		113 patients with sustained VA showed that patients who had a mean VA rate of	
		≥200 bpm had a 65% incidence of syncope or near-syncope, compared with only	
		15% among patients with a rate <200 bpm (273). Of the patients with VA ≥200	
		bpm, 34% did not experience syncope or presyncope. The risk of recurrent	
		syncope and the overall long-term prognosis of patients with VA depend on the	
		severity of the underlying cardiac disease substrates. Indications for ICDs in	
		patients with syncope and suspected VA are predicated on the documentation of or	
		the risk of developing lethal VA (12).	

#### 4.2. Structural Conditions

Syncope occurs not infrequently in patients with underlying heart diseases. Comprehensive guidelines exist for diagnosis and management of many of these diseases, including sections on syncope. In this section, management of syncope is discussed in patients with underlying structural heart disease. The disease-specific ACC/AHA guidelines were assessed first, and then a comprehensive review of literature published since publication of these disease-specific guidelines was performed to ensure that prior recommendations about

syncope remained current. If new published data were available, they were incorporated into the present document.

### 4.2.1. Ischemic and Nonischemic Cardiomyopathy: Recommendation

Recommen	Recommendation for Ischemic and Nonischemic Cardiomyopathy		
COR	LOE	Recommendation	
I	С-ЕО	In patients with syncope associated with ischemic and nonischemic	
		cardiomyopathy, GDMT is recommended (12,13).	
		Evaluation of syncope in patients with ischemic and nonischemic cardiomyopathy	
		encompasses diagnosis and prognosis. Treatment of syncope is based on the	
		specific cause of syncope, whereas treatment for the underlying cardiomyopathy	
		impacts the long-term prognosis. A review of evidence supports previously	
N	/A	published recommendations for patients with syncope in the presence of	
1,	,	underlying cardiomyopathy. An ICD is recommended in patients with syncope of	
		undetermined origin with clinically relevant and significant VA induced at the time	
		of an EPS (28). ICD therapy is also reasonable for patients with unexplained	
		syncope and nonischemic dilated cardiomyopathy with significant LV dysfunction	
		(12,13,28).	

#### 4.2.2. Valvular Heart Disease: Recommendation

Recomme	Recommendation for Valvular Heart Disease		
COR	LOE	Recommendation	
I	С-ЕО	In patients with syncope associated with valvular heart disease, GDMT is recommended (11).	
		Patients with aortic stenosis may experience syncope during exertion. The	
		mechanism is often hemodynamic, as opposed to arrhythmic, because of inability	
		to augment and sustain cardiac output. In patients with valvular heart disease	
1	N/A	causing syncope, treatment is recommended by the latest guidelines (11).	
		Specifically, aortic valve replacement is recommended in patients with severe	
		aortic stenosis and syncope after other causes of syncope are also considered and	
		excluded.	

### 4.2.3. Hypertrophic Cardiomyopathy: Recommendation

Recommen	Recommendation for HCM				
COR	LOE	Recommendation			
I	С-ЕО	In patients with syncope associated with HCM, GDMT is recommended (20).			
N/A		A MEDLINE search and review of papers on syncope and HCM has been performed since the last guideline was published in 2011 (20). There are no new data that would alter the 2011 recommendations. Thus, the writing committee supports the previous recommendations pertaining to syncope in patients with HCM. Although there are no randomized trials, data from registries have shown consistently that unexplained syncope is an independent predictor for SCD and			

appropriate ICD discharges. The present writing committee concurs that ICD
implantation is reasonable in patients with HCM presenting with ≥1 recent
episodes of syncope suspected to be of arrhythmic nature.

## 4.2.4. Arrhythmogenic Right Ventricular Cardiomyopathy: Recommendation

Recommen	Recommendation for ARVC			
COR	LOE	Recommendations		
I	B-NR	ICD implantation is recommended in patients with ARVC who present with syncope and have a documented sustained VA (274-278).		
See Onli Supplen		ICD indications in patients with ARVC and sustained VA are no different than guidelines-based indications for secondary prevention of SCD in other diseases (12).		
IIa	B-NR	ICD implantation is reasonable in patients with ARVC who present with syncope of suspected arrhythmic etiology (274,275,277-279).		
See Online Data Supplement 17.		Unexplained or arrhythmic-appearing syncope in patients with ARVC has consistently been associated with increased risk of SCD or appropriate therapy after ICD implantation in multiple observational studies (274-279).		

#### 4.2.5. Cardiac Sarcoidosis: Recommendations

Recommer	dations for	Cardiac Sarcoidosis
COR	LOE	Recommendations
I	B-NR	ICD implantation is recommended in patients with cardiac sarcoidosis presenting with syncope and documented spontaneous sustained VA (12,280-286).
See Online Data Supplement 18.		ICD indications in patients with cardiac sarcoidosis and sustained VA are no different than guidelines- or consensus-based indications for secondary prevention of SCD (12,286). Macroreentry around the granulomas is the most common mechanism of VA in patients with cardiac sarcoidosis (280,281). Other mechanisms include triggered activity and abnormal automaticity due to myocardial inflammation (282). Unlike AV block, the results of immunosuppression in patients with VA are controversial. Some studies have shown improvement with immunosuppression (283), whereas others have shown no benefit and even harm due to worsening VA and aneurysm formation (284,285).
I	С-ЕО	In patients with cardiac sarcoidosis presenting with syncope and conduction abnormalities, GDMT is recommended (12,286-289).
See Online Data Supplement 18.		Patients with cardiac sarcoidosis and conduction abnormalities should be treated according to the most recent guidelines for cardiac pacing (12). Patients with cardiac sarcoidosis and conduction abnormalities have a worse prognosis than that of patients with idiopathic AV block (286,287). Immunosuppression can result in transient reversal of AV block; however, the reversibility is unpredictable (287-289). As such, it is recommended to proceed with pacing according the most recent guidelines regardless of AV block reversibility.
IIa	B-NR	ICD implantation is reasonable in patients with cardiac sarcoidosis and syncope of suspected arrhythmic origin, particularly with LV dysfunction or pacing indication (290-293).

Heart

See Online Data Supplement 18.		The presence of myocardial noncaseating granulomas and inflammation puts patients at risk of having both AV block and VA, particularly in the presence of LV dysfunction. Patients with cardiac sarcoidosis and mild-to-moderate LV dysfunction have a substantial risk of developing VA (290-293). In a multicenter study including 235 patients with cardiac sarcoidosis who received ICD therapy for primary or secondary prevention, including patients with syncope, 36% of patients received appropriate ICD therapy. Patients who received appropriate ICD therapies were more likely to be male and to have a history of syncope, lower LV ejection fraction, ventricular pacing on baseline ECG, and a secondary prevention indication than were those who did not receive appropriate ICD therapies (292). Therefore, given the presence of a substrate for VA in patients with cardiac sarcoidosis, ICD implantation is reasonable in patients presenting with syncope suspected to be of arrhythmic origin.
IIa	B-NR	EPS is reasonable in patients with cardiac sarcoidosis and syncope of suspected arrhythmic etiology (294).
See Online Data Supplement 18.		In patients with cardiac sarcoidosis, programmed electrical stimulation may help identify patients at risk of having VA. According to a study of 76 patients with cardiac sarcoidosis and no cardiac symptoms, 8 (11%) had inducible sustained VA. During a median follow-up of 5 years, 6 of 8 had VA or died, versus 1 of 68 in the noninducible group (294).

## 4.3. Inheritable Arrhythmic Conditions

The prevalence of inherited arrhythmic conditions is low, rendering the clinical significance of an abnormal test a challenge. Few syncope-specific studies exist. Most studies of patients with inherited arrhythmias are open label or not randomized and often are uncontrolled. Most of the publications included other cardiac events, such as cardiac arrest and death, either at enrollment or as an outcome. Syncope of suspected arrhythmic cause has been correlated with increased risk of SCD, cardiac arrest, or overall cardiac death. Although ICD is effective in aborting cardiac arrest and presumably reducing risk of death in the patients with inheritable rhythm disorders, its impact on syncope recurrence is unknown (25,26,220).

#### 4.3.1. Brugada Syndrome: Recommendations

Brugada syndrome is defined as a genetic disease characterized by an increased risk of SCD and ST elevation with type 1 morphology  $\ge 2$  mm in  $\ge 1$  lead among the right precordial leads V1 and V2, occurring either spontaneously or after intravenous administration of Class I antiarrhythmic drugs. The prevalence is higher in Asian countries than in North America or Western Europe, ranging from 0.01% to 1%, with a significant male predominance (295).

Recommendations for Brugada ECG Pattern and Syncope			
COR	COR LOE Recommendations		
IIa	B-NR	ICD implantation is reasonable in patients with Brugada ECG pattern and	
Па		syncope of suspected arrhythmic etiology (296-300).	

See Online Data Supplement 19.		Syncope is a risk factor for cardiac arrhythmic events in patients with Brugada syndrome (296,297). ICD implantation is reasonable in these patients; however, the benefit seems to be limited to patients with suspected arrhythmic syncope (298). Patients with syncope consistent with a reflex-mediated mechanism should not undergo the implantation of an ICD.  In a meta-analysis, the relative risk of cardiac events (SCD, syncope, or ICD shock) among patients with a history of syncope or SCD was approximately 3 times higher than among patients without a prior history of syncope or SCD (296). Data from an international registry showed that the cardiac event rate per year was 7.7% in patients with aborted SCD, 1.9% in patients with syncope, and 0.5% in asymptomatic patients (297). In a cohort including 203 patients with Brugada, VA occurred only in patients with syncope suspected to be arrhythmic in origin, at a rate of 5.5% per year. No SCD occurred in patients with nonarrhythmic syncope or with syncope of doubtful origin (298).
IIb	B-NR	Invasive EPS may be considered in patients with Brugada ECG pattern and syncope of suspected arrhythmic etiology (297,301,302).
See Online Data Supplement 19.		The value of EPS in assessing the mechanism of syncope in patients with Brugada is unknown. In large registries of patients with Brugada (PRELUDE and FINGER) (297,301), inducibility of VA was higher among patients with autor prior history of syncope or SCD. However, the value of EPS in predicting prognosis in patients with Brugada is essentially unknown in patients with syncope. The role of inducibility of VA in identifying high-risk patients remains controversial (301,302). Therefore, EPS may be considered only in patients with syncope suspected to be due to an arrhythmia and is not recommended in patients with reflex syncope.
III: No Benefit	B-NR	ICD implantation is not recommended in patients with Brugada ECG pattern and reflex-mediated syncope in the absence of other risk factors (303,304).
See Online Data Supplement 19.		In a retrospective multicenter study, appropriate ICD therapy was limited to survivors of cardiac arrest, whereas none of the other patients with syncope and/or inducible ventricular fibrillation (VF) suffered an arrhythmic event (303,304). Given the lack of benefit of ICD therapy in patients with reflex syncope and the known rate of inappropriate shocks and ICD complications in patients who receive an ICD (51), ICD implantation is not recommended when the syncope mechanism is believed to be reflex mediated.

## 4.3.2. Short-QT Syndrome: Recommendation

Short-QT syndrome is a genetic disease characterized by palpitations, syncope, and increased risk of SCD, associated with a QTc interval  $\leq$ 340 ms (25,26). It is a rare condition. Limited data are available about its prognostic significance, particularly in the absence of documented VA. Invasive EPS has shown increased vulnerability to VF induction in most patients, yet the clinical significance of this finding remains unknown (305). Quinidine therapy might provide some protection against VA; however, there are insufficient data to make any recommendations (305,306).

Recommer	Recommendation for Short-QT Syncope		
COR	LOE	Recommendation	
IIb	С-ЕО	ICD implantation may be considered in patients with short-QT pattern and	
110	C-EO	syncope of suspected arrhythmic etiology.	
See Online Data Supplement 20.		The prevalence of short-QT syndrome is very low, ranging from 0.02% to 1.63% (305,307-312). There is no evidence that syncope in patients with short-QT pattern is a risk factor for cardiac arrest in the absence of documented VT or VF. Therefore, ICD implantation may be limited to patients with suspected arrhythmic syncope, particularly in the presence of a family history of SCD (306).	

#### 4.3.3. Long-QT Syndrome: Recommendations

LQTS is diagnosed in the presence of QTc ≥500 ms or LQTS risk score ≥3.5 when secondary causes have been excluded or in the presence of a pathogenic mutation in 1 of the LQTS genes. It can also be diagnosed when the QTc is 480 to 499 ms in a patient presenting with syncope (25). There are several genetic forms of LQTS, which affect presentation and response to therapy. Given that syncope is often the result of an arrhythmic event in patients with LQTS, early recognition and treatment are needed to avoid recurrences, which could present as cardiac arrest or SCD. This is particularly true in the pediatric population, where significant overlap exists in the clinical presentation of patients with VVS and arrhythmic syncope (313,314). Attention to the triggers and presence of palpitations preceding syncope onset have been helpful in diagnosing an arrhythmic etiology (315).

Patients with LQTS and syncope should adhere to the lifestyle changes previously published, including avoidance of strenuous activity in LQT1, and drugs known to prolong QT interval in all patients with LQTS (25).

Recommen	Recommendations for LQTS		
COR	LOE	Recommendations	
I	B-NR	Beta-blocker therapy, in the absence of contraindications, is indicated as a first-line therapy in patients with LQTS and suspected arrhythmic syncope (316-318).	
See Online Data Supplement 21.		In the International Long QT Registry, patients who experienced ≥1 episode of syncope had a 6- to 12-fold increase in the risk of subsequent fatal/near-fatal events, independent of QTc duration. Beta-blocker therapy was associated with a significant reduction in the risk of recurrent syncope and subsequent fatal/near-fatal events. The response to beta blockers depends on the genotype, and not all beta blockers are the same (316,319). Patients with LQT1 appear to respond better than patients with LQTS2 and LQTS3 (316,320).	
IIa	B-NR	ICD implantation is reasonable in patients with LQTS and suspected arrhythmic syncope who are on beta-blocker therapy or are intolerant to beta-blocker therapy (317,320-324).	
See Online Data Supplement 21.		Cardiac events can occur in patients receiving beta-blocker therapy, with a prevalence ranging from 10% to 32%, depending on the genotype (316,317). Many patients who appear to not respond to beta blockers are poorly compliant or do not tolerate the medication (317). Therefore, ICD implantation is reasonable in patients with LQTS who continue to have syncope despite beta-blocker therapy and in those who cannot tolerate beta-blocker therapy. In a study	

		of 459 patients with genetically confirmed LQTS who received an ICD, syncope
1		was a predictor of appropriate therapy (322).
IIa	C-LD	Left cardiac sympathetic denervation (LCSD) is reasonable in patients with LQTS and recurrent syncope of suspected arrhythmic mechanism who are intolerant to beta-blocker therapy or for whom beta-blocker therapy has failed (325-327).
See Online Data Supplement 21.		LCSD has been shown to be associated with a large and significant clinical benefit in patients with symptomatic LQTS who are either refractory or intolerant to beta-blocker therapy (325,326). LCSD also reduces shocks in patients with an ICD during arrhythmia storms. Therefore, LCSD can be beneficial in patients with recurrent syncope despite beta blockade, in those who cannot tolerate beta-blocker therapy, and in those with frequent shocks from their ICD. However, LCSD alone does not completely prevent cardiac events, including SCD, during long-term follow-up.

#### 4.3.4. Catecholaminergic Polymorphic Ventricular Tachycardia: Recommendations

CPVT is characterized by the presence of catecholamine-induced (often exertional) bidirectional VT or polymorphic VT in the setting of a structurally normal heart and normal resting ECG (328,329). In patients with CPVT, 60% have a mutation in either the gene encoding the cardiac ryanodine receptor (*RyR2*) (autosomal dominant) or in the cardiac calsequestrin gene (*CASQ2*) (autosomal recessive) (330-333). The prevalence of the disease is estimated to be around 0.1 per 1,000 patients. Patients usually present in the first or second decade of life with stress-induced syncope (25).

Recommendations for CPVT		
COR	LOE	Recommendations
т	C-LD	Exercise restriction is recommended in patients with CPVT presenting with
1	C-LD	syncope of suspected arrhythmic etiology (328,334,335).
		The presence of VA in patients with CPVT has been shown to correlate with
See Onli	ine Data	increases in heart rate, highlighting the role of the sympathetic nervous system in
Supplemen	nts 22 and	arrhythmogenesis (328,334). Therefore, exercise restriction, including avoidance
23	3.	of heavy exercise and competitive sports, is recommended in all patients with
		CPVT (335).
т	C-LD	Beta blockers lacking intrinsic sympathomimetic activity are recommended
1	C-LD	in patients with CPVT and stress-induced syncope (329,334,336-339).
		Beta blockers should be first-line therapy in patients with CPVT, as they have
	_	been shown to suppress exercise-induced arrhythmias. However, they are not
See Onli		always completely protective (329,334,336). The variability in outcome with
Supplemer 23		beta-blocker therapy is due to multiple factors, including dosing and compliance
2.	J.	(337,338). Repeat exercise testing and cardiac monitoring to document
		arrhythmia suppression can be reassuring (334,339).
IIa	C-LD	Flecainide is reasonable in patients with CPVT who continue to have
Ha	C-LD	syncope of suspected VA despite beta-blocker therapy (319,320).
See Online Data		Despite beta-blocker therapy, breakthrough arrhythmias occur in patients with
Supplements 22 and 23.		CPVT because of treatment failure, noncompliance, and subtherapeutic dosing.
		The addition of flecainide to conventional therapy has been shown to partly or

		1,1
		completely suppress exercise-induced VA (340). In patients intolerant of beta-
		blocker therapy, flecainide is useful as monotherapy (341).
		ICD therapy is reasonable in patients with CPVT and a history of exercise-
IIa	B-NR	or stress-induced syncope despite use of optimal medical therapy or LCSD
		(271,342,343).
		ICD therapy appears to reduce mortality rate in patients with CPVT and syncope
		or VA refractory to medical therapy. However, VT storms in patients with CPVT
		may not always respond to ICD shocks (344), and shocks may precipitate early
		recurrence of arrhythmia because of their painful nature with resultant adrenergic
	ine Data	state. Furthermore, the effectiveness of ICD shock therapy in CPVT depends on
	nts 22 and	the mechanism of the VA, with greater success noted when shocks are delivered
2	3.	for VF (345). ICD implantation should be performed in conjunction with beta
		blocker therapy or LCSD when available (342). Careful programming, including
		long detection intervals with high cutoff rate, is recommended to decrease the
		prevalence of inappropriate shocks (342,343).
		In patients with CPVT who continue to experience syncope or VA,
IIb	C-LD	verapamil with or without beta-blocker therapy may be considered
22.0	0 22	(346,347). American
See Onl	ine Data	Verapamil alone or in combination with beta blockers helps suppress
Suppleme	nts 22 and	arrhythmias in patients with CPVT (347), including delaying the onset of
2	3.	exercise-induced ventricular ectopy (346,347).
TII.	C-LD	LCSD may be reasonable in patients with CPVT, syncope, and symptomatic
IIb	C-LD	VA despite optimal medical therapy (348-350).
See Online Data		When syncope occurs despite optimal medical therapy, LCSD may be a
		reasonable therapy (348-350). In a worldwide cohort study, the percentage of
Supplements 22 and 23.		patients with major cardiac events despite optimal medical therapy was reduced
		68% after LCSD (349).

#### 4.3.5. Early Repolarization Pattern: Recommendations

Early repolarization pattern is characterized by a distinct J point and ST elevation in the lateral or inferolateral leads. The pattern is more prevalent in young athletes, particularly African Americans, with 70% of the subjects being male (351). Early repolarization ECG pattern (>1 mm) in the inferior/lateral leads occurs in 1% to 13% of the general population and in 15% to 70% of idiopathic VF cases (352-354). Furthermore, it has been shown in population-based studies to be associated with increased risk of cardiac death (352,353,355-357). One study showed that the presence of a J wave increased the risk of VF from 3.4/100,000 to 11/100,000 (353). However, given the low incidence of VF in the general population, the absolute risk in patients with early repolarization remains low. In patients with syncope, the clinical significance of the early repolarization pattern is unknown.

Recommendations for Early Repolarization Pattern		
COR	LOE	Recommendations
IIb	С-ЕО	ICD implantation may be considered in patients with early repolarization pattern and suspected arrhythmic syncope in the presence of a family history of early repolarization pattern with cardiac arrest.

N/A	ICD implantation may be considered in patients with early repolarization pattern and suspected cardiac syncope if they have a family history of unexplained SCD, VF, or polymorphic VT with documented early repolarization pattern in the affected family member (358,359).
III: B-NR	EPS should not be performed in patients with early repolarization pattern
Harm	and history of syncope in the absence of other indications (359).
See Online Data Supplement 24.	In a multicenter study including 81 patients with early repolarization syndrome and aborted SCD who underwent EPS, VF was inducible in only 22% of cases. The VF recurrence rate was similar in patients who were inducible and in those who were noninducible (359). Given the high prevalence of early repolarization, the possibility of inducing VF in healthy individuals, and the limited value of ventricular programmed stimulation in risk stratification, EPS is not recommended in patients with early repolarization and syncope in the absence of other cardiac indications (352,353,360).

#### 5. Reflex Conditions



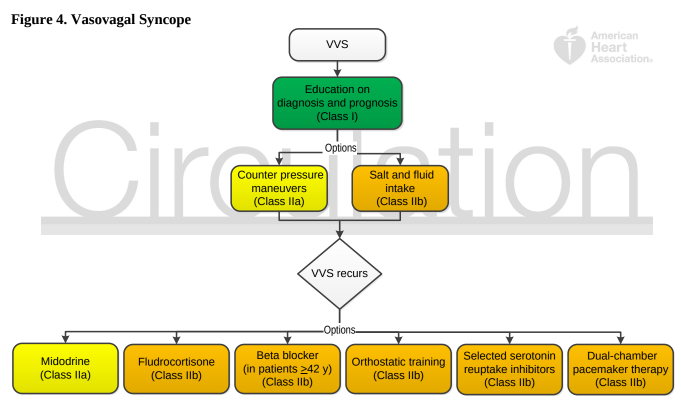
## **5.1. Vasovagal Syncope: Recommendations**

VVS is the most common cause of syncope and a frequent reason for ED visits (66). The underlying pathophysiology of VVS results from a reflex causing hypotension and bradycardia, triggered by prolonged standing or exposure to emotional stress, pain, or medical procedures (361-365). An episode of VVS is typically associated with a prodrome of diaphoresis, warmth, and pallor, with fatigue after the event. Given the benign nature of VVS and its frequent remissions, medical treatment is usually not required unless conservative measures are unsatisfactory. In some patients, effective treatment is needed, as syncopal events may result in injury and an impaired quality of life (QoL) (366-368). Despite the need and substantial efforts by investigators, there are limited evidence-based therapeutic options (369). Preliminary data from cardiac ganglia plexi ablation in treating selected patients with VVS are encouraging but still insufficient to make recommendations at this time (370-372). See Figure 4 for the algorithm for treatment of VVS.

Recommend	Recommendations for VVS		
COR	LOE	Recommendations	
I	С-ЕО	Patient education on the diagnosis and prognosis of VVS is recommended.	
See Onlin Supplement 26	its 25 and	In all patients with the common faint or VVS, an explanation of the diagnosis, education targeting awareness of and possible avoidance of triggers (e.g., prolonged standing, warm environments, coping with dental and medical settings), and reassurance about the benign nature of the condition should be provided.	
IIa	B-R	Physical counter-pressure maneuvers can be useful in patients with VVS who have a sufficiently long prodromal period (373-375).	

See Online Data Supplements 25 and 26.		Patients with a syncope prodrome should be instructed to assume a supine position to prevent a faint and minimize possible injury. In patients with a sufficiently long prodrome, physical counter-maneuvers (e.g., leg crossing, limb and/or abdominal contraction, squatting) are a core management strategy. In a randomized, parallel, openlabel trial, leg crossing with conventional therapy (i.e., fluid, salt intake, counseling, and avoidance) was superior to conventional therapy in preventing syncope recurrence (375).
IIa	B-R	Midodrine is reasonable in patients with recurrent VVS with no history of hypertension, HF, or urinary retention (376-380).
See Onlin Supplemen 26	nts 25 and	Midodrine is a prodrug that is metabolized to desglymidodrine, which is a peripherally active alpha-agonist used to ameliorate the reduction in peripheral sympathetic neural outflow responsible for venous pooling and vasodepression in VVS. Studies on the efficacy of midodrine support its use. In a meta-analysis of 5 RCTs in adults and children, midodrine was associated with a 43% reduction in syncope recurrence (318,376,378,379,381).
IIb	B-R	The usefulness of orthostatic training is uncertain in patients with frequent VVS (382-386).
See Onli Supplemen 26	nts 25 and	There are 2 main methods of orthostatic training. Patients undergo repetitive tilt-table tests in a monitored setting until a negative tilt-table test occurs and then are encouraged to stand quietly against a wall for 30 to 60 minutes daily, or patients simply standing quietly against a wall at home for a prolonged period of time daily. RCTs have not shown a sustained benefit in reducing episodes of syncope recurrence with either option (382,383,385,387).
IIb	B-R	Fludrocortisone might be reasonable for patients with recurrent VVS and inadequate response to salt and fluid intake, unless contraindicated (388,389).
See Online Data Supplements 25 and 26.		Fludrocortisone has mineralocorticoid activity resulting in sodium and water retention and potassium excretion, which results in increased blood volume. In a pediatric population, an RCT found more recurrent symptoms in the fludrocortisone arm than in the placebo arm (389). Serum potassium level should be monitored because of potential drug-induced hypokalemia. POST II (Prevention of Syncope Trial II) reported a marginally insignificant 31% risk reduction in adults with moderately frequent VVS, which was significant in patients after a 2-week dose stabilization period (388).
IIb	B-NR	Beta blockers might be reasonable in patients 42 years of age or older with recurrent VVS (390-393).
See Online Data Supplements 25 and 26.		RCTs on the efficacy and effectiveness of beta blockers for the prevention of syncope have been negative (64,390-393). However, in a meta-analysis of a prespecified, prestratified substudy of POST I and a large observational study, an age-dependent benefit of beta blockers among patients ≥42 years of age was found, compared with those of younger age (394,395).
IIb	C-LD	Encouraging increased salt and fluid intake may be reasonable in selected patients with VVS, unless contraindicated (396-399).
N/A		Evidence for the effectiveness of salt and fluid intake for patients with VVS is limited.  Nonetheless, in patients with recurrent VVS and no clear contraindication, such as a history of hypertension, renal disease, HF, or cardiac dysfunction, it may be reasonable to encourage ingestion of 2 to 3 L of fluid per day and a total of 6 to 9 g (100 to 150

		mmol) of salt per day, or about 1 to 2 heaping teaspoonfuls. The long-term balance of risks and benefits of a strategy of increasing salt and water intake is unknown.
IIb	C-LD	In selected patients with VVS, it may be reasonable to reduce or withdraw
11.0	C ZZ	medications that cause hypotension when appropriate (400).
		A careful examination of the patient's history for medications that may lower blood
		pressure (hypotensive agents) should be performed. Care should be taken to withdraw
N/	A	or reduce medications only where safe to do so and in conjunction with the prescribing
		healthcare provider.
	C-LD	In patients with recurrent VVS, a selective serotonin reuptake inhibitor might be
IIb		considered (393,401,402).
		Serotonin has central neurophysiological effects on blood pressure and heart rate and
See Online Data Supplements 25 and 26.		acutely induces syncope during tilt-table testing (403). Three small RCTs on selective
		serotonin reuptake inhibitors have been conducted on the effectiveness of fluoxetine
		and paroxetine in preventing syncope, with contradictory evidence of effectiveness
		(393,401,402).



Colors correspond to Class of Recommendation in Table 1. VVS indicates vasovagal syncope.

## 5.2. Pacemakers in Vasovagal Syncope: Recommendation

Pacemakers might seem to be an obvious therapy for VVS, given that bradycardia and asystole are present during some spells. Numerous observational studies and RCTs have assessed whether pacemakers are efficacious in preventing syncope (404-409). It is becoming clear that strict patient selection on the basis of documented

asystole during clinical syncope is important, and that observation combined with a tilt-table test that induces minimal or no vasodepressor response may increase the likelihood of a response to pacing. This is because a positive tilt-table test might identify patients who are likely to also have a vasodepressor response during VVS and therefore not respond as well to permanent pacing. As noted in Section 1.1, the recommendation in this section was based on a separately commissioned systematic review of the available evidence, the results of which were used to frame our decision making. Full details are provided in the ERC's systematic review report (9).

Recommendation for Pacemakers in VVS		
COR	LOE	Recommendation
IIb F	B-R <sup>SR</sup>	Dual-chamber pacing might be reasonable in a select population of patients 40 years of age or older with recurrent VVS and prolonged spontaneous pauses (404-408,410).
See Online Supplement and 28.	ts 27	Among patients with a positive tilt-table test, a benefit of pacing for treatment of recurrent syncope was evident as compared with medical or no therapy in open-label trials (52,404,406,410-412), but this result must be interpreted with caution because of the possibility of outcome ascertainment bias. In 2 RCTs, there was no statistically significant benefit seen with active pacing (407,408). However, in a select population of patients >40 years of age with recurrent syncope and documented spontaneous pauses $\geq$ 3 seconds correlated with syncope or an asymptomatic pause $\geq$ 6 seconds, dual-chamber pacing reduced syncope recurrence. There was less benefit in patients with a positive tilt-table test that induced a vasodepressor response (405).

SR indicates systematic review.

### 5.3. Carotid Sinus Syndrome: Recommendations

Carotid sinus syndrome is associated with mechanical manipulation of the carotid sinus, either spontaneously or with carotid sinus massage. It is diagnosed by the reproduction of clinical syncope during carotid sinus massage, with a cardioinhibitory response if asystole is >3 seconds or if there is AV block, or a significant vasodepressor response if there is ≥50 mm Hg drop in systolic blood pressure, or a mixed cardioinhibitory and vasodepressor response. It occurs more commonly in men >40 years of age (413,414) and is due to an abnormal reflex attributed to baroreceptor and possibly medulla dysfunction (415,416). Carotid sinus massage should be performed sequentially over the right and left carotid artery sinus in both the supine and upright positions for 5 seconds each, with continuous beat-to-beat heart rate monitoring and blood pressure measurement (417). Contraindications to performing carotid sinus massage include auscultation of carotid bruit and transient ischemic attack, stroke, or myocardial infarction within the prior 3 months, except if carotid Doppler excludes significant stenosis (418).

Recommendations for Carotid Sinus Syndrome			
COR	LOE	Recommendations	
IIa	B-R	Permanent cardiac pacing is reasonable in patients with carotid sinus syndrome that is cardioinhibitory or mixed (413,419-426).	

See Online Data		Syncope recurred in fewer patients treated with pacing than in untreated patients,
		with observation periods up to 5 years (420,423). In 3 controlled, open-label
Supplements 29–32.		trials, the relative risk reduction of syncope recurrence with pacemaker
		implantation was 76% (409,427-429). There are no large RCTs.
IIb	B-R	It may be reasonable to implant a dual-chamber pacemaker in patients with
110	D-K	carotid sinus syndrome who require permanent pacing (427-430).
		Evidence for dual-chamber pacing versus single-chamber pacing in carotid sinus
		hypersensitivity is limited to a few small RCTs and limited observational data
See On	line Data	(409,418,427-429). Although mixed, the data suggest dual-chamber pacing may
Supplements 29–32.		prevent hemodynamic compromise and improve symptom recurrence in older
		adults who may have concomitant sinus node dysfunction or conduction system
		disease.

#### **5.4. Other Reflex Conditions**

Situational syncope is defined as syncope occurring only in certain distinct and usually memorable circumstances, including micturition syncope, defecation syncope, cough syncope, laugh syncope, and swallow syncope (431-437). Appropriate investigations should be undertaken to determine an underlying etiology, including causes that may be reversible (431,433-436). Evidence for treatment is limited mainly to case reports, small case series, and small observational studies (431,433-436). Treatment of most types of situational syncope relies heavily on avoidance or elimination of a triggering event. This may not always be possible, so increased fluid and salt consumption and reduction or removal of hypotensive drugs and diuretics are encouraged where appropriate and safe (436).

## 6. Orthostatic Hypotension

#### 6.1. Neurogenic Orthostatic Hypotension: Recommendations

OH involves excessive pooling of blood volume in the splanchnic and leg circulations. With standing, venous return to the heart drops, with a resultant decrease in cardiac output (31). Normally, the autonomic nervous system provides compensatory changes in vascular tone, heart rate, and cardiac contractility. In some individuals, this response may be defective or inadequate (31). In neurogenic OH, the vasoconstrictor mechanisms of vascular tone may be inadequate because of neurodegenerative disorders, such as multiple system atrophy, pure autonomic failure, Parkinson's disease, and autonomic peripheral neuropathies, such as those due to diabetes mellitus and other systemic diseases (31). Neurogenic OH may present clinically as classic or delayed OH. Most commonly, OH is due to dehydration or medications, such as diuretics and vasodilators. Syncope caused by OH conditions occurs in the upright position. See Figure 5 for the algorithm for treatment of OH.

Recommendations for Neurogenic OH			
COR	LOE	Recommendations	

I	B-R	Acute water ingestion is recommended in patients with syncope caused by neurogenic OH for occasional, temporary relief (438,439).
See Online Data Supplements 33 and 34.		In neurogenic OH, acute water ingestion can temporarily restore orthostatic tolerance (438-444). The pressor effect of water is most likely sympathetically driven, with the peak effect occurring 30 minutes after ingestion of ≥240 mL and additional benefit seen with ≥480 mL (398,441,442). The presence of glucose or salt may reduce this effect by splanchnic vasodilatation or a decreased osmopressor response, respectively (397,439). Acute water ingestion for temporary relief of OH is not intended for routine or long-term use (24).
IIa	C-LD	Physical counter-pressure maneuvers can be beneficial in patients with neurogenic OH with syncope (374,445-450).
Suppleme	ine Data nts 33 and 4.	Isometric contraction, such as by leg crossing, lower body muscle tensing, and maximal force handgrip, can increase blood pressure, with the largest effect occurring with squatting versus other counter-pressure maneuvers (374,445-450). Leg crossing increases cardiac output in patients with neurogenic hypotension (447). Similar or larger benefit would be expected with squatting and other isometric contraction (449). The benefit is limited to patients with sufficient prodrome and the ability to perform these maneuvers adequately and safely (449).
IIa	C-LD	Compression garments can be beneficial in patients with syncope and OH (451-455).
Suppleme	ine Data nts 33 and 4.	In patients with OH, including older adult patients and those with neurogenic etiologies, compression garments can improve orthostatic symptoms and blunt associated decreases in blood pressure (451-456). The garments should be at least thigh high and preferably include the abdomen, as shorter garments have not been proved to be beneficial (457).
IIa	B-R	Midodrine can be beneficial in patients with syncope due to neurogenic OH (458-467).
	ine Data nts 33 and 4.	Midodrine improves symptoms of OH in patients with neurogenic OH (458-467). There is a dose-dependent effect, usually corresponding to an increase in standing blood pressure (459,460,462,463,466,467). Its use may be limited by supine hypertension, and other common side effects include scalp tingling, piloerection, and urinary retention (459,460,463,467).
IIa	B-R	Droxidopa can be beneficial in patients with syncope due to neurogenic OH (380,468-471).
See Online Data Supplements 33 and 34.		Droxidopa improves symptoms of neurogenic OH due to Parkinson disease, pure autonomic failure, and multiple system atrophy (380,468,470,471). Droxidopa might reduce falls, according to small studies (472). Use of carbidopa in patients with Parkinson disease may decrease the effectiveness of droxidopa (380). Use and titration of droxidopa may be limited by supine hypertension (380,469), headache, dizziness, and nausea (468,470-472).
IIa	C-LD	Fludrocortisone can be beneficial in patients with syncope due to neurogenic OH (473-476).
See Online Data Supplements 33 and 34.		Fludrocortisone increases plasma volume, with a resultant improvement in symptoms of OH (473,477,478). When taken regularly, fludrocortisone may prevent OH, at least in astronauts after space flight (476). Supine hypertension may be a limiting factor. When supine hypertension is present, other medications should be used before fludrocortisone. Other side effects commonly seen include

edema, hypokalemia, and headache, but more serious adverse reactions, such as adrenal suppression and immunosuppression, can also occur with doses >0.3 mg daily (479,480).  Encouraging increased salt and fluid intake may be reasonable in selected patients with neurogenic OH (396,398,441,443,444).  Although the data are limited for salt and fluid supplementation in patients with OH, these 2 treatments may improve blood pressure while decreasing symptoms from OH (396,398,439-444). Salt supplementation (e.g., 6 to 9 g [100 to 150 mmol; about 1 to 2 teaspoons] of salt per day) increases plasma volume, with limited benefit in patients with already high salt intake (396). Water ingestion increases the blood pressure via a pressor effect, most likely mediated by sympathetic activation, with a peak effect approximately 30 minutes after ingestion (398,439,441-443). This additional salt and fluid intake may not be beneficial in patients with history of hypertension, renal disease, HF, or cardiac dysfunction, and the long-term effects of these treatments, including the benefits and risks, is unknown.  Pyridostigmine may be beneficial in patients with syncope due to neurogenic OH who are refractory to other treatments (466,481,482).  In patients with autonomic failure and neurogenic OH, pyridostigmine is able to improve orthostatic tolerance through increases in peripheral vascular resistance
daily (479,480).  Encouraging increased salt and fluid intake may be reasonable in selected patients with neurogenic OH (396,398,441,443,444).  Although the data are limited for salt and fluid supplementation in patients with OH, these 2 treatments may improve blood pressure while decreasing symptoms from OH (396,398,439-444). Salt supplementation (e.g., 6 to 9 g [100 to 150 mmol; about 1 to 2 teaspoons] of salt per day) increases plasma volume, with limited benefit in patients with already high salt intake (396). Water ingestion increases the blood pressure via a pressor effect, most likely mediated by sympathetic activation, with a peak effect approximately 30 minutes after ingestion (398,439,441-443). This additional salt and fluid intake may not be beneficial in patients with history of hypertension, renal disease, HF, or cardiac dysfunction, and the long-term effects of these treatments, including the benefits and risks, is unknown.  Pyridostigmine may be beneficial in patients with syncope due to neurogenic OH who are refractory to other treatments (466,481,482).  In patients with autonomic failure and neurogenic OH, pyridostigmine is able to
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See Online Data Supplements 33 and 34.  limited benefit in patients with already high salt intake (396). Water ingestion increases the blood pressure via a pressor effect, most likely mediated by sympathetic activation, with a peak effect approximately 30 minutes after ingestion (398,439,441-443). This additional salt and fluid intake may not be beneficial in patients with history of hypertension, renal disease, HF, or cardiac dysfunction, and the long-term effects of these treatments, including the benefits and risks, is unknown.  Pyridostigmine may be beneficial in patients with syncope due to neurogenic OH who are refractory to other treatments (466,481,482).  In patients with autonomic failure and neurogenic OH, pyridostigmine is able to
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In patients with autonomic failure and neurogenic OH, pyridostigmine is able to
See Online Data   improve orthografic tolorence through increases in peripheral vescular registance
miprove orthostatic tolerance unrough increases in peripheral vascular resistance
Supplements 33 and and blood pressure (481,482). Side effects include nausea, vomiting, abdominal
cramping, sweating, salivation, and urinary incontinence (483).
Octreotide may be beneficial in patients with syncope and refractory
recurrent postprandial or neurogenic OH (484-487).
Splanchnic circulation pooling can contribute to OH, and this pooling can worsen  See Online Data  Splanchnic circulation pooling can contribute to OH, and this pooling can worsen  See Online Data
I III the posidiandial deriod (484-487). Octreolide reduces spiancinic blood flow by
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Supplements 33 and approximately 20% (486), which prevents postprandial hypotension, increases blood pressure, and improves orthostatic tolerance (484-487).

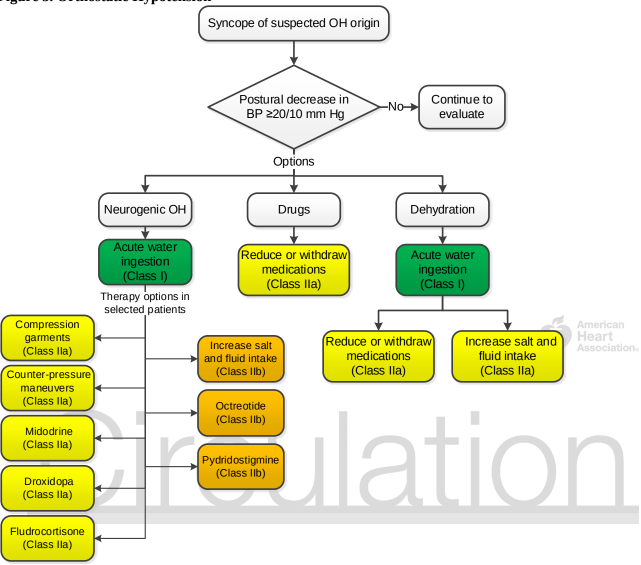
## **6.2. Dehydration and Drugs: Recommendations**

Syncope related to medication becomes prevalent particularly in older adults, who frequently have multiple comorbidities requiring treatment and are prone to polypharmacy effects (488-490). Cessation of offending medications is usually key for symptomatic improvement, but often feasibility of cessation of medications is limited by the necessity of the treatments (491-493). Dehydration may manifest along a spectrum of symptoms, ranging from tachycardia to shock, depending on whether a person has compensated or uncompensated hypovolemia (494). Orthostatic tolerance worsens with dehydration and is exacerbated by heat stress, which promotes vasodilation (495-497). Rehydration, whether by intravenous or oral formulation, should include sodium supplementation for more rapid recovery (21,498-501).

Recommendations for Dehydration and Drugs				
COR	LOE	Recommendations		
Т	C-LD	Fluid resuscitation via oral or intravenous bolus is recommended in		
1	C-LD	patients with syncope due to acute dehydration (438,499,501-504).		

	Fluid resuscitation is recommended for syncope secondary to both dehydration	
	and exercise-associated hypotension. The latter is likely due to peripheral	
	vasodilation and vasovagal physiology (438,495,504,505). Both dehydration	
ne Data	and heat stress worsen orthostatic tolerance (495-497). Oral fluid bolus may	
	require less volume than intravenous fluid infusion to have a similar treatment	
36.	effect because oral fluid loading has a pressor effect (398,438,440-444,502).	
	Beverages with increased sodium concentration (closer to normal body	
	osmolality) rehydrate faster than beverages with lower sodium concentration	
	or increased osmolality (e.g., because of glucose content) (498-501,503,506).	
	Reducing or withdrawing medications that may cause hypotension can be	
B-NR	beneficial in selected patients with syncope (488-490,492,507-510).	
	Syncope is a commonly reported adverse drug reaction, often resulting in	
	hospital admission (488,489). The prevalence of medication-related syncope	
appears higher in older patients (491,492,507,510). Several drug classes have		
na Data	been implicated in syncope, including diuretics, vasodilators, venodilators,	
	negative chronotropes, and sedatives (488-490,492,507-510). Close	
	supervision during adjustment of medications is frequently required because of	
30.	potential worsening of preexisting supine hypertension or cardiac arrhythmias	
	(491-493,511). Other factors to consider include frailty, HF and/or cardiac	
	dysfunction, and the use of a large number of medications causing adverse	
	effects because of drug-drug interactions (488,507,511-513).	
C-I D	In selected patients with syncope due to dehydration, it is reasonable to	
C-LD	encourage increased salt and fluid intake (396,498-501,503).	
	In patients with dehydration, sodium supplementation improves plasma	
	volume and improves orthostatic tolerance (396,499,503). This additional	
ne Data	dietary sodium may be provided as sodium tablets or sodium already dissolved	
	in beverages (396,498-500,503). Higher-sodium-content beverages with	
	osmolality comparable to normal body osmolality may rehydrate faster than	
	lower-sodium-content beverages (498-501,503). This treatment option is not	
	appropriate for patients with cardiac dysfunction or HF, uncontrolled	
	hypertension, or chronic kidney disease (19).	
	ne Data ents 35 36.  B-NR  ne Data ents 35 36.  C-LD  ne Data ents 35 36.	

Figure 5. Orthostatic Hypotension



Colors correspond to Class of Recommendation in Table 1. BP indicates blood pressure; OH, orthostatic hypotension.

#### 7. Orthostatic Intolerance

Orthostatic intolerance is a general term referring to frequent, recurrent, or persistent symptoms that develop upon standing (usually with a change in position from sitting or lying to an upright position) and are relieved by sitting or lying (38). Most commonly, the symptoms include lightheadedness, palpitations, tremulousness, generalized weakness, blurred vision, exercise intolerance, and fatigue. These symptoms may be accompanied by hemodynamic disturbances, including blood pressure decrease, which may or may not meet criteria for OH, and heart rate increase, which may be inadequate or compensatory (38). The pathophysiology is quite varied. One condition of note is POTS, in which upright posture results in an apparently inappropriate tachycardia, usually with heart rates >120 bpm (24).

Although syncope occurs in patients with POTS, it is relatively infrequent, and there is little evidence that the syncope is due to POTS (24,514). Treatments that improve symptoms of POTS might decrease the occurrence of syncope, although this is unknown (24,514-523). For further guidance on the management of POTS, we refer readers to the HRS consensus statement (24).

### 8. Pseudosyncope: Recommendations

Psychogenic pseudosyncope is a syndrome of apparent loss of consciousness occurring in the absence of impaired cerebral perfusion or function. Psychogenic pseudosyncope is believed to be a conversion disorder—in essence, an external somatic manifestation or response to internal psychological stresses. It is an involuntary response and should not be confused with malingering or Munchausen syndrome. Psychogenic pseudosyncope and pseudoseizures may be the same condition. The clinical distinction between the 2 is based on whether prominent jerky muscle movements simulating seizure activity are reported by witnesses. In the absence of associated jerky movements, the patient is likely to be referred for evaluation of syncope (30,229,524).

Psychogenic pseudosyncope does not result in a true loss of consciousness, but it is included in the present document because patients appear to exhibit syncope and therefore are referred for evaluation of syncope.

Several key clinical features are suggestive of the diagnosis of psychogenic pseudosyncope. None alone, however, provides a definitive diagnosis. Patients with psychogenic pseudosyncope are often young females with a higher prevalence of preexisting VVS or a history of physical and/or sexual abuse (229,525). The apparent duration of loss of consciousness is often long (5 to 20 minutes), and episodes are frequent (525). Some common characteristics include closed eyes, lack of pallor and diaphoresis, and usually little physical harm (526). A normal pulse, blood pressure, or EEG during a psychogenic pseudosyncope episode can be documented (229). Although many patients with pseudosyncope can be diagnosed with a careful history, occasionally tilt-table testing with or without transcranial Doppler and monitoring of an EEG is helpful.

Recommen	Recommendations for the Treatment of Pseudosyncope		
COR LOE Recommendations		Recommendations	
IIb	C-LD	In patients with suspected pseudosyncope, a candid discussion with the patient about the diagnosis may be reasonable (30,527-529).	
See Online Data		Some reports suggest that patients benefit from being informed of the suspected	
Supplements 37 and 38.		diagnosis in a clear but sympathetic manner that also acknowledges the	
		involuntary nature of the attacks (30,527,528).	
IIb C-LD Cognitive behavioral therapy may be beneficial in patients with pseudosyncope (530-532).			
See Online Data Supplements 37 and		Uncontrolled studies suggest that psychotherapy, particularly cognitive	
		behavioral therapy, may be beneficial in conversion disorders (530-532). One	
		RCT reported that cognitive behavioral therapy provided a non-statistically	
3	8.	significant trend toward improvement in pseudosyncope at 3 months (530).	
		There are no data that support significant benefit from pharmacotherapy (529).	

### 9. Uncommon Conditions Associated With Syncope

Syncope has been reported in many uncommon diseases, according to case reports. However, specific conditions may predispose the patient to various types of syncope. Table 9 provides a list of less common conditions associated with syncope. It is not intended as a reference for differential diagnosis or a complete synopsis of all conditions associated with syncope. Furthermore, it is not necessary to fully evaluate for all these causes when the etiology remains elusive. Most of these presentations rarely cause syncope, and data are sparse. If the cause for syncope is unclear, these conditions could be included in the differential diagnosis on the basis of other clinical characteristics and/or historical features.

**Table 9. Conditions Uncommonly Associated With Syncope** 

Condition	Clinical Characteristics	Notes
Cardiovascular and Cardio	pulmonary	
Cardiac tamponade	Hypotension, tachycardia, cardiogenic shock.	Often tachycardia and hypotension; may be hypotensive and bradycardic acutely.
Constrictive pericarditis (533-535)	Severe HF symptoms, including edema, exertional dyspnea, orthopnea.	May be associated with cough syncope.
LV noncompaction (536-539)	Cardiomyopathy characterized by prominent LV trabeculae and deep intertrabecular recesses, due to embryologic perturbation.	Syncope reported in 5%–9% of both adult and pediatric patients. The mechanism may be a tachyarrhythmia.
Takotsubo cardiomyopathy (540,541)	Apical ballooning and basal hypercontractility, often due to stress. Chest pain and ECG changes consistent with ischemia are commonly seen.	Syncope is uncommon and may be multifactorial.
Pulmonary embolus (128,542,543)	Hypoxemia, tachycardia; hypotension and shock leading to pulseless electrical activity cardiac arrest in severe cases.	Syncope due to bradycardia and/or hypotension.
		One study showed higher prevalence of pulmonary embolus in older patients with first episode of syncope after admission to the hospital. Further confirmation of this finding in the older populations is warranted.
Pulmonary arterial hypertension	Occurs more often during exertion in younger patients.	Syncope due to inability to augment or sustain cardiac output during exertion, followed by vasodilatation.
Infiltrative		
Fabry's disease (544,545)	Lysosomal storage disorder with neuropathic pain, renal failure concentric LVH, and HF.	Syncope usually due to AV block.
Amyloidosis (546,547)	Systemic disease due to amyloid deposition. Light chain amyloidosis affects the kidneys, heart, and peripheral and autonomic nervous systems.	Syncope may be due to conduction system disease, arrhythmias, impaired cardiac output from restrictive cardiomyopathy, or neurological involvement. AV block is the likely cause, although VA may occur with myocardial involvement.
Hemochromatosis (548)	Systemic iron deposition causing liver disease, skin pigmentation, diabetes mellitus, arthropathy, impotence, and dilated cardiomyopathy.	Myocardial involvement more common than sick sinus syndrome and AV conduction disease.

Infectious		
Myocarditis	Chest pain, arrhythmias, or profound LV	VT and AV block are the likely causes
(413,549-553)	systolic dysfunction. Hemodynamic	of syncope; transient hemodynamic
,	collapse may occur.	collapse is possible.
Lyme disease	Lyme myocarditis with classical features	Syncope may be due to AV block, but
(554)	of Lyme disease, including erythema	many patients manifest VVS (554,555).
(334)	migraines and neurological manifestations.	many patients mannest v v 5 (554,555).
Cl 1'		C
Chagas disease	Chagasic cardiomyopathy caused by	Syncope and sudden death associated
(556-559)	trypanosomiasis.	with ventricular tachyarrhythmias. AV
		block also occurs.
Neuromuscular		
Myotonic dystrophy	Autosomal dominant inheritance with	Both bradyarrhythmia and
(12,560,561)	multiple organ systems affected. Grip	tachyarrhythmias.
	myotonia, weakness, temporal wasting,	
	alopecia, cataracts, glucose intolerance,	
	and daytime somnolence.	
Friedreich ataxia	Autosomal recessive inheritance with limb	Syncope can be bradycardic or
(562,563)	and gait ataxia, bladder dysfunction, and	tachycardic. SCD is known to occur.
(202,203)	daytime somnolence. Diffuse interstitial	tacifycaraic. See is known to occur.
	fibrosis and HCM.	
Vacama Carma		Many patients develop significant His
Kearns Sayre	Mitochondrial myopathy. Chronic	
(564,565)	progressive external ophthalmoplegia;	Purkinje disease.
	pigmentary retinopathy.	Heart
Erb dystrophy	Limb girdle muscular dystrophy,	AV conduction disease, dilated
(566)	manifesting as scapulohymeral and/or	cardiomyopathy.
	pelvifemoral weakness and atrophy.	
Anatomic		
Lev's and Lenegre's	Progressive fibrosis and sclerosis of	Syncope is usually due to high-grade
diseases	cardiac conduction system, including the	AV block.
(567-571)	cardiac skeleton, including the aortic and	
(88, 8, 1)	mitral rings.	1 1 1 / 1 / 1
Cardiac tumors	Triad of obstruction, embolic, and	Syncope is often due to obstruction to
(572)	systemic signs and symptoms.	blood flow.
Prosthetic valve thrombosis	Ranges from asymptomatic to profound	May have similar presentation to a
	• • • • • • • • • • • • • • • • • • • •	
(573-575)	HF.	cardiac tumor, with a high risk of
		embolic phenomenon and obstruction.
Anomalous coronary artery	Common cause of exertional syncope or	Syncope can be due to Bezold Jarisch
(576-579)	SCD, classically in young athletes.	reflex, hypotension, VT, or AV block.
Aortic dissection	Aortic dissection may manifest with	The risk of in-hospital death, tamponade
(580-582)	neurological symptoms, myocardial	and neurological deficits is higher in
	infarction, and HF. Syncope can occur in	patients with syncope. Otherwise,
	as many as 13% of aortic dissections.	syncope alone does not appear to
		increase the risk of death.
Subclavian steal	The phenomenon of flow reversal in a	Syncope is generally associated with
(583-587)	vertebral artery ipsilateral to a	upper-extremity activity.
(303 201)	hemodynamically significant stenosis of	apper extremity detivity.
	the subclavian artery. Severe cases	
	resulting in vertebrobasilar ischemia may	
G	rarely result in syncope.	A
Coarctation of the aorta	If severe, it can result in HF or aortic	Associated bicuspid aortic valve stenosis
(588)	dissection.	may be considered with syncope.
Rheumatoid arthritis	Chronic, autoimmune inflammatory	Rarely associated with complete heart
(589)	disorder with systemic manifestations.	block and syncope.
	Arnold Chiari malformations are the most	Syringomyelia-induced disruption of
Syringomyelia	Arnold Chiari manormations are the most	Syllingolliy olla lilaacea alsraptioli ol
Syringomyelia (590-597) Chiari malformation	common form of syringomyelia.	sympathetic fibers in the thoracic spinal cord is a rare mechanism of syncope

Neck/vagal tumor (600,601)	Recurrent syncope is an uncommon complication of neck malignancy.	The mechanism may be invasion of the carotid sinus or the afferent nerve fibers of the glossopharyngeal nerve.
Endocrine		
Carcinoid syndrome (602) Pheochromocytoma (602,603) Mastocytosis (602-609) Vasoactive intestinal peptide tumor	These tumors can release vasoactive peptides and cause vasodilation, flushing, pruritus, and gastrointestinal symptoms.	Syncope is usually due to transient hypotension.
Hematologic		
Beta thalassemia major (610)	Severe anemia, multiple organ failure, and dilated cardiomyopathy due to iron overload.	Syncope may be arrhythmic.
Neurological disorders		
Seizure induced bradycardia/hypotension (611-614)	Generally due to temporal lobe epilepsy.	Postictal bradyarrhythmia is uncommon and likely originates from the temporal lobe or limbic system.
Migraine (615,616)	Migraine headaches are statistically associated with syncope.	Syncope may be vasovagal or due to orthostatic intolerance.

ACC indicates American College of Cardiology; AHA, American Heart Association; AV, atrioventricular; ECG, electrocardiogram; HCM, hypertrophic cardiomyopathy; HF, heart failure; HRS, Heart Rhythm Society; LV, left can ventricular; LVH, left ventricular hypertrophy; SCD, sudden cardiac death; VA, ventricular arrhythmias; VT, ventricular tachycardia; and VVS, vasovagal syncope.

### 10. Age, Lifestyle, and Special Populations

## 10.1. Pediatric Syncope: Recommendations

Syncope is common in the pediatric population. By 18 years of age, it is estimated that 30% to 50% of children experience at least 1 fainting episode, and syncope accounts for 3% of all pediatric ED visits (617-622). The incidence is higher in females and peaks between 15 to 19 years of age (617). Neurally mediated syncope accounts for 75% of pediatric syncope, followed by psychogenic or unexplained syncope in 8% to 15% of cases (623). Breath-holding spells are a form of syncope unique to the pediatric population. Cyanotic breath-holding spells typically occur from age 6 months to age 5 years and may be due to desaturation caused by forced expiration during crying. Pallid breath-holding spells are seen in the first 1 to 2 years of age and may be an early form of VVS. The latter episodes are associated with significant bradycardia and prolonged asystole. Pediatric cardiac syncope may result from obstruction to blood flow (HCM, aortic stenosis, pulmonary hypertension), myocardial dysfunction (myocarditis, cardiomyopathy, congenital coronary anomaly, or post–Kawasaki disease) or a primary arrhythmic etiology (LQTS, CPVT, Brugada syndrome, ARVC, or Wolff-Parkinson-White syndrome).

A detailed history with careful attention to the events leading up to the syncope and a complete physical examination can guide practitioners in differentiating the life-threatening causes of syncope (with potential for injury or SCD) from the more common and benign neurally mediated syncope. A detailed family history, with particular attention to premature SCD among first- and second-degree relatives and the manner in which those

deaths occurred, is helpful. Given that many of the causes of non-CHD cardiac syncope in children who do not have a form of CHD are similar to those experienced in an adult cohort (LQTS, HCM, Wolff-Parkinson-White, Brugada, and ARVC), interventions recommended for adults with similar conditions presenting with syncope can be applied in children.

Recommendations for Pediatric Syncope		
COR	LOE	Recommendations
I	C-LD	VVS evaluation, including a detailed medical history, physical examination, family history, and a 12-lead ECG, should be performed in all pediatric patients presenting with syncope (315,618,620,624-630).
See Online Data Supplement 40.		Although VVS is the most common cause of pediatric syncope, cardiac syncope does represent 1.5% to 6% of pediatric cases (usually defined as up to 18 years of age) (617,619,620,629,631,632). Characteristics of presenting signs and symptoms differentiating VVS from cardiac causes of syncope are generally similar to those in adults. A family history of VVS and early SCD should be sought. VVS occurs in 33% to 80% of children with syncope (624,628). Risk factors that raise suspicion of a cardiac etiology include the absence of prodromal symptoms, presence of preceding palpitations within seconds of loss of consciousness, lack of a prolonged upright posture, syncope during exercise or in response to auditory or emotional triggers, family history of SCD, abnormal physical examination, and abnormal ECG (626,627), although the specificity is modest (618,627,630,633). It should be remembered that children may not be able to clearly communicate specific symptoms. Exertional syncope has been associated with LQTS and CPVT (315,318,337,630,634). Regardless of symptoms, exertional syncope, especially mid-exertional syncope, should result in a high index of suspicion for a cardiac etiology (633).
		Noninvasive diagnostic testing should be performed in pediatric patients
I C-LD		presenting with syncope and suspected CHD, cardiomyopathy, or primary rhythm disorder (315,318,618,625,627,630,633).
See Online Data Supplement 40.		Channelopathies are major causes of cardiac-related syncope in young people. They may be associated with a family history of SCD, and they increase the risk of SCD in these patients (315,337,630,632,634,635). Exercise stress testing may be helpful in the diagnosis of channelopathies, such as LQTS and CPVT, which have adrenergically mediated arrhythmias. Extended monitoring is reasonable when an arrhythmia diagnosis is suspected. The types of monitoring devices, their clinical utility, and their limitations are available in Table 8. Prolonged heart rhythm monitoring can often provide a correlation between symptoms and an arrhythmia. In 5 retrospective studies of prolonged monitoring in 87 children with either syncope or presyncope, the mean diagnostic yield was 43% (636-640). Bradyarrhythmias and high-grade AV block or asystole, as well as tachyarrhythmias, SVT, and polymorphic VT, were documented (636-640). The diagnostic yield of an ICM is higher if the clinical indication was exertional syncope or the patient had underlying CHD (637,639,640).
I	С-ЕО	Education on symptom awareness of prodromes and reassurance are indicated in pediatric patients with VVS.

See Online Data Supplement 40.		Management of children with VVS should include reassurance about the generally benign nature of this condition (641,642). Treatment should emphasize symptom awareness and avoidance of precipitating factors that might worsen the condition, such as dehydration, standing for prolonged periods of time, hot crowded environments, and diuretic intake.
IIa	when the diagnosis is unclear (024,029,045-050).	
See Online Data Supplement 40.		Tilt-table testing has a diminishing role in the diagnosis of children with unexplained syncope. The sensitivity of tilt-table testing ranges from 20% to 90% (624,629,643,644,647,648,651,652), and the specificity ranges from 83% to 100% (624,643,652). Pediatric patients with episodes of VVS may exhibit convulsive movements during loss of consciousness that mimic epileptic seizures. In children with syncope and convulsions on tilt-table testing, 64% exhibited cardiac asystole with pauses >3 seconds (645). Upright tilt-table testing combined with a graded isoproterenol infusion identified 42% to 67% of patients previously thought to have a primary seizure disorder (223,649). A combined cardiology and neurology evaluation may be warranted in this group of patients with syncope and seizure-like activity.
		In pediatric patients with VVS not responding to lifestyle measures, it is ciation
IIa	B-R	reasonable to prescribe midodrine (381,620,653).
See Onli Supplen	ine Data nent 40.	In a single-center prospective case series, pseudoephedrine reduced clinical symptoms in 94% of children with recurrent neurally mediated syncope (653). In an RCT comparing patients receiving conventional therapy (health education, tilt-table training, and salt) and midodrine with patients receiving conventional therapy alone, the recurrence rate of syncope decreased from 80% to 22% (381). In 2 prospective studies, side effects from midodrine were rare (381,653).
IIb	B-R	Encouraging increased salt and fluid intake may be reasonable in selected
See Onli Supplen		pediatric patients with VVS (642).  In an RCT, conventional therapy and oral rehydration salts resulted in no further recurrence of syncope in 56% of patients, versus 39% in the placebo arm (p<0.05) (642).
IIb	C-LD	The effectiveness of fludrocortisone is uncertain in pediatric patients with OH associated with syncope (389,654,655).
See Online Data Supplement 40.		In 2 single-center prospective case series of 0.1 mg of fludrocortisone, 83% of subjects demonstrated improvement or resolution of symptoms (654,655). In the only pediatric RCT, children with recurrent syncope did better on placebo than on fludrocortisone (389).
IIb	B-NR	Cardiac pacing may be considered in pediatric patients with severe neurally mediated syncope secondary to pallid breath-holding spells (656,657).
See Onli Supplen		In 2 separate studies of 22 predominantly infants and toddlers with reflex anoxic seizures, pallid breath-holding spells, and documented prolonged asystole (pauses >4 s), 86% had either complete resolution or a significant reduction in the number of syncopal events with pacing (656,657). Although the studies were not powered to address the specifics of pacing programming, either single- or dual-chamber pacing significantly reduced the number of syncopal episodes compared with a sensing-only strategy (656,657). Single-chamber pacing with

		hysteresis appears as effective as dual-chamber pacing with rate drop response
		for the prevention of syncope and seizures. The beneficial response to pacing in
		these studies cannot exclude a placebo effect from pacemaker implantation itself;
		however, the young age of the patients with pallid breath-holding spells makes
		placebo effect less likely. The long-term outcome with pacing in this population
		has not been reported. Finally, it is important to remember that pallid breath-
		holding syncope does end, although some patients do present again at a later age
		with classic VVS. This should be balanced against the known complications of
		permanent cardiac pacing.
III: No	B-R	Beta blockers are not beneficial in pediatric patients with VVS (655,658).
Benefit	D-K	
See Online Data		In an RCT comparing metoprolol and conventional therapy, the treatment group
		actually had a higher recurrence rate. Side effects of beta blockers occur
Supplement 40.		frequently in children (655,659).

#### 10.2. Adult Congenital Heart Disease: Recommendations

Patients with ACHD are at risk for syncope as a result not only of the underlying structural disease, but also as a result of a previous palliative or corrective surgery. These patients may present with syncope of both hemodynamic and either bradycardic or tachycardic origin. Care by a physician with experience in management of CHD can be beneficial. The entire spectrum of arrhythmias may be seen in adults with CHD, including bradyarrhythmias secondary to sinus or AV nodal disease, atrial arrhythmias, and VA. By age 50, approximately 38% of patients with ACHD will develop an atrial arrhythmia, and by age 65, >50% of patients with severe CHD will develop atrial arrhythmias (660). The prevalence of VT after tetralogy of Fallot repair is 3% to 14% (661,662).

Recommen	Recommendations for ACHD		
COR	LOE	Recommendations	
IIa	С-ЕО	For evaluation of patients with ACHD and syncope, referral to a specialist with expertise in ACHD can be beneficial.	
		The care of the expanding population of ACHD survivors is complex, especially	
		in patients with moderate-to-severe ACHD. Care providers must be	
		knowledgeable in the anatomy and repair; be vigilant in the recognition and	
		management of HF, arrhythmias, and pulmonary hypertension; and have a deep	
N	/ <b>A</b>	understanding of noncardiac comorbidities. Delivery of ACHD care in highly	
IN/	/A	specialized centers has been shown to reduce mortality rate (663). In a	
		population-based retrospective study of 71,467 patients with ACHD from	
		Quebec, Canada, between 1990 and 2005, care in a specialized referral center for	
		ACHD care, compared with other care, was independently associated with	
		reduced mortality rate, particularly in those with severe ACHD (663).	
TT-	D ND	EPS is reasonable in patients with moderate or severe ACHD and	
IIa	B-NR	unexplained syncope (664,665).	
See Onl	ine Data	SCD is a leading cause of death in the patient with ACHD. Unexplained syncope	
Supplement 40.		is a concerning event. In a cohort of 252 patients with repaired tetralogy of Fallot	

undergoing risk stratification with programmed ventricular stimulation, induction of either monomorphic or polymorphic VT predicted future clinical VT and SCD (664). Patients with tetralogy of Fallot and inducible monomorphic or polymorphic VT were more likely to have a history of syncope (42.9%) than were those without inducible VT (13.4%) (664). In a cohort study of ICD recipients with transposition of the great arteries after an atrial baffle procedure, 35% of patients with primary-prevention ICDs presented with syncope. In 50% of patients receiving appropriate ICD shocks, atrial tachyarrhythmias preceded or coexisted with VT (665). It is reasonable to exclude atrial arrhythmias in patients with syncope and a CHD substrate at risk of atrial arrhythmias (e.g., Mustard, Senning, Fontan, Ebstein anomaly, and tetralogy of Fallot) (665).

#### 10.3. Geriatric Patients: Recommendations

The management of syncope in older adults is particularly challenging: The incidence is high; the differential diagnosis is broad; the diagnosis is imprecise because of amnesia, falls, lack of witnesses, and polypharmacy; and secondary morbidity is high because of comorbidities, physical injury, and frailty (35,45,666-675). The vulnerability of older adults to syncope increases because of age-associated cardiovascular and autonomic changes, decreased fluid conservation (45,671,676-678), and an increased probability of developing multiple concurrent morbidities (with their associated pharmacological treatments) that can overwhelm homeostasis. In many instances, a syncopal event in an older adult is multifactorial, with many predisposing factors present simultaneously.

Older patients (>75 years of age) who present with syncope tend to have poor outcomes, both fatal and nonfatal (109,679,680). Although some of the risk is attributable to the aspects of syncope described in this guideline, among older adults such risks are usually compounded by multiple morbidities and frailty, which add to age-related vulnerability to syncope (671,681,682), and by the physical injuries associated with falls, collisions, or trauma, which more commonly result from syncope in old age (670). Furthermore, recurrent syncope can lead to nursing home admission and a devastating loss of independence (683). Given the multifactorial etiologies and high risks associated with syncope, a comprehensive and multidisciplinary approach is often necessary to assess for multiple morbidities, frailty, trauma, and other dimensions of health (including cognition and medications) pertinent to diagnosis and management (77,188,684,685). A thorough history and physical examination, including orthostatic vital signs, is particularly important in older patients (77).

Recommen	Recommendations for Geriatric Patients		
COR	COR LOE Recommendations		
		For the assessment and management of older adults with syncope, a	
IIa	C-EO	comprehensive approach in collaboration with an expert in geriatric care	
		can be beneficial.	
·		A multidisciplinary approach helps to facilitate diagnosis of frailty and other	
N/A		factors that predispose to syncope and poor outcome in older adults. The goal is	
		to make management decisions in which older patients are well informed,	
		therapeutic choices are tailored to each patient's needs and goals of care, and	

		decision making is successfully shared between patients and providers.					
		Diagnostic and therapeutic approaches to syncope should incorporate					
		considerations of age, comorbid illness, physical and cognitive functions, patient					
		preferences, and severity of symptoms. Assessment is required of underlying					
		cardiovascular and noncardiovascular diseases; use of medications (e.g.,					
		polypharmacy, drug-drug interaction, age-related reduction in hepatic and renal					
		clearance); the potential to reduce medications that might lower blood pressure;					
		and circumstantial factors, such as dehydration, infection, or fever.					
		Consideration of frailty is particularly relevant. Characteristics of frailty					
		include weight loss, weakness, exhaustion, reduced physical activity, physical					
		slowing, and cognitive decline, with cumulative severity and impact that					
		typically vary between patients and even in 1 patient over time.					
II. D	ND	It is reasonable to consider syncope as a cause of nonaccidental falls in older					
IIa B	8-NR	adults (666-669,686).					
		Approximately 30% of older adults who present with nonaccidental falls may					
See Online Data Supplement 41.		have had syncope (687). Amnesia is commonly associated with both falls and					
		loss of consciousness, which diminishes the effectiveness of the history.					
		Cognitive impairment is also frequently present in older adults, even in those					
		without a formal diagnosis of dementia (688-690), and this too can reduce the					
· ·		without a formal diagnosis of dementia (688-690), and this too can reduce the					
		Approximately 30% of older adults who present with nonaccidental falls may have had syncope (687). Amnesia is commonly associated with both falls and loss of consciousness, which diminishes the effectiveness of the history. Cognitive impairment is also frequently present in older adults, even in those					

#### 10.4. Driving and Syncope: Recommendation

The assessment of medical fitness to drive is a common issue for practitioners caring for patients with syncope. The main concern is the risk of causing injury or death to the driver or others as a result of recurrent syncope (691). Factors to consider in assessing the risk of syncope while driving are summarized in a formula developed by the Canadian Cardiovascular Society 25 years ago (692) that estimates the risk that a driver will suddenly become incapacitated. The acceptable level of risk then becomes a societal decision.

Balancing the need to minimize risk from drivers fainting is the need for patients to drive to meet the demands of family and work. Society recognizes that certain groups, such as younger and older adults, are allowed to drive despite their higher risk of causing harm for reasons other than syncope (693). The societally acceptable risk of injury and death due to motor vehicle accidents has been quantified from an analysis of accident data collected in the United States, United Kingdom, and Canada (694). In the general population, the yearly risk of serious injury and death is 0.067%, or 1 in 1,500 (694). The 418 patients in POST I and POST II had a median of 3 vasovagal faints in 1 year but had no serious injuries or deaths and only 2 minor accidents in the subsequent year (694). This provides an estimated yearly risk of serious injury and death in the VVS population of <0.0017%, less than the Risk of Harm formula predicted (692). However, for patients with other etiologies of syncope or those in whom syncope occurred without prodrome or warning, the risk of causing harm may be higher than for patients with VVS. Public policies, laws, and regulations have not been adapted to these results, and providers caring for patients with syncope should be aware of pertinent local driving laws and restrictions. Although untreated syncope may disqualify patients from driving, effective treatment reduces the © 2017 by the American College of Cardiology Foundation, American Heart Association, Inc., and Heart Rhythm Society

risk enough to permit driving after a period of observation has elapsed without recurrent syncope. Regulatory agencies are more likely to disqualify commercial drivers than private drivers because of the amount of driving and the impact of accidents (i.e., commercial drivers typically operate vehicles heavier than private automobiles). As the risk of recurrent syncope decreases with treatment or with the natural history of a disease process, the risk of harm may become low enough for private drivers to resuming driving, but not necessarily for commercial drivers because of the higher risk of harm. The suggestions in Table 10 provide general guidance for private drivers. Most suggestions are based on expert opinion and supported by limited data. Commercial driving in the United States is governed by federal law and administered by the U.S. Department of Transportation (695).

Recommendation for Driving and Syncope							
COR	LOE	Recommendation					
		It can be beneficial for healthcare providers managing patients with					
IIa	C-EO	syncope to know the driving laws and restrictions in their regions and					
		discuss implications with the patient.					
		The writing committee encourages healthcare providers who care for patients					
		with syncope to know pertinent driving laws and restrictions in their region (e.g.,					
		states or provinces), as well as the duty of drivers or physicians to report Heart					
		inability of an individual to drive a motor vehicle. The Risk of Harm formula					
		simply estimates risk and does not supersede local driving regulations (692). In					
		the United States, private driving is state regulated, but commercial driving					
		requiring a U.S. Department of Transportation commercial driver's license is					
		federally regulated. Recommendations about commercial driving are more a					
N/	A	legal than a medical matter, and are not within the purview of this guideline.					
		Physicians providing care to commercial drivers should be familiar with U.S.					
		Department of Transportation policy (695).					
		Individual states may require reporting of drivers who faint. Many					
		patients do not stop driving despite advice to do so, regardless of the duration of					
		restriction (696,697). Although physicians have an obligation to maintain					
		confidentiality, if a patient's condition poses a significant risk to others, then th					
information should be reported as specific laws require.							

**Table 10. Avoidance of Private Driving After an Episode of Syncope: Suggested Symptom-Free Waiting Times for Various Conditions** 

vialing Times for Various Conditions							
Condition	Symptom-Free Waiting Time*						
OH	1 month						
VVS, no syncope in prior year (698)	No restriction						
VVS, 1–6 syncope per year (694)	1 month						
VVS, >6 syncope per year (694,698)	Not fit to drive until symptoms resolved						
Situational syncope other than cough syncope	1 month						
Cough syncope, untreated	Not fit to drive						
Cough syncope, treated with cough suppression	1 month						
Carotid sinus syncope, untreated (698)	Not fit to drive						
Carotid sinus syncope, treated with permanent pacemaker (698)	1 week						
Syncope due to nonreflex bradycardia, untreated (698)	Not fit to drive						
Syncope due to nonreflex bradycardia, treated with permanent	1 week						
pacemaker (12,698)							
Syncope due to SVT, untreated (698)	Not fit to drive						

Syncope due to SVT, pharmacologically suppressed (698)	1 month
Syncope due to SVT, treated with ablation (698)	1 week
Syncope with LVEF <35% and a presumed arrhythmic etiology	Not fit to drive
without an ICD (699,700)	
Syncope with LVEF <35% and presumed arrhythmic etiology	3 months
with an ICD (701,702)	
Syncope presumed due to VT/VF, structural heart disease, and	Not fit to drive
LVEF ≥35%, untreated	
Syncope presumed due to VT/VF, structural heart disease, and	3 months
LVEF ≥35%, treated with an ICD and guideline-directed drug	
therapy (701,702)	
Syncope presumed due to VT with a genetic cause, untreated	Not fit to drive
Syncope presumed due to VT with a genetic cause, treated with an	3 months
ICD or guideline-directed drug therapy	
Syncope presumed due to a nonstructural heart disease VT, such	Not fit to drive
as RVOT or LVOT, untreated	
Syncope presumed due to a nonstructural heart disease VT, such	3 months
as RVOT or LVOT, treated successfully with ablation or	
suppressed pharmacologically (698)	
Syncope of undetermined etiology	1 month

<sup>\*</sup>It may be prudent to wait and observe for this time without a syncope spell before resuming driving. ICD indicates implantable cardioverter-defibrillator; LVEF, left ventricular ejection fraction; LVOT, left ventricular outflow tract; OH, orthostatic hypotension; RVOT, right ventricular outflow tract; SVT, supraventricular tachycardia; VF, ventricular fibrillation; VT, ventricular tachycardia; and VVS, vasovagal syncope.

#### 10.5. Athletes: Recommendations

Syncope occurring in the athlete is predominantly of vasovagal origin, but underlying cardiac conditions may place athletes at undue risk for adverse events (703). Syncope during exercise is associated with increased probability of cardiac causes of syncope (Table 4). A thorough history, differentiating syncope occurring during exercise from syncope occurring after exercise or at other times, with typical characteristics of dehydration or VVS, is critically important during initial evaluation. The definition of an athlete is imprecise, but *athlete* can be defined as someone who engages in routine vigorous training (e.g., >150 minutes per week) and is skilled in exercises, sports, or games requiring physical strength, agility, or stamina (704). More importantly, cardiac adaptations to high levels of exercise may lead to the "athlete's heart" and thus alter the myocardial substrate (705). Primary or secondary prevention of syncope, morbidity, and mortality in at-risk athletes is a major consideration, but current strategies are largely inadequate (706). The current evidence base is insufficient to support general screening with ECG or echocardiography at baseline (706,707).

Several approved therapeutics, especially macrolide antibiotics and antihistamines/decongestants, have been associated with syncopal episodes (708). Performance-enhancing agents, such as somatotrophic compounds and amphetamine-like stimulants, are associated with precipitous collapse. A careful history is required in the athlete with syncope to rule out exposure to any of these agents (709). Similarly, before drugs are prescribed to highly competitive athletes, it is prudent to determine whether the drug or its metabolites are on lists of banned substances.

#### **Recommendations for Athletes**

COR	LOE	Recommendations					
_	C EO	Cardiovascular assessment by a care provider experienced in treating					
I C-EO		athletes with syncope is recommended prior to resuming competitive sports.					
N/A		A thorough history and physical examination should be completed by an experienced provider, including an assessment for OH and evidence of underlying cardiovascular disease (709-711). Cardiovascular causes account for 75% of sport-related deaths in young athletes (709,710). Syncope that occurs after exercise is often of benign origin and may be due to abdominal venous pooling. However, syncope during exercise is a much more compelling symptom and can be a harbinger of SCD (712,713). Syncopal episodes first require a personal and family history to evaluate precipitating causes and benign conditions, particularly volume depletion and vasovagal activity. Concomitant illnesses, especially viral infections, should be investigated and an ECG obtained (709,710).					
IIa	C-LD	Assessment by a specialist with disease-specific expertise is reasonable for athletes with syncope and high-risk markers (706,714).					
N/A		Syncope in the competitive athlete requires an evaluation for potentially fatal causes of syncope, especially when evidence of HCM, LQTS, Wolff-Parkinson-White syndrome, ARVC, ventricular noncompaction, symptomatic mitral valve prolapse, Marfan syndrome, congenital coronary anomalies, or other at-risk conditions is present (706,709,715,716). Any suspected cardiovascular pathology requires further evaluation, and family counseling and/or genetic testing is advised for those conditions with a known familial tendency.					
IIa	C-LD	Extended monitoring can be beneficial for athletes with unexplained exertional syncope after an initial cardiovascular evaluation (717,718).					
N/A		For those with a suspected cardiovascular etiology of syncope, an evaluation includes an ECG, tilt-table testing, and imaging as clinically indicated (Figure 3) (719). Imaging may include echocardiography or MRI as required. Exercise stress testing, unless contraindicated, can be helpful. For persistent unexplained syncope, extended arrhythmia monitoring can be used, as appropriate. This is a rapidly evolving field, with no firm data on the best device and optimum monitoring period (720).					
Participation in competitive sports is not recommended for athletes v syncope and phenotype-positive HCM, CPVT, LQTS1, or ARVC bef evaluation by a specialist (704,721-724).							
See Online Data Supplement 42.		In the absence of vagal mechanisms, VA in patients with HCM, CPVT, LQTS1, or ARVC is catecholamine sensitive. Participation in competitive sports in that circumstance in these patients is not recommended (704,715,716).					

### 11. Quality of Life and Healthcare Cost of Syncope

### 11.1. Impact of Syncope on Quality of Life

QoL is reduced with recurrent syncope (725-733), as demonstrated in studies that compared patients with and without syncope (727,731). QoL associated with recurrent syncope was equivalent to severe rheumatoid arthritis and chronic low-back pain in an adult population (728). Similarly, pediatric patients with recurrent syncope reported worse QoL than individuals with diabetes mellitus and equivalent QoL to individuals with asthma, end-stage renal disease, and structural heart disease (725). In a hospital-based cohort of patients with a prior episode of syncope, 33% reported syncope-related functional impairments with daily activities, such as driving or working (732). Those with more frequent syncope have reported poorer QoL (726,729,730,732). There is consistent evidence that syncope is associated with worse function on multiple domains of QoL, such as perceptions of low overall physical health (725,730,734); perception of mental health, including increased fear, somatization, depression, and anxiety (725,727,728,731,734); and impairment in activities of daily living, such as driving, working, and attending school.

QoL impairments associated with syncope improve over time (733). In the Fainting Assessment Study (733), general and syncope-specific QoL improved over a 1-year period. Predictors of worse QoL over time include advanced age, recurrent syncope, neurological or psychogenic reason for syncope, and greater comorbidity at baseline (733). Syncope-related QoL can be improved through effective diagnosis and treatment. In 1 study, use of an implantable loop recorder increased diagnostic rate, reduced syncope recurrence, and improved QoL as compared with patients who received a conventional diagnostic workup (164). In a second study, nonpharmacological treatment of recurrent syncope was associated with reductions in recurrent syncope and improvements in QoL (729).

## 11.2. Healthcare Costs Associated With Syncope

High healthcare costs are associated with the evaluation and management of syncope. Costs are defined as the resources needed to produce a set of services and are distinct from charges billed by facilities and healthcare providers (735). Most studies have focused on facility costs and excluded professional fees and patient copays. These high costs have been estimated both in the United States and abroad. In the U.S. Healthcare Utilization Project, total annual hospital costs exceeded \$4.1 billion in 2014 dollars, with a mean cost of \$9,400 per admission (736). Total costs and costs per admission for presumptive undiagnosed syncope were \$1.6 billion and \$7,200, respectively (736). Single-center studies from multiple countries, including Austria, the United Kingdom, Israel, and Spain, confirm similarly high costs associated with the hospital evaluation of syncope (122,737,738).

Several investigators have estimated the costs per clinically meaningful test result. Physician reviewers determined whether the results of a diagnostic test affected clinical management at a U.S. tertiary referral hospital after an episode of syncope (77). The cost per informative diagnosis (as ordered in routine practice)

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affecting clinical management varied widely by specific diagnostic test, from postural blood pressure (\$50) through telemetry (\$1,100) to EEG (\$32,973) (77). Similar high costs per actionable diagnosis occur in children admitted for new-onset syncope. Finally, mean costs per diagnostic result were also high in an outpatient (\$19,900) specialty clinic for unexplained recurrent syncope (163).

#### 12. Emerging Technology, Evidence Gaps, and Future Directions

The writing committee created a list of key areas in which knowledge gaps are present in the evaluation and management of patients presenting with syncope. These knowledge gaps present opportunities for future research to ultimately improve clinical outcomes and effectiveness of healthcare delivery.

### 12.1. Definition, Classification, and Epidemiology

Reported incidence and prevalence of syncope vary significantly because of several confounders: variable definitions for syncope versus transient loss of consciousness, different populations, different clinical settings, and different study methodologies. Definition and classification of syncope provided in this document will set the standard for future research. Standardized national registries and large sample databases are needed to gather data on a continuous basis to understand the true incidence and prevalence of syncope, understand patient risk, inform driving policies, improve patient outcomes, and improve and streamline health service delivery.

### 12.2. Risk Stratification and Clinical Outcomes

At a patient's presentation, several key questions follow: What is the likely cause of syncope? Does the patient have significant underlying heart disease and/or comorbid medical illnesses? If the cause of syncope is determined, is there an effective therapy to prevent recurrent syncope, prevent syncope-related nonfatal outcomes (injury, diminished health care—related QoL, lost workdays), or improve survival? What are the predictors of short- and long-term clinical outcomes? What are the key outcomes relevant to patients with syncope, including recurrent syncope? When the cause of syncope is unknown, what is the standard of care for this group of patients?

- Studies are needed to determine whether syncope is an independent predictor of nonfatal or fatal outcomes in selected patient populations.
- Studies are needed to develop risk scores to be prospectively validated in a given clinical setting with predefined endpoints from short- and long-term follow-up.
- Prospective and well-designed studies are needed to define relevant clinical outcomes with regard to
  recurrent syncope, nonfatal outcomes such as injury, and fatal outcomes. Future studies should
  incorporate QoL, work loss, and functional capacity as additional clinical endpoints.

- Prospective studies are needed to differentiate cardiac and noncardiac clinical outcomes in different clinical settings and with different follow-up durations.
- Among patients without identifiable causes of syncope, studies are needed to determine short- and long-term outcomes to guide the overall management of these patients.

#### 12.3. Evaluation and Diagnosis

Because of the concerns that patients presenting with syncope are at higher risk for an impending catastrophic event, overuse and inappropriate use of testing and hospital admission are common. Answers to the following question will improve the effectiveness of patient evaluation: How should the initial evaluation and subsequent follow-up vary by risk (low, intermediate, or high) to assess clinical outcomes?

- Studies are needed to better understand the interaction and relationships among the presenting symptom of syncope, the cause of syncope, the underlying disease condition, and their effect on clinical outcomes.
- Investigations are needed to understand the key components of clinical characteristics during the initial evaluation and to develop standardization tools to guide the evaluation by healthcare team.
- RCTs are needed to develop structured protocols to evaluate patients with syncope who are at intermediate risk without an immediate presumptive diagnosis. In addition to the endpoints of diagnostic yield and healthcare utilization, relevant clinical endpoints of nonfatal and fatal outcomes and recurrence of syncope are to be included.
- RCTs are needed to determine the features of syncope-specialized facilities that are necessary to
  achieve beneficial outcomes for patient care and to improve efficiency and effectiveness of healthcare
  delivery.
- As technology advances, studies are needed to determine the value of new technology in the evaluation and management of patients with syncope.

#### 12.4. Management of Specific Conditions

- Although potential causes of syncope are multiple, a treatment decision is usually fairly
  straightforward for patients with cardiac causes of syncope or orthostatic causes. Vasovagal syncope is
  the most common cause of syncope in the general population. Treatment remains challenging in
  patients who have recurrences despite conservative therapy. Studies are needed to differentiate
  "arrhythmic syncope" versus "nonarrhythmic syncope" versus "aborted SCD" in patients with
  inheritable arrhythmic conditions
- Prospectively designed multicenter or national registries are needed to gather clinical information from patients with reflex syncope to better our understanding on other associated conditions, plausible

mechanisms, effectiveness of therapeutic interventions, and natural history of these uncommon conditions.

 RCTs are needed to continue the identification of effective treatment approaches to patients with recurrent reflex syncope.

### 12.5. Special Populations

Each population in Section 6 is unique with regard to syncope, and within each of them we identified several key areas that are important for future research considerations.

- Questions and research about risk stratification, evaluation, and management outlined above for the adult population are needed in the pediatric population, geriatric population, and athletes.
- Prospective national registries and big databases are needed to determine risk associated with driving among different populations with syncope.
- Prospective and randomized studies are needed to assess the usefulness of specialized syncope units in different clinical settings.



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Richard A. Chazal, MD, FACC, President

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**Key Words**: AHA Scientific Statements ■ syncope ■ risk assessment ■ diagnosis ■ prognosis ■ cardiac syncope ■ reflex syncope ■ vasovagal syncope ■ orthostatic hypotension ■ neurogenic syncope ■ dehydration ■ pediatrics ■ adult congenital heart disease ■ geriatrics ■ driving ■ athletes

## **Appendix 1. Author Relationships With Industry and Other Entities (Relevant)**—2017 ACC/AHA/HRS

Guideline for the Evaluation and Management of Patients With Syncope (March 2015)

Committee Member	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness	Voting Recusals by Section*
Win-Kuang Shen (Chair)	Mayo Clinic Arizona— Professor of Medicine; Mayo Clinic College of Medicine—Chair, Department of Cardiovascular Diseases	None	None	None	None	None	None American	None
Robert S. Sheldon (Vice Chair)	University of Calgary Department of Medicine— Professor	None	None	None	None	None	None L Association	None
David G. Benditt	University of Minnesota Medical School, Cardiovascular Division— Professor of Medicine	Medtronic†     St. Jude     Medical†	None	None	None	None	None	3.2, 3.2.3, 3.2.5, 4.1.1– 4.1.3, 4.2.1– 4.2.5, 4.3.1– 4.3.5, 5.1–5.3, 10.1, 10.2, 10.3, 10.5, 12
Mitchell I. Cohen	University of Arizona School of Medicine- Phoenix—Clinical Professor of Child Health; Phoenix Children's Heart Center— Co-Director; Phoenix Children's Hospital, Pediatric Cardiology— Chief	None	None	None	None	None	None	None
Daniel E. Forman	University of Pittsburgh— Professor of Medicine; University of Pittsburgh Medical Center—Chair, Geriatric Cardiology Section; VA Pittsburg Healthcare Systems— Director, Cardiac Rehabilitation	None	None	None	None	None	None	None

Roy Freeman§	Harvard Medical School— Professor of Neurology; Beth Israel Deaconess Medical Center, Center for Autonomic and Peripheral Nerve Disorders—Director	• Lundbeck†	None	None	None	None	None	4.3.1–4.3.5, 5.1, 6.1, 10.1, 10.3, 10.5, 12
Zachary D. Goldberger	University of Washington School of Medicine, Harborview Medical Center Division of Cardiology— Assistant Professor of Medicine	None	None	None	None	None	None	None
Blair P. Grubb	University of Toledo Medical Center, Medicine and Pediatrics—Professor	Biotronik     Medtronic	None	None	None	None	None can Heart Association	3.2, 3.2.3, 3.2.5, 4.1.1– 4.1.3, 4.2.1– 4.2.5, 4.3.1– 4.3.5, 5.1–5.3, 10.1, 10.2, 10.3, 10.5, 12
Mohamed H. Hamdan	University of Wisconsin School of Medicine, Cardiovascular Medicine— Professor and Chief of Cardiovascular Medicine	None	None	• F2 Solutions	None	None	None	2.3.3, 2.3.4, 12
Andrew D. Krahn	The University of British Columbia, Division of Cardiology—Professor of Medicine and Head of Division	Medtronic	None	None	None	<ul><li>Boston Scientific†</li><li>Medtronic†</li></ul>	None	3.2, 3.2.3, 3.2.5, 4.1.1– 4.1.3, 4.2.1– 4.2.5, 4.3.1– 4.3.5, 5.1–5.3, 10.1, 10.2, 10.3, 10.5, 12
Mark S. Link	University of Texas Southwestern Medical Center, Department of Medicine, Division of Cardiology—Director, Cardiac Electrophysiology; Professor of Medicine	None	None	None	None	None	None	None
Brian Olshansky	University of Iowa Carver College of Medicine, Cardiovascular Medicine—	• Lundbeck†	None	None	None	None	None	None

	Emeritus Professor of Internal Medicine; Mercy Hospital North Iowa— Electrophysiologist							
Satish R. Raj	University of Calgary, Cardiac Sciences— Associate Professor	GE Healthcare     Lundbeck†	None	None	Medtronic	None	None	2.3.2, 2.3.4, 3.2–3.2.5, 3.3.2, 4.1.1– 4.1.3, 4.2.1– 4.2.5, 4.3.1– 4.3.5, 5.1–5.3, 6.1, 7, 10.1– 10.3, 10.5, 12
Roopinder Kaur Sandhu	University of Alberta, Medical Division of Cardiology—Assistant Professor of Medicine	None	None	None	None	None	ANonecan Heart Association	None
Dan Sorajja	Mayo Clinic Arizona, Cardiovascular Diseases— Assistant Professor of medicine	None	None	None	None	None	None	None
Benjamin C. Sun	Oregon Health & Science University—Associate Professor	None	None	None	None	None	None	None
Clyde W. Yancy	Northwestern University Feinberg School of Medicine, Division of Cardiology—Professor of Medicine and Chief; Diversity & Inclusion— Vice Dean	None	None	None	None	None	None	None

This table represents the relationships of committee members with industry and other entities that were determined to be relevant to this document. These relationships were reviewed and updated in conjunction with all meetings and/or conference calls of the writing committee during the document development process. The table does not necessarily reflect relationships with industry at the time of publication. A person is deemed to have a significant interest in a business if the interest represents ownership of  $\geq$ 5% of the voting stock or share of the business entity, or ownership of  $\geq$ 55,000 of the fair market value of the business entity; or if funds received by the person from the business entity exceed 5% of the person's gross income for the previous year. Relationships that exist with no financial benefit are also included for the purpose of transparency. Relationships in this table are modest unless otherwise noted.

According to the ACC/AHA, a person has a *relevant* relationship IF: a) the *relationship or interest* relates to the same or similar subject matter, intellectual property or asset, topic, or issue addressed in the *document*; or b) the *company/entity* (with whom the relationship exists) makes a drug, drug class, or device addressed in the *document*, or makes a competing drug or device addressed in the *document*; or c) the *person or a member of the person's household*, has a reasonable potential for financial, professional or other personal gain or loss as a result of the issues/content addressed in the *document*.

\*Writing committee members are required to recuse themselves from voting on sections to which their specific relationships with industry and other entities may apply †Significant relationship.

‡No financial benefit.

§Dr. Roy Freeman, the official representative of the American Academy of Neurology, resigned from the writing committee in November 2016, before the final balloting process; recusals noted are from the initial round of balloting. We thank him for his contributions.





# **Appendix 2. Reviewer Relationships With Industry and Other Entities (Comprehensive)**—2017 ACC/AHA/HRS Guideline for the Evaluation and Management of Patients With Syncope (June 2016)

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
Italo Biaggioni	Official Reviewer—AHA	Vanderbilt University School of Medicine— Professor of Medicine	• Lundbeck* • Shire Pharmaceuticals* • Theravance*	None	None	<ul> <li>Astellas Pharma (DSMB)</li> <li>AstraZeneca*</li> <li>Forest Pharmaceuticals*</li> <li>Janssen Pharmaceuticals (DSMB)</li> <li>Lundbeck*</li> <li>Theravance*</li> </ul>	American Heart Association	None
Joaquin E. Cigarroa	Official Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Oregon Health & Science University— Clinical Professor of Medicine	None	None	None	None	<ul> <li>NIH†</li> <li>AHA†</li> <li>SCAI†</li> <li>ASA†</li> <li>Catheterization and Cardiovascular Intervention†</li> </ul>	None
Kenneth A. Ellenbogen	Official Reviewer—AHA	VCU Medical Center—Director, Clinical EP Laboratory	• AHA • Atricure* • Biosense Webster* • Biotronik* • Boston Science* • HRS* • Janssen Pharmaceuticals • Medtronic* • Pfizer* • Sentra Heart • St. Jude Medical*	None	None	• Atricure* • Boston Science • Biosense Webster • Daiichi-Sankyo* • Medtronic (DSMB) • Medtronic • NIH • Sanofi-aventis	• AHA • American Heart Journal • Biosense Webster* • Boston Science* • HRS • JCE • Medtronic* • PACE • Sanofi-aventis	<ul> <li>Defendant, Catheter ablation complication , 2015</li> <li>Plantiff, Lead extraction complication , 2015</li> </ul>
Rakesh Gopinathannair	Official Reviewer—HRS	University of Louisville School of	<ul><li>Boston Scientific</li><li>Health Trust PG</li></ul>	• AHA	None	None	None	None

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
		Medicine and Jewish Hospital Division of Cardiovascular Medicine—Associate Professor of Medicine, Director of Cardiac EP	• St. Jude Medical*	<ul><li>Bristol- Myers Squibb</li><li>Pfizer*</li><li>Zoll Medical</li></ul>				
Robert Helm	Official Reviewer—HRS	Boston University School of Medicine— Assistant Professor of Medicine, Assistant Professor of Radiology	None	None	None	None	Boston     Scientific     St. Jude Medical     Association	None
Dhanunjaya Lakkireddy	Official Reviewer—ACC Board of Governors	University of Kansas Medical Center— Professor of Medicine; Center for Excellence in AF and Complex Arrhythmias— Director	Biosense Webster     St. Jude Medical	Boehringer Ingleheim     Bristol-Meyer Squibb     Janssen Pharmaceutic als     Pfizer	None	None	None	None
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Christopher Gibbons	Organizational Reviewer—AAN	Beth Israel Deaconess Medical Center Neuropathy Clinic— Director	• Lundbeck	None	None	• Astellas Pharma (DSMB) • Janssen Pharmaceuticals (DSMB)	None	None
Kaushal H. Shah	Organizational Reviewer— ACEP/SAEM	The Mount Sinai Hospital—Associate Professor of Emergency Medicine	None	None	None	None	None	None
Mike Silka	Organizational Reviewer— PACES	Children's Hospital Los Angeles—	None	None	None	None	None	• Defendant,

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
		Professor of Pediatrics, Cardiology						SCD in CPVT patient, 2016
Sana M. Al- Khatib	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Duke Clinical Research Institute— Professor of Medicine	None	None	None	• FDA* • NHLBI* • PCORI* • VA Health System (DSMB)	• Elsevier* • AHA  American	None
Kim K. Birtcher	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	University of Houston College of Pharmacy—Clinical Professor	• Jones & Bartlett Learning	None	None	None	None Part Association	None
Michele Brignole	Content Reviewer	Arrhythmologic Centre, Ospedali del Tigullio—Head of Cardiology	None	None	• F2 Solutions†	None	None	None
Hugh Calkins	Content Reviewer—ACC EP Section Leadership Council	Johns Hopkins Hospital—Professor of Medicine, Director of EP	Abbott     Atricure     Boehringer     Ingelheim*     Medtronic*	None	None	Boehringer     Ingelheim†     St. Jude Medical*	Abbott     Laboratories	• Defendant, SCD, 2015
Coletta Barrett	Content Reviewer—Lay Reviewer	Our Lady of the Lake Regional Medical Center—Vice President	None	None	None	None	None	None
Lin Yee Chen	Content Reviewer	University of Minnesota Medical School—Associate Professor of Medicine	None	None	None	None	• NIH*	None
Andrew Epstein	Content Reviewer	University of Pennsylvania Hospital and the Veteran's Administration	None	None	None	Biosense     Webster*     Biotronik*	None	None

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
		Medical Center— Professor of Medicine				<ul> <li>Boston Scientific* (DSMB)</li> <li>Boston Scientific*</li> <li>C.R. Bard*</li> <li>Medtronic (DSMB)</li> <li>Medtronic*</li> <li>St. Jude Medical* (DSMB)</li> <li>St. Jude Medical</li> </ul>	American <b>Heart</b> Association	
Susan Etheridge	Content Reviewer—ACC EP Section Leadership Council	University of Utah— Training Program Director	None	None	None	• SADS Foundation • PACES†	• Up-to-Date†	None
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Samuel S. Gidding	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Nemours/Alfred I. duPont Hospital for Children—Chief, Division of Pediatric Cardiology	• FH Foundation† International • FH Foundation†	None	None	• FH Foundation† • NIH*	None	None
Bulent Gorenek	Content Reviewer— ACC EP Section Leadership Council	Eskisehir Osmangazi University Cardiology Department—Chair	None	None	None	None	None	None
Paul LeLorier	Content Reviewer—ACC Heart Failure and	LSU Health Sciences Center—Associate Professor of Medicine	None	None	None	• Medtronic*	• Medtronic*	None

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
	Transplant Section Leadership Council	and Neurology; EP Service—Director						
Patrick McBride	Content Reviewer	University of Wisconsin School of Medicine & Public Health—Professor of Medicine and Family Medicine; Dean for Faculty Affairs— Associate; Prevention Cardiology— Associate Director	None	None	None	None	American Heart Association	None
Carlos Morillo	Content Reviewer	Cumming School of Medicine—Professor Department of Cardiac Sciences; University of Calgary—Section Chief Division of Cardiology, Libin Cardiovascular Institute	<ul> <li>Bayer Healthcare</li> <li>Boston Scientific</li> <li>Boehringer Ingelheim</li> <li>Boston Scientific</li> </ul>	None	None	Biosense Webster     Canadian     Institutes for     Health Research†     Medtronic†     Merck     Pfizer     St. Jude Medical	Biotronik     Pfizer	None
Rick Nishimura	Content Reviewer	Mayo Clinic Division of Cardiovascular Disease—Professor of Medicine	None	None	None	None	None	None
Richard Page	Content Reviewer	University of Wisconsin School of Medicine & Public Health—Chair, Department of Medicine	None	None	None	None	• FDA	None
Antonio Raviele	Content Reviewer	Alliance to Fight Atrial Fibrillation— President; Venice	None	None	None	None	None	None

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
		Arrhythmias— President						
Marwan Refaat	Content Reviewer—ACC EP Section Leadership Council	American University of Beirut—Faculty of Medicine and Medical Center	None	None	None	None	None	None
Melissa Robinson	Content Reviewer	University of Washington— Assistant Professor of Medicine; Director, Ventricular Arrhythmia Program	• Medtronic*	None	None	None	Nonenerican Heart Association	None
Paola Sandroni	Content Reviewer	Mayo Clinic— Professor of Neurology, Practice Chair of Neurology	None	None	None	None	None	None
Colette Seifer	Content Reviewer	University of Manitoba—Associate Professor, Section of Cardiology,	None	None	None	None	None	None
Monica Solbiati	Content Reviewer	Fondazione IRCCS CA' Granda, Ospedale Maggiore Policlinico, Milano—Senior Physician	None	None	None	None	None	None
Richard Sutton	Content Reviewer	National Heart & Lung Institute, Imperial College— Emeritus Professor	• Medtronic*	• St. Jude Medical*	Boston     Scientific*     Edwards     Lifesciences     *      Shire     Pharma     AstraZenec     a	• Medtronic*	None	• Defendant, Fatal car accident caused by VVS patient, 3 trials in 2016*

Reviewer	Representation	Employment	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational, or Other Financial Benefit	Expert Witness
Gaurav Upadhyay	Content Reviewer—ACC EP Section Leadership Council	University of Chicago—Assistant Professor of Medicine	<ul> <li>Biosense Webster</li> <li>Biotronik</li> <li>Boston Scientific</li> <li>Medtronic</li> <li>St. Jude Medical</li> <li>Zoll Medical</li> </ul>	None	None	• Biotronik* • Medtronic* • Biosense Webster	None	None
Paul Varosy	Content Reviewer	University of Colorado Hospital, Clinical Cardiac EP Training program— Associate Program Director; VA Eastern Colorado Healthcare System—Director of Cardiovascular EP	None	None	None	• AHA† • VA Office of Health Services Research and Development (PI)*	None American Heart Association	None

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AAN indicates American Academy of Neurology; ACC; American College of Cardiology; ACEP, American College of Emergency Physicians; AHA, American Heart Association; ASA, American Stroke Association; DSMB, data safety monitoring board; CPVT, catecholaminergic polymorphic ventricular tachycardia; EP, electrophysiology; FDA, U.S. Food and Drug Administration; FH, familial hypercholesterolemia; HRS, Heart Rhythm Society; ICD, implantable cardioverter-defibrillator; JCE, *Journal of Cardiovascular Electrophysiology*; LSU, Louisiana State University; NHLBI, National Heart, Lung, and Blood Institute; PACE, Partners in Advanced Cardiac Evaluation; PACES, Pediatric and Congenital Electrophysiology Society; PCORI; Patient-Centered Outcomes Research Institute; PI, principal investigator; SADS, Sudden Arrhythmia Death Syndromes Foundation; SAEM, Society for Academic Emergency Medicine; SCAI, Society for Cardiovascular Angiography and Interventions; SCD, sudden cardiac death; VA, Veterans Affairs; VCU, Virginia Commonwealth University; and VVS, vasovagal syncope.

<sup>\*</sup>Significant relationship.

<sup>†</sup>No financial benefit.

#### **Appendix 3. Abbreviations**

ACHD = adult congenital heart disease

ARVC = arrhythmogenic right ventricular cardiomyopathy

AV = atrioventricular

CHD = congenital heart disease

CPVT = catecholaminergic polymorphic ventricular tachycardia

CT = computed tomography

ECG = electrocardiogram/electrocardiographic

ED = emergency department

EEG = electroencephalogram/electroencephalography

EPS = electrophysiological study

GDMT = guideline-directed management and therapy

HCM = hypertrophic cardiomyopathy

HF = heart failure

ICD = implantable cardioverter-defibrillator

ICM = implantable cardiac monitor

LCSD = left cardiac sympathetic denervation

LQTS = long-QT syndrome

LV = left ventricular

MRI = magnetic resonance imaging

OH = orthostatic hypotension

QoL= quality of life

RCT = randomized controlled trial

POTS = postural tachycardia syndrome

SCD = sudden cardiac death

SVT = supraventricular tachycardia

VA = ventricular arrhythmia

VF = ventricular fibrillation

VT = ventricular tachycardia

VVS = vasovagal syncope



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#### 2017 ACC/AHA/HRS Guideline for the Evaluation and Management of Patients With Syncope: A Report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines, and the Heart Rhythm Society

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Author Relationships With Industry and Other Entities (Comprehensive)—2017 ACC/AHA/HRS Guideline for the Evaluation and Management of

Patients With Syncope (March 2015)

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<sup>\*</sup>Significant relationship.

<sup>†</sup>No financial benefit.

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§Dr. Roy Freeman, the official representative of the American Academy of Neurology, resigned from the writing committee in November 2016, prior to the final balloting process; recusals noted are from the initial round of balloting. We thank him for his contributions.
ACC indicates American College of Cardiology; AHA, American Heart association; CANet, Cardiac Arrhythmia Network of Canada; CIHR, Canadian Institute of Health Research; DSMB, data safety monitoring board; ED, emergency department; HRS, Heart Rhythm Society; NIH, National Institutes of Health; PI, principal investigator, and VA, Veteran's Affairs.
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## 2017 ACC/AHA/HRS Guideline for the Evaluation and Management of Patients With Syncope - Data Supplement

(Section numbers correspond to the full-text guideline.)

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#### Methodology and Evidence Review

The recommendations listed in this guideline are, whenever possible, evidence based. An extensive evidence review was conducted from July through October 2015, that included literature published through October 2015. Other selected references published through May 2016 were incorporated by the writing committee. Literature included was derived from research involving human subjects, published in English, and indexed in MEDLINE (through PubMed), EMBASE, the Cochrane Library, the Agency for Healthcare Research and Quality, and other selected databases relevant to this guideline. Key search words included but were not limited to the following: adverse, aged, aging, ambulatory monitor, arrhythmogenic right ventricular cardiomyopathy, arrhythmogenic right ventricular dysplasia, athletes, AV block, b-blockers, biomarkers, blood pressure, bradycardia, breath-holding, Brugada Syndrome, cardiovascular disease, carotid sinus hypersensitivity, carotid sinus massage, carotid sinus syndrome, catecholaminergic polymorphic ventricular tachycardia, children, consciousness, dehydration, diagnosis, drug, early repolarization syndrome, echocardiogram, electrocardiogram, electrocardiography, electrophysiologic, electrophysiological, falls, florinef, fludrocortisone, fluoxetine, functional neurologic symptoms, heart rate, holter monitor, holter, hypertrophic cardiomyopathy, hypotension, ICD, idiopathic AV block, implantable cardioverter defibrillator, implantable loop recorder, laboratory testing, left cardiac sympathetic denervation, long QT Syndrome, loop monitor, loop recorder, medication, midodrine, mode of pacing, monitor, non-epileptic pseudo seizures, orthostatic, pacemaker, pacing, pediatrics, postural, pressure counter maneuvers, presyncope, psychogenic non-epileptic seizure, psychogenic pseudoseizures, psychogenic pseudoseizures,

Abbreviations 1° indicates primary; 2°, secondary; AAD, antiarrhythmic dug; AAI, atrioventricular interval; ACA, aborted cardiac arrest; ACS, acute coronary syndrome; ADE, indicates adverse drug events; AF, atrial fibrillation; AMI, acute myocardial infarction; ARVC, arrhythmogenic right ventricular cardiomyopathy; ARVC/D, arrhythmogenic right ventricular dysplasia/cardiomyopathy: ARVD, arrhythmogenic right ventricular dysplasia; AS, aortic stenosis; ASR, Anatolian Syncope Rule; AUC, appropriate use criteria; AV, atrioventricular; AVB, atrioventricular block; BB, beta blocker; BBB, bundle branch block; BID, two times a day; BNP, brain natriuretic peptide; BP, blood pressure; BS, Brugada syndrome; BSC, Boston Syncope Criteria; CA, cardiac arrest; CAA, carotid artery angioplasty; CAD, coronary artery disease; CBT, cognitive behavioral therapy; CCU, coronary care unit; CHD, congenital heart disease; CHF, congestive heart failure; CI, confidence interval; CLS, closed loop stimulation; CO, cardiac output; COPD, chronic obstructive pulmonary disease; CPR, cardiopulmonary resuscitation; CPVT, catecholaminergic polymorphic ventricular tachycardia; CS, carotid sarcoidosis; CSH, carotid sinus hypersensitivity; CSM, carotid sinus massage; CSR, carotid sinus reaction; CSS, Carotid Sinus Syndrome; CSSS, Calgary Syncope Symptom Score; CT, computed tomography; cTnThs, high-sensitivity cardiac troponin T; CV, cardiovascular; CVD, cardiovascular disease; CXR, chest x-ray; DBP, diastolic blood pressure; DDD, dual chamber pacing; DM, diabetes mellitus; DS, defecation syncope; DVI, dual chamber pacing; ECG, electrocardiogram; ED, emergency department; EDOSP, emergency department observation syncope protocol; EEG, electroencephalogram; EF, ejection fraction; EGSYS, evaluation of guidelines of syncope study; ELR, external loop recorder; EP, ejectrophysiological; EPS, electrophysiological study; ER, early repolarization; ERP, early repolarization pattern; EST, exercise stress test; FINGER, France, Italy, Netherlands, Germany, Registry; GERD, gastroesophageal reflux disease; GFR, glomerular filtration rate; GTN, glyceryl trinitrate; H&P, history and physical exam; HCM, hypertrophic cardiomyopathy; HF, heart failure; HR, hazard ratio; HTN, hypertension; HUTT, head-up tilt test; Hx, history; ICD, implantable cardioverter-defibrillator; ICU, intensive care unit; IDCM, idiopathic dilated cardiomyopathy; ILR, implantable loop recorder; IV, intravenous fluid: IVCD, intraventricular conduction disturbances; KM, Kaplan-Meier; LBBB, left bundle branch block; LBNP, lower body negative pressure; LCS, left cervicothoracic sympathectomy; LCSD, left cardiac sympathetic denervation; LOC, loss of consciousness; LOS, length of stay; LQTS, long QT syndrome; LV, left ventricular non-compaction; MACE, major adverse cardiac event; MAP, mean arterial pressure; MCA, middle cerebral artery blood velocity; MCOT, mobile cardiac outpatient telemetry; MD, doctor of medicine; MI, myocardial infarction; MRI, magnetic resonance imaging; MS, micturition syncope; MSA, multiple systems atrophy; N/A, not available; NICM, nonischemic dilated cardiomyopathy; NMS, neurally mediated syncope; NPV, negative predictive value; NS, not significant; NSVT, nonsustained ventricular tachycardia; NYHA, New York Heart Association classification for heart failure; ODO, sensing without pacing; OESIL, Osservatorio Epidemiologico sulla Sincope nel Lazio; OH, orthostatic hypotension; OHDAS, Orthostatic Hypotension Daily Activity Scale; OHQ, orthostatic hypotension guestionnaire; OHSA, Orthostatic Hypotension Symptom Assessment; OI, orthostatic intolerance; OR, odds ratio; OT, Oral Fluid and Trendelenburg position; OT, orthostatic tachycardia; PAF, pure autonomic failure; PCA, posterior cerebral artery blood velocity; PCI, percutaneous coronary intervention; PCM,

physical counter pressure maneuvers; PD, Parkinson disease; PE, physical examination; PES, programmed electrical stimulation; PM, pacemaker; PMVT, polymorphic ventricular arrhythmias; PNES, psychogenic nonepileptic seizures; POST, Prevention of Syncope trial; POTS, postural (orthostatic) tachycardia syndrome; PPM, permanent pacemaker; PPS, psychogenic pseudosyncope; PVC, premature ventricular contractions; PVD, peripheral vascular disease; QoL, quality of life; RCT, randomized controlled trials; RDBPCT, randomized, double blind, placebo-controlled trial; ROSE, risk stratification of Syncope in the Emergency Department; RR, relative risk; RRR relative risk ratio; RyR2, Ryanodine receptor type 2; S/P, strategies primary; SA, sinoatrial; SBP, systolic blood pressure; SCD, sudden cardiac death; SCI, spinal cord injury; SD, sudden death; SFSR, San Francisco Syncope Rule; SHD, structural heart disease; SN, sinus node; SND, sinus node dysfunction; SNRT, sinus node recovery time; SNS, sympathetic nervous system; SQTS, short QT syndrome; SUO, syncope of unknown origin; SV, stroke volume; SVT, supraventricular tachycardia; TCA, trichloroacetic acid; TIA, transient ischemic attack; TLOC, transient loss of consciousness; TOF, tetralogy of Fallot; TPR, total peripheral resistance; TST, thermoregulatory sweat test; TTT, tilt-table test; VA, ventricular arrhythmias; VATS, video-assisted thoracic surgery; VF, ventricular fibrillation; VFL, ventricular flutter; VHD, valvular heart disease; VS, vital signs; VT, ventricular tachycardia; VVI, ventricular pacing; VVS, vasovagal syncope; and WPW, Wolff-Parkinson-White.

## Data Supplement 1. Nonrandomized Trials, Observational Studies, and/or Registries for History and Physical Exam – (Section 2.3.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Calkins, et al. 1995 7709949 (1)	Aim: Identify +quantitate symptoms assoc. with VVS, AVB, or VT  Study type: Prospective  Size: n=80 pts (16 AVB,32 VT, 32 VVS)	Inclusion criteria: 80 pts with established hx of VT, VVS, or AVB  Exclusion criteria: N/A	Results: Features suggestive of AVB or VT  • Male gender  • Age >54  • <2 episodes of syncope  Features suggestive of VVS  • Before syncope: blurred vision, nausea, diaphoresis, palpitations  • After syncope: nausea, warmth, diaphoresis, fatigue	Clinical history is of value in distinguishing pts with these 3 causes of syncope
Alboni P, et al. 2001 11401133 (2)	Aim: Establish the historical findings predictive of the cause of syncope  Study type: Prospective study  Size: n=341 pts analyzed Cardiac cause 78 (23%) VVS 199 (58%) Neuro/Psych 4 (1%) Unexplained 60 (18%)	Inclusion criteria: Pts with syncope  Exclusion criteria: N/A	Results: Only heart disease was an independent predictor of a cardiac cause of syncope (sensitivity: 95%; specificity: 45%)	Absence of heart disease allowed an exclusion of a cardiac cause in 97%
Alboni P, et al. 2004 14697727 (3)	Aim: Establish the clinical features of VVS  Study type: Prospective Study  Size: n=461 pts prospectively evaluated. 280 had VVS:  Typical VVS n=39  HUTT induced n=142  Complex (CSH+VVS) n=31	Inclusion criteria: Pts with syncope  Exclusion criteria: N/A	Results: VVS differed from other neutrally mediated syncopes in precipitating factors and clinical features, including lower age and prevalence of organic heart disease, higher prevalence and duration of prodrome, Low prevalence of trauma	Considerable overlap between different Neurally medicated syndromes

Sheldon, et al. 2006 <u>16223744</u> (4)	Aim: Establish historical criteria for diagnosis of VVS  Study type: Prospective, used a Questioner of 118 items	Inclusion criteria: Pts with syncope and no apparent structural heart disease  Exclusion criteria: N/A	Results: The point score correctly classified 90 % of pts with an 89% sensitivity and 91 % specificity	The point scoring system can distinguish VVS from other causes of syncope with a high sensitivity and specificity
	Size: n=418 pts 235 syncope and positive HUTT n=95 no apparent cause (-HUTT) n=88 pts secondary syncope n=42 pts with CHB n=21 pts with SVT n=6 pts with VT n=5 pts with AS			
Sheldon, et al. 2002 12103268 (5)	Aim: Develop criteria that distinguish syncope due to VT from VVS in pts with SHD      Study type: Prospective analysis      Size: n=671 pts with a history of TLOC completed a 118 item historical questionnaire	Inclusion criteria: Pts with syncope and SHD  Exclusion criteria: N/A	Results:  Cause of TLOC known in 539 pts Seizures in 102 pts: Complex partial in 50 pts; Primary Generalized in 52 pts Syncope in 437 pts: VVS in 267 pts; VT in 90 pts; Other in 80 pts	The point score based on symptoms alone correctly classified 94% of pts, diagnosing seizures with a 94% sensitivity and 94% specificity
FAST Van Dijk, et al. 2008 <u>17916139</u> (6)	Aim: Assess yield and accuracy of an initial evaluation using: History, PE, and ECG  Study type: Prospective analysis then a 2 y follow-up by an expert committee  Size: n=503 pts (with a 2 y follow-up in 99%)	Inclusion criteria: Adults presenting with TLOC to the Academic Medical Center Amsterdam between February 2000 and May 2002  Exclusion criteria: N/A	Results: At initial evaluation:  • 119 pts (24%) certain diagnosis  • 199 pts (40%) had a highly likely diagnosis  • Overall diagnostic accuracy was 88%	Attending physicians can make a diagnosis in 63% of pts with TLOC, with a diagnostic accuracy of 88%

Romme, et al. 2009 19687157	Aim: Evaluate the Calgary Syncope Symptom Score	Inclusion criteria: Pts with TLOC Exclusion criteria: N/A	Results: Sensitivity of Calgary score was 87% but the Specificity was 32%	Sensitivity of the Calgary score similar to original study but the specificity less
Sheldon, et al. 2010	Study type: Prospective trial  Size: n=380 pts with TLOC: 237 pts (55%) were diagnosed with VVS using Calgary Score and then compared after 2 y of follow-up  Aim: Evaluate evidence based criteria to distinguish syncope due to VT from	Inclusion criteria: Pts with syncope and SHD	Results: • 21 pts with HUTT+VVS	Factors predicting VT were male gender and >35 y of age
<u>20586825</u> (8)	VVS in pts with structural heart disease  Study type: Prospective. 118 item questionnaire and an invasive and non-invasive diagnostic assessment  Size: n=134 pts	Exclusion criteria: N/A	<ul> <li>78 pts with clinical or EPS Induced VT</li> <li>35 pts with no cause identified</li> </ul>	<ul> <li>Factors predicting VVS were Prolonged sitting or standing, presyncope preceded by stress, headaches and fatigue after syncope lasting &gt;1 min</li> <li>The point score identified 92% of pts correctly, diagnosing VT with 99% sensitivity and 68% specificity, negative predictive value of &gt;96%</li> </ul>
PLOS Berecki-Gisolf, et al. 2013 24223233 (9)	Aim: Develop a model for symptoms that associate with cardiac causes of syncope  Study type: Literature based review  Size: n=7 studies	Inclusion criteria:  • 2 Pubmed searches using the following key words:  1. Diagnosis; signs and symptoms; vasovagal syncope  2. Clinical history; diagnosis; syncope  • Pts with ≥1 transient loss of consciousness  • A diagnosis of cardiac syncope vs. other causes  • Degree of evidence accepted in each paper  • Studies reporting ≥2 predictors of cardiac syncope  Exclusion criteria: N/A	Results: A total of 10 variables were found associated with cardiac syncope:  1. Age >60 y 2. Male gender 3. Structural heart disease 4. Low number of spells 5. Brief or absent prodrome 6. Supine syncope 7. Effort syncope 8. Absence of nausea 9. Absence of blurred vision	A model with 5 variables was as effective with moderate accuracy:  • >60 y of age  • Male gender  • Structural heart disease  • Low number of spells  • Lack of prodromal symptoms

#### Data Supplement 2. Nonrandomized Trials, Observational Studies, and/or Registries of Electrocardiography – (Section 2.3.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Recchia D, et al. 1995 <u>8770716</u> (10)	Study type: Retrospective observational  Size: n=128 pts	Inclusion criteria: All pts admitted to hospital due to syncope	1° endpoint: frequency of use ofechocardiogram to evaluate pts admitted with syncope	Hx, physical and ECG provided information to diagnosis a cause of syncope in 77% of pts (33 of 48 pts for whom a cause of syncope was felt to be ultimately determined)
		Exclusion criteria: Pts with syncope with known cause, pts with near syncope, vertigo, seizure, or pts referred to EP testing	Results: 90% of pts underwent cardiac testing; 64% of pts had echocardiogram which did not help elucidate cause of syncope, and echocardiogram; the ECG was normal for 52% of pts	<ul> <li>For pts with suspected cardiac disease, echocardiogram confirmed suspected diagnosis for 48% and ruled out suspected cause for remaining 52%.</li> </ul>
Perez-Rodon J, et al. 2014 24993462 (11)	Study type: Multicenter, prospective, observational  Size: n=524 pts	Inclusion criteria: Pts with syncope, readable ECG and 12 mo f/u  Exclusion criteria: N/A	1° endpoint: Mortality  Results: 344 pts (65.6%) had abnormal ECG, 33 pts (6.3%) died during f/u. AF OR: 6.8; 95% CI: 1.5–26.3 p=0.011. Ventricular pacing: OR: 21.8; 95% CI:4.1–115.3, p=0.001. left ventricular hypertrophy ECG criteria OR: 6.3; 95% CI:1.5–26.3; p=0.011. Intraventricular conduction disturbances OR: 3.8; 95% CI: 1.7–8.3; p=0.001	Only the presence of AF, intraventricular conduction disturbances, left ventricular hypertrophy ECG and ventricular pacing is associated with 1 y all cause mortality

#### Data Supplement 3. Nonrandomized Trials, Observational Studies, and/or Registries of Risk Stratification - Short-Term Outcomes - (Section 2.3.3)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
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Grossman SA, et al. 2012 22981659 (12)	Study type: Prospective observational  Size: n=244 ED pts with presyncope	Inclusion criteria: Presyncope, >18 y of age  Exclusion criteria: None	1° endpoint: Adverse outcomes ( death, cardiac arrest, pulmonary embolus, stroke, severe infection/sepsis, ventricular dysrhythmia, atrial dysrhythmia (including SVT and AF with rapid ventricular response), intracranial bleed, hemorrhage, MI, CHF, acute renal failure, or life-threatening sequelae of syncope (i.e., rhabdomyolysis, long bone or cervical spine fractures)  Results: 11 pts admitted with 49 adverse outcomes. If BSC had been followed 41 additional pts admitted and 34 pts discharged.	If BSC had been followed strictly, another 41 pts with risk factors would have been admitted and 34 discharged, a 3% increase in admission rate. However, using the modified criteria, only 68 pts would have required admission, a 38% reduction in admission, with no missed adverse outcomes on follow-up.
Colivicchi F, et al. 2003 12727148 (13)	Study type: Prospective observational  Size: Derivation cohort n=270 pts, Validation cohort n=328 pts	Inclusion criteria: Pts >12 y of age presenting for syncope to one of 6 ED's  Exclusion criteria: Seizure, presyncope, dizziness, vertigo	1° endpoint: 1 y all-cause mortality  Results: Primary outcome occurred in 31 (11.5%) pts in derivation cohort and 28 (8.5% in the validation cohort. "OESIL" score predictors include pts >65 y of age; Hx of CV disease; no prodrome; abnormal ECG	No "OESIL" risk factors associated with 0%, 1 y mortality, may identify low-risk subgroup that can be discharged Quantitative, attempts at reproducing difficult
Costantino G, et al. 2014 24862309 (14)	Study type: Patient level meta-analysis  Size: n=3,681 pts	Inclusion criteria: Patient level data from 6 prospective observational studies  Exclusion criteria: N/A	1º endpoint: 30 d combined death, arrhythmia, severe outflow tract obstruction, MI, CPR, pulmonary embolism, aortic dissection, hemorrhage, syncope resulting in major trauma  Results: "OESIL", "SFSR," "EGSYS" risk scores had similar sensitivity and specificity as clinical judgment.	Unclear whether these specific risk scores add value to clinical evaluation     Value of risk scores, etiology important to consider
Costantino G, et al. 2008 18206736 (15)	Study type: Prospective observational  Size: n=676 pts	Inclusion criteria: >18 y of age presenting to one of 4 ED's  Exclusion criteria: Dangerous condition identified in ED; head injury as cause of loss of consciousness; nonspontaneous return to consciousness; light-headedness, vertigo, coma, shock, seizure; terminal illness; substance abuse;	1° endpoint: 10 d combined death, CPR, pacemaker/ICD placement, ICU admission, acute anti-arrhythmic therapy, readmission  Results: Predictors of short-term outcomes (n=41 pts; 6.1%) included abnormal ECG, concomitant trauma, no prodrome, and male gender.	These criteria may identify pts who might benefit from hospital admission

		refusal to provide consent		
D'Ascenzo F, et al. 2013 22192287 (16)	Study type: Pooled meta- analysis  Size: n=11 studies	Inclusion criteria: Presentation of syncope to an ED  Exclusion criteria: N/A	1º endpoint: Combined death,     hospitalization/intervention related to arrhythmia,     ischemic heart disease, or VHD.      Results: Strongest predictors of an adverse     outcome included palpitations preceding syncope,     exertional syncope, history of HF or ischemic heart     disease, evidence of bleeding	These criteria may identify pts who might benefit from hospital admission
Da Costa A, et al. 2006 <u>15975670</u> (17)	Study type: Prospective observational  Size: n=305 pts	Inclusion criteria: Normal EPS after first onset of syncope or near-syncope  Exclusion criteria: None	1º endpoint: Combined symptomatic AV block, conduction abnormalities requiring pacemaker therapy, sustained ventricular arrhythmia, sudden death  Results: ECG is only independent predictor of long term adverse events	5% event rate at 2.5 y; normal EPS does not rule out dangerous conduction problems as cause of syncope
Del Rosso A, et al. 2008 <u>18519550</u> (18)	Study type: Prospective observational  Size: Derivation n=260 pts, validation n=256 pts	Inclusion criteria: Presentation of unexplained syncope to one of 14 ED's  Exclusion criteria: None	1° endpoint: Cardiac cause of syncope  Results: "EGSYS" risk score predictors include palpitations prior to syncope (+4), heart disease and/ or abnormal ECG (+3), exertional syncope (+3), supine syncope (+2), precipitating factors (-1), autonomic prodrome (-1)	• Risk of cardiac cause is <3% if EGSYS score <3, and >17% if EGSYS score ≥3
Derose S, et al. 2012 22594351 (19)	Study type: Retrospective observational  Size: n=22,189 pts	Inclusion criteria: Primary ED diagnosis of syncope or near-syncope in an integrated health system  Exclusion criteria: None	1° endpoint: 30 d mortality  Results: Predictors of short term mortality included increasing age, male gender, recent visit for syncope, history of HF, DM, seizure, and dementia	Pts without history of HF and <60 y of age had less than 0.2% risk of 30 d mortality
Dipaola F, et al. 2010 <u>20466221</u> (20)	Study type: Prospective observational  Size: n=488 pts	Inclusion criteria: >18 y of age presenting to one of 2 EDs with syncope of unknown cause  Exclusion criteria: None	1° endpoint: 10 d combined death, CPR, pacemaker/ ICD placement, ICU admission, acute anti-arrhythmic therapy, readmission  Results: Compared to the "OESIL" and "SFSR" risk scores, unstructured clinical judgment had similar sensitivity and higher specificity.	Unclear whether these specific risk scores add value to clinical evaluation

Exposito V, et al. 2013 23478089 (21)	Study type: Prospective observational  Size: n=180 pts	Inclusion criteria: >60 y of age with suspected VVS and undergoing tilt test  Exclusion criteria: None	1° endpoint: Positive tilt test  Results: CSSS score ≥ -2 has sensitivity of 50% and specificity of 73%	Calgary Syncope Symptom score for VVS has lower sensitivity and specificity in elderly population than previously reported
Gabayan G, et al. 2010 <u>20102895</u> (22)	Study type: Retrospective observational  Size: n=35,330 pts	Inclusion criteria: Primary ED diagnosis of syncope or near-syncope in an integrated health system  Exclusion criteria: None	1º endpoint: 7 d death, hospitalization, or procedure related to ischemic heart disease, VHD, or arrhythmia  Results: Predictors included >60 y of age, male gender, Hx of HF, ischemic heart disease, arrhythmia, and VHD.	Increasing age and presence of cardiac co- morbidities is associated with short term serious cardiac outcomes.
Grossman S, et al. 2007 17976548 (23)	Study type: Prospective observational  Size: n=362 pts	Inclusion criteria: >18 y of age presenting to an ED with syncope  Exclusion criteria: None	1º endpoint: 30 d pacemaker/ ICD placement, PCI, cardiac urgery, blood transfusion, CPR, change in anti-arrhythmic therapy, death, pulmonary embolus, stroke, sepsis, arrhythmia, intranial bleed, MI  Results: Low risk pts (<3% event rate) had none of the following: 1. suspicion for ACS; 2. signs of conduction disease; 3. worrisome cardiac history; 4. VHD; 5. family Hx of sudden death; 6. persistent abnormal vital signs in ED; 7. volume depletion; 8. primary central nervous system event	These criteria may identify low risk pts for whom discharge can be considered
Kayayurt K, et al. 2012 22520447 (24)	Study type: Prospective observational  Size: n=231 pts	Inclusion criteria: >18 y of age presenting to one of 2 ED's with syncope of unknown cause  Exclusion criteria: None	1º endpoint: 7 d rehospitalization, death, CPR, pacemaker/ ICD implantation, ICU admission, antiarrhythmic therapy  Results: The "ASR" risk score includes dyspnea (+1), OH (+1), precipitating cause for syncope (+1), pts >58 y of age (+1), Hx of CHF (+1), abnormal ECG (+2). ASR at a cut-point of >2 appears to similar test characteristics as the "OESIL,", "SFSR," and "EGSYS" risk scores.	These criteria may identify pts who might benefit from hospital admission
Martin T, et al. 1997 <u>9095005</u> (25)	Study type: Prospective observational  Size: Derivation n=252, validation n=374l	Inclusion criteria: Presentation of syncope to an ED  Exclusion criteria: None	1º endpoint: 1 y mortality or arrhythmia  Results: Predictors include abnormal ECG, Hx of ventricular arrhythmia, >45 y of age, Hx of CHF. Pts without any of these risk factors had <8% risk of the	These criteria may identify pts who might benefit from hospital admission or close outpatient follow-up.

			outcome.	
Moazez F, et al. 1991 <u>1985382</u> (26)	Study type: Prospective observational  Size: n=91 pts	Inclusion criteria: Syncope of unknown origin referred for EPS  Exclusion criteria: None	1º endpoint: Inducible sustained monomorphic VT     Results: Risk factors included abnormal signal averaged ECG; abnormal LVEF; prior sustained monomorphic VT	These criteria may be used to identify pts who might benefit from EPS.
Numeroso F, et al. 2010 20515909 (27)	Study type: Retrospective observational  Size: n=200 pts	Inclusion criteria: >18 y of age hospitalized for syncope  Exclusion criteria: None	1° endpoint: Cardiac cause of syncope  Results: OESIL score <2 had NPV of 98% to exclude cardiogenic cause. Prior syncope episodes and lack of prodrome were associated with increased risk of cardiogenic cause.	These criteria may identify pts who might benefit from hospital admission
Oh J, et al. 1999 10030311 (28)	Study type: Prospective observational  Size: n=275 pts	Inclusion criteria: >18 y of age with syncope of unknown origin after initial evaluation  Exclusion criteria: None	1° endpoint: Arrhythmic syncope  Results: Risk factors included absence of nausea/vomiting prior to syncope, and ECG abnormalities	These criteria may identify pts requiring who might benefit from cardiac monitoring
Quinn J, et al. 2004 14747812 (29)	Study type: Prospective observational  Size: n=684 visits	Inclusion criteria: Presentation of syncope to an ED  Exclusion criteria: None	1º endpoint: 7 d combined death, MI, arrhythmia, pulmonary embolism, stroke, subarachnoid hemorrhage, significant hemorrhage, or any condition causing a return ED visit and hospitalization for a related event  Results: In this derivation study, "SFSR" risk score	Using a cutpoint of 0 risk scores, the "SFSR" risk score has 96% sensitivity and 62% specificity. Use of the "SFSR" in the derivation cohort may have reduced hospitalizations by 10%.
			predictors include abnormal ECG, shortness of breath, hematocrit <30%, SBP <90 mmHg, Hx of CHF.	
Quinn J, et al. 2006 <u>16631985</u> (30)	Study type: Prospective observational  Size: n=791 consecutive visits	Inclusion criteria: Presentation of syncope to an ED  Exclusion criteria: None	1° endpoint: 30 d combined death, MI, arrhythmia, pulmonary embolism, stroke, subarachnoid hemorrhage, significant hemorrhage, or any condition causing a return ED visit and hospitalization for a related event	Application of the "SFSR" risk score may have decreased hospitalizations by 7%
			Results: In this validation cohort, the "SFSR" risk score was 98% sensitive and 56% specific.	

Reed M, et al. 2010 20170806 (31)	Study type: Prospective observational  Size: n=550 pts	Inclusion criteria: >16 y of age presenting with syncope to an ED  Exclusion criteria: None	1° endpoint: 30 d combined acute MI, dangerous arrhythmia, pacemaker/ICD placement, pulmonary embolus, neurologic event, hemorrhage requiring transfusion, emergent surgical or endoscopic procedure  Results: The "ROSE" risk score predictors include BNP≥300, bradycardia ≤ 50, rectal exam with fecal occult blood, hemoglobin ≤ 90 g/l, chest pain, ECG with q waves, oxygen saturation ≤ 94% on room air. The validation cohort demonstrated sensitivity of 87% and specificity of 66%.	These criteria may identify pts who might benefit from hospital admission
Ruwald M, et al. 2013 <u>23450502</u> (32)	Study type: Retrospective registry  Size: n=37,705 pts	Inclusion criteria: Discharged from an ED with first time diagnosis of syncope  Exclusion criteria: None	1° endpoint: All-cause mortality  Results: The CHADS2 score (HF [+1], hypertension [+1], age ≥75 [+1], DM [+1], prior TIA/ stroke [+2] associated with all-cause mortality.	CHADS2=0 is associated with 1.5% 1 y mortality rate
Saccilotto R, et al. 2011 21948723 (33)	Study type: Meta-analysis  Size: n=12 studies; n=5,316 pts	Inclusion criteria: External validation study of "SFSR" risk score  Exclusion criteria: N/A	1º endpoint: Combined serious outcomes, definition varied by specific study  Results: The "SFSR" risk score has a pooled sensitivity of 87% and specificity of 52%. Significant between-study heterogeneity was observed	"SFSR" risk score appears to be less sensitive and specific in external validation studies than originally reported
Sarasin F, et al. 2003 <u>14644781</u> (34)	Study type: Prospective observational  Size: n=175 pts cohort to develop and cross-validate the risk score; 269 pts cohort to validate the system	Inclusion criteria: Unexplained syncope after ED evaluation  Exclusion criteria: None	1° endpoint: Arrhythmic syncope  Results: Predictors include abnormal ECG, Hx of CHF, ≥65 y of age	Pts without any risk factors had <2% risk of arrhythmic syncope
Serrano L, et al. 2010 <u>20868906</u> (35)	Study type: Meta-analysis Size: n=18 eligible studies	Inclusion criteria: ED cohort study of syncope/ near-syncope study for risk score derivation or validation  Exclusion criteria: N/A	1° endpoint: Combined serious outcomes, definition varied by specific study  Results: The "OESIL" risk score has a pooled sensitivity of 95% and specificity of 31%. The "SFSR" risk score has a pooled sensitivity of 86% and specificity of 49%. Large variations were noted in methodological quality of studies.	These criteria may identify pts who might benefit from hospital admission

Sheldon R, et al. 2006 16223744 (4)	Study type: Prospective observational Size: n=418 pts	Inclusion criteria: Prior episode of syncope evaluated in cardiology clinic or hospital cardiology wards  Exclusion criteria: None	Positive tilt test  Results: CSSS risk score predictors include: any of bifasciular block, asystole, SVT, DM (-5); blue color at time of event (-4); age at first syncope ≥35 (-3), intact memory of event (-2); presyncope/ syncope with standing (+1); sweating/ warm feeling before episode (+2); episode associated with pain or procedure (+3)	CSSS ≥-2 has sensitivity of 89% and specificity of 91% for identifying tilt-positive syncope
Sule S, et al. 2012 22878409 (36)	Study type: Prospective observational  Size: n=242 consecutive pts	Inclusion criteria: Hospitalized for syncope  Exclusion criteria: None	1° endpoint: Mortality  Results: Predictors included unexplained etiology, SFSR risk score, lack of hypertension, GFR (higher value reduces risk)	These criteria may identify pts who might benefit from hospital admission
Sun B, et al. 2009 <u>19766355</u> (37)	Study type: Retrospective observational  Size: n=2,871 pts	Inclusion criteria: >60 y of age with unexplained syncope or near-syncope after ED evaluation  Exclusion criteria: None	1° endpoint: 30 d combined death, arrhythmia, MI, new diagnosis of severe SHD, pulmonary embolism, aortic dissection, stroke/TIA, cerebral hemorrhage, significant anemia requiring blood transfusion  Results: Risk predictors include age >90 (+1), male gender (+1), history of arrhythmia (+1), triage SBP >160 mmHg (+1), abnormal ECG (+1), abnormal troponin result (+1), complaint of near syncope (-1). Score of <1 was associated with 2.5% event rate	These criteria may identify pts who might benefit from hospital admission

## Data Supplement 4. Nonrandomized Trials, Observational Studies, and/or Registries of Risk Stratification - Long-Term Outcomes - (Section 2.3.3)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Numeroso F, et al.	Study type: Prospective	Inclusion criteria: ED syncope	1° endpoint: Recurrent syncope, trauma, major	◆ N/A
2014	observational		procedures, CV events, death	
<u>24489075</u>		Exclusion criteria: None		
(38)	Size: n=200 consecutive pts		Results: Any heart disease not associated with	
			endpoints, but high risk heart disease (CAD, CHF,	
			AS, cardiomyopathies, primary arrhythmic	
			diseases) was.	

Ungar A, et al. 2010 20167743 (39)	Study type: Prospective observational Size: n=380 pts	Inclusion criteria: ED syncope  Exclusion criteria: None	1º endpoint: Death  Results: Predictors for recurrent syncope were prodromes and palpitations prior to syncope	Incidence of syncope recurrence not related to mechanism of syncope or EGSYS score < or ≥3.
Sule S, et al. 2011 <u>21259276</u> (40)	Study type: Observational Size: n=325 pts	Inclusion criteria: Hospitalized for syncope  Exclusion criteria: None	1° endpoint: Recurrent syncope  Results: Associated with recurrent hospitalized syncope were DM, AF and smoking	Syncope etiology found in 74%
Sumner G, et al. 2010 <u>20662990</u> (41)	Study type: Observational Size: n=208 pts	Inclusion criteria: NCS with Positive tilt and > lifetime syncope  Exclusion criteria: None	1° endpoint: Recurrent syncope  Results: Number of syncope in prior y better predicted syncope recurrence compared to lifetime syncope episodes	• Syncope recurred in 22% of those with <2 episodes in the prior y compared to 69% in those with >6 episodes.
Koechl B, et al. 2012 <u>22722821</u> (42)	Study type: Observational Size: n=242 pts	Inclusion criteria: Syncope  Exclusion criteria: None	1° endpoint: Recurrent syncope  Results: Increased syncope with age and disability	Syncope recurrence was 32.5%
Khera S, et al. 2013 <u>23332735</u> (43)	Study type: Observational retrospective  Size: n=352 pts	Inclusion criteria: ED syncope  Exclusion criteria: None	1° endpoint: Admission for syncope  Results: 3% readmitted; CHF and ACS were risk factors	Etiology of syncope found in 69%
Sorajja D, et al. 2009 <u>19720940</u> (44)	Study type: Case control  Size: n=3877 pts with syncope; of which 9.8% had syncope while driving	Inclusion criteria: Syncope  Exclusion criteria: None	1° endpoint: Syncope while driving in followup  Results: In the syncope while driving group (n=381 pts) 72 pts had recurrent syncope, including 10 while driving.	Etiology of syncope while driving included neutrally mediated (37%) and arrhythmic (12%)
Lee S, et al. 2014 <u>25402339</u> (45)	Study type: Observational Size: n=289 pts	Inclusion criteria: Syncope  Exclusion criteria: None	1º endpoint: Recurrent syncope  Results: 6.6% with recurrent syncope in 1 y.  Syncope more common in those with ≥6 prior episodes and unexplained syncope	Etiology of initial syncope 63%     NMS, 12% OH, 12% cardiac, 12%     unexplained

Ruwald MH, et al.	Study type: Nationwide	Inclusion criteria: Syncope	1° endpoint: Recurrent syncope	• N/A	
2013	administrative registries				
<u>24035171</u>		Exclusion criteria: None	<b>Results:</b> Predictors of recurrent syncope include:		
(46)	Size: n=5141 pts >85 y of age		AS, kidney disease, AV or LBBB, Male, COPD,		
	n=23,454 <85 y of age		CHF, AF, Age, orthostatic medications		
	· -			•	

## Data Supplement 5. Nonrandomized Trials, Observational Studies, and/or Registries of Disposition After Initial Evaluation – (Section 2.3.4)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and 95% CI)	Summary/Conclusion Comment(s)
Sun, et al. 2012 <u>22687184</u> (47)	Aim: Create standardized reporting guidelines, including serious outcomes, for syncope research  Study type: Expert consensus  Size: n=24 panelists	Inclusion criteria: Convenience sample of 24 panelists with clinical or methodological expertise relevant to syncope research	N/A  Results: Modified Delphi consensus process identified final guideline elements from 183 candidate elements	23 serious conditions identified for research reporting
Daccarett, et al. 2011 21757485 (48)	Study type: Retrospective observational  Size: n= 254 pts	Inclusion criteria: ED visit for syncope identified by ICD code 780.2  Exclusion criteria: Pts with secondary diagnosis of syncope	1º endpoint: Admission rate  Results: Retrospective application of the Utah Faint-Algorithm would have reduced admissions by 52%. Algorithm explicitly defined conditions or high risk criteria for which admission would be indicated. The 7-d serious event rate in pts who should have been discharged per the algorithm (3%) was similar to those who were actually discharged (4%).	A standardized evaluation algorithm that explicitly identified serious conditions which requires admission appears to be safe and reduces resource use.
Framingham Cohort Study Soteriades, et al. 2002 12239256 (49)	Aim: Describe prognosis of syncope in general population  Study type: Prospective cohort	Inclusion criteria: Participants in the original Framingham Heart Study and the Framingham Offspring Study  Exclusion criteria: N/A	All-cause mortality Over 25 y follow-up period, pts with presumptive VVS had similar riskadjusted mortality risk as pts without syncope	Syncope of vasovagal etiology does not appear to increase mortality risk

	<u>Size</u> : n=7,814 pts			
Morag, et al. 2004 <u>15498613</u> (50)	Aim: Assess diagnostic benefit of admission for unexplained syncope  Study type: Prospective cohort  Size: n=45 pts	Inclusion criteria: ED visit for syncope, undergoing structured evaluation, age ≥ 50  Exclusion criteria: Intoxicated with drugs or alcohol, had antecedent head trauma prompting symptoms, had witnessed seizure activity with a history of seizures, or if their loss of consciousness promptly responded to medical management (administration of glucose or naloxone)	1° endpoint: Life threatening event or significant therapeutic intervention  Results: Of 30 admitted pts, none experienced the primary endpoint as inpatient or at 30 d follow up	Yield of diagnostic admission appears to be low
Shiyovich, et al. 2008 18432020 (51)	Aim: Assess diagnostic evaluation, costs, and prognosis of pts admitted for syncope  Study type: Retrospective cohort  Size: n=376 pts	Inclusion criteria: Hospital admission for evaluation of syncope  Exclusion criteria: pre syncope, seizure, malignant arrhythmia	1º endpoint: Diagnostic evaluation, costs, 1 y mortality  Results: 38% had no clear diagnosis at discharge.	A significant proportion of pts have an unrevealing evaluation
Schillinger, et al. 2000 11098534 (52)	Aim: Assess evaluation and prognosis of pts admitted for syncope  Study type: Retrospective cohort  Size: n=127 pts	Inclusion criteria: Hospital admission for evaluation of syncope  Exclusion criteria: Not admitted after ED evaluation	1° endpoint:  No patient has inpatient death or recurrent syncope as inpatient. 2% of pts died within 30 days, all from known pre-existing disease  Results: Of 376 pts, 48% had no clear diagnosis at discharge. Long term mortality was higher for pts with cardiac and neurologic etiology.	Hospital evaluation had modest diagnostic yield; population had low short term mortality risk.
Ungar, et al. 2015 <u>25976905</u> (53)	Study type: Observational Size: n= 362 pts	Inclusion criteria: ED evaluation for TLOC  Exclusion criteria: N/A	1° endpoint: Disposition  Results: Disposition included 29% admitted; 20%	Presence of ED observation unit and hospital based syncope unit is associated with lower hospitalization rates compared to historical experience

			ED observation unit; 20% referred to hospital based syncope unit; 31% discharged. No 1 y death after evaluation in any setting appeared to be related to TLOC	
Shin, et al. 2013	Study type: Quasi experimental, pre-post w/o control, assess	Inclusion criteria: >18 y of age with syncope evaluated in ED	1° endpoint: Admission rate	<ul> <li>Standardized evaluation, including risk stratification and use of an observation unit,</li> </ul>
<u>23918559</u>	implementation of standard			reduced admissions, costs, and LOS
(54)	approach including risk	Exclusion criteria: inability to	Results: In the 1-y post-period	
	stratification, hospital order set, and ED observation unit	consent, prior enrollment in other	compared to the 1-y pre- period, there	
	ED observation unit	studies, non-syncope syndromes	were reductions in admissions (8.3%), costs (30%), and LOS (35%)	
	<u>Size</u> : n= 244 pts		60313 (30 /0), and £00 (30 /0)	

## Data Supplement 6. RCTs for Disposition After Initial Evaluation – Serious Conditions – (Section 2.3.4)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
SEEDS Shen, et al. 2004 15536093 (55)	Aim: Assess whether designated syncope unit in ED improves diagnostic yield and reduces admission  Study type: 1 site RCT  Size: n=103 pts	Inclusion criteria: Syncope of undetermined cause after ED evaluation, AND intermediate risk by semistructured criteria  Exclusion criteria: 1. Identified cause of syncope; 2. Dangerous condition requiring admission; 3. Nonsyncope syndrome such as light-headedness	Intervention: Syncope unit: continuous cardiac monitoring up to 6 h; hourly VS/ orthostatic BP; ECG for abnormal heart sounds or ECG; recommended tilt-table testing for selected pts; outpatient EP consult, echocardiogram, tilt-table testing available within 72 h after discharge  Comparator: Standard care (default was admission to hospital)	1° endpoint: Admission rate: 43% in intervention, 98% in control  1° Safety endpoint (if relevant): No differences in survival or recurrent syncope	Hospital d: 64 in intervention, 140 in control     Presumptive diagnosis: 67% in intervention, 10% in control      Summary:     Structured syncope unit in ED reduced hospital admission and length of stay without affecting mortality or recurrent syncope rates.

EDOSP	Aim: Assess whether	Inclusion criteria: Pts	Intervention:	1° endpoint:	Index hospital costs: \$629 less in EDOSP
Sun, et al.	EDOSP reduces resource	>50 y of age, AND	12–24 h of cardiac	LOS: 29 h in EDOSP, 47 h in	vs. control
2014	use without adversely	intermediate risk for	monitoring;	control	
<u>24239341</u>	affecting patient oriented	serious short-term events	echocardiogram for cardiac		
(56)	outcomes	by semi-structured	murmur; serial troponin	1° Safety endpoint (if	Summary:
		criteria		relevant):	EDOSP reduced resource use with no
	Study type:		<u>Comparator</u> : Admission to	No differences in 30 d serious	difference in outcomes, quality-of-life, or
	5-site RCT	Exclusion criteria: 1.	inpatient service	outcome rates, quality-of-life	patient satisfaction.
		Dangerous condition		scores, patient satisfaction	
	Size: n=124 pts	requiring admission; 2.			
		non-syncope syndrome			
		such as seizure			

## Data Supplement 7. Nonrandomized Trials, Observational Studies, and/or Registries Comparing Blood Testing – (Section 3.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type*; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Pfister R, et al. 2009 18237792 (57)	Aim: Determine NT-pro-BNP role in the differential diagnosis of pts with syncope.  Study type: Observational cohort	Inclusion criteria: Consecutive pts in the emergency room  Exclusion criteria: None	Intervention: None  Comparator: Between subsequently diagnosed groups	1° endpoint: NT-pro BNP levels in different etiology of syncope groups	<ul> <li>Post hoc determination of levels after diagnosis obtained</li> <li>No gold standard for most diagnostic categories</li> </ul>
Thiruganasambanda moorthy V, et al. 2015 26498335 (58)	Size: n=61 pts  Aim: Prognostic value of cardiac biomarkers in the risk stratification of syncope  Study type: Systematic review  Size: N/A	Inclusion criteria: Adult syncope pts during acute management  Exclusion criteria: Case reports, children	Intervention: None  Comparator: None	1° endpoint: MACE: death, CPR, MI, structural heart disease, PE, significant hemorrhage, cardiac intervention. High sensitivity Troponin and natriuretic peptides showed good sensitivity and specificity for MACE	Relationship of syncope to MACE and biomarkers is unclear

<b>GESINUR</b> Pérez-Rodon, et al. 2014 24993462 (11)	Aim: Determine outcome predictors on resting ECG  Study type: Multicenter, prospective, retrospective observational cohort  Size: n=524 pts	Inclusion criteria: Syncope in the ER with 1 y follow-up	Intervention: None	1º endpoint: Mortality: 33 total deaths (6.6%), 1 SCD	Summary: AF, IVCD, LVH and ventricular pacing an independent risk factors for mortality
Chiu DT, et al. 2014 24698512 (59)	Aim: Determine the yield of standard diagnostic tests  Study type: Prospective, observational, cohort study of consecutive ED  Size: n=570 pts	Inclusion criteria: ER presentation syncope  Exclusion criteria: None	Intervention: None  Comparator: None	1º endpoint: Yield of 3 diagnostic tests in those pts that had the test (no structured indication for why tests were performed).  Safety endpoint (if relevant): None	Summary: Diagnosis in 73 pts (8%). Yield: echo 22%, telemetry 3%, troponin 3%.
<b>SYSTEMA</b> Fedorowski, et al. 2013 23510366 (60)	Aim: Determine role of biomarkers in pts with syncope  Study type: Observational cohort  Size: n=270 pts	Inclusion criteria: Unexplained syncope	Intervention: Tilt with CSM and biomarker analysis	1º endpoint: Levels of C-terminal pro-arginine vasopressin (CT-proAVP), C-terminal endothelin-1 precursor fragment (CT-proET-1), midregional fragments of pro-atrial natriuretic peptide (MR-proANP) and pro-adrenomedullin (MR-proADM)	Summary: Biomarkers divided into quartiles, CT-proET-1 and MR-proANP were associated with diagnoses of OH, carotid sinus hypersensitivity and VVS.
Reed, et al. 2012 <u>22962048</u> (61)	Aim: Assess whether plasma troponin concentration can predict 1 mo and 1 y serious outcome, or all-cause death  Study type: Prospective observational cohort  Size: n=261 pts	Inclusion criteria: Admitted pts with syncope	Intervention: None	1° endpoint: The proportion of pts with a composite serious outcome increased across pts stratified into quintiles based on peak troponin concentration at 1 mo (0%, 9%, 13%, 26%, 70%) and at 1 y (10%, 22%, 26%, 52%, 85%).	Summary: Troponin concentrations were above the limit of detection in 261 (77%) pts. Peak troponin concentration was associated with increasing risk of serious outcome and death, which increases with higher troponin concentrations.
Grossman, et al. 2003 14630890 (62)	Aim: Determine role of cardiac enzymes in elderly pts with syncope  Study type: Retrospective chart	Inclusion criteria: Consecutive pts 65 y of age and older with syncope in an urban teaching hospital ED	Intervention: None	1º endpoint: 3 of 141 pts, or 2.1% (95% CI: 0.04%–6.09%), had positive cardiac enzymes during their hospitalization (CPK, not Tpl study)	Summary: Author conclusion: Cardiac enzymes may be of little additional value if drawn routinely on elderly pts with syncope

	review Size: n= 319 pts				
Pfister, R, et al. 2009 18237792 (57)	Aim: determine NT-pro-BNP values between cardiac and non-cardiac syncope  Study type: Observational cohort  Size: n=61 pts	Inclusion criteria: ED syncope  Exclusion criteria: none	Intervention: None	1° endpoint: Pts with cardiac syncope had significantly higher NT-pro-BNP values (514 IQR 286–1154 pg/ml) than pts with noncardiac cause (182 IQR 70–378 pg/ml, p=0.001). NT-pro-BNP at a cut-off of 164 pg/ml identified pts with cardiac syncope with a sensitivity of 90% and 93.8%, a specificity of 48.8% and 46.7% and a negative predictive value of 91% and 95.5%	Summary: NT-pro-BNP assessment was helpful in differentiating cardiac from non-cardiac syncope
Goble MM, et al. 2008 18082784 (63)	Aim: To evaluate ED management of childhood syncope, focusing on diagnostic tests ordered  Study type: Retrospective chart review  Size: n=113 pts	Inclusion criteria: <18 y of age, pediatric ED syncope	Intervention: None	1º endpoint: Most commonly ordered tests in the ED in order of decreasing frequency were electrolytes (90%), ECG (85%), complete blood count (80%), urinalysis, urinary drug screen, or urinary human chorionic gonadotropin 76%, head CT, 58%, and chest x-ray 37%	<u>Summary</u> : Nearly 100% admitted because of automated or non-expert ECG interpretation, weak descriptive study.

## Data Supplement 8. Nonrandomized Trials, Observational Studies, and/or Registries of Blood Testing – (Section 3.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Tanimoto K, et al.	Study type:	Inclusion criteria: Pts with syncope	1° endpoint:	Limitations:
2003	Retrospective	. ,	To evaluate the feasibility of measuring BNP to	<ul> <li>Retrospective study, and "unknown"</li> </ul>
<u>14715356</u>	observational	Exclusion criteria: AF; renal failure; and	identify cardiac syncope.	causes could be cardiac
(64)		who died within 24 h after admission		
	Size: n=118 pts		Results:	Conclusions:

			BNP concentrations in the cardiac syncope group (118±42 pg/ml) were significantly higher than those with reflex-mediated, neurologic, or unknown causes of syncope (p<0.01).	Measurement of BNP concentrations may help confirm cardiac causes of syncope
			At a cut-off value of 40 pg/ml used to determine a cardiac cause of syncope, the	
			sensitivity and specificity identifying cardiac syncope were 82% and 92%, respectively	
Christ M, et al.	Study type: Prospective	Inclusion criteria: Consecutive pts	1° endpoint: Diagnostic and predictive value of	Limitations:
2015 25447619	observational	presenting to ED with syncope or near syncope	cTnThs in pts with syncope.	<ul> <li>Post hoc analysis of a single-center trial— not all syncopal pts had troponins.</li> </ul>
(65)	<b>Size</b> : n=360 pts		Results:	Possible bias in selecting pts for whom
		Exclusion criteria: Persistent altered mental status or illicit drug-related loss of	Cardiac syncope present in 22% of pts.	treating physicians ordered cTnThs
		consciousness; seizure; coma;	<ul> <li>Diagnostic accuracy for cTnThs levels AUC:</li> </ul>	Conclusions:
		hypoglycemia; transient loss of	(0.77; CI:0.72–0.83; p<0.001). Comparable AUC	<ul> <li>cTnThs levels show a limited diagnostic</li> </ul>
		consciousness caused by head injury; no	(0.78; CI:0.73–0.83; p<0.001) obtained for	and predictive accuracy for the identification
		phlebotomy or troponin	predictive value of cTnThs levels within 30 d.	of pts with syncope at high risk

#### Data Supplement 9. Nonrandomized Trials, Observational Studies, and/or Registries of Cardiac Imaging – (Section 3.2.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Chiu DT, et al. 2014 <u>24698512</u> (59)	Study type: Prospective observational  Size: n=570 pts presenting to ED with syncope	Inclusion criteria: ≥18 y of age with syncope  Exclusion criteria: Altered mental status; substance-induced LOC; seizure; coma; hypoglycemia; TLOC due to head trauma; near syncope	1º endpoint: Finding on diagnostic test (echocardiogram, troponin [suspected AMI], telemetry, ambulatory monitor) while inpatient or follow-up that identified etiology of syncope.  Results: 73 positive tests (12.8%) Echo: 33/150 (22%), telemetry: 19/330 (5.7%), ambulatory ECG: 2/56 (3.6%), troponin: 19/317 (6%)	Limitations: Single-center study; small sample; no long-term follow-up; kappa rarely >0.80.  Conclusions: Routing testing common, but diagnostic yield low, although they uncover significant causes of syncope.  Echo the highest yield (low LVEF most common etiology of syncope).
Recchia D, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:

1995	Retrospective	Admission for syncope	Frequency echocardiography used in evaluation of pts	Single-center study; small sample
<u>8770716</u>	observational		admitted because of syncope and to examine the	
(10)	<b>C:</b>	Exclusion criteria:	diagnostic information, over and above that provided	Conclusions:
	<u>Size</u> : n=128 pts	Syncope of a known cause, near- syncope or vertigo, clinically	by the initial H&P, and electrocardiography	<ul> <li>For pts without suspected cardiac disease after H&amp;P, and ECG, the echocardiogram did not appear</li> </ul>
		obvious seizure, or referred for	Results:	to provide additional useful information, suggesting
		ECG testing	Echocardiogram normal for 52% pts	that syncope alone may not be an indication for
		J T T T T T T T T T T T T T T T T T T T	Echocardiograms of pts with syncope and no clinical	echocardiography.
			evidence of heart disease by H&P, or	<ul> <li>For pts with suspected heart disease,</li> </ul>
			electrocardiography were normal (63%) or provided no	echocardiography served to confirm or refute the
			useful additional information for arriving at a diagnosis (37%).	suspicious in equal proportions.
			<ul> <li>Among pts for whom cardiac disease was suspected</li> </ul>	
			after H&P, or ECG, the echocardiogram confirmed the	
			suspected diagnosis for 48% and ruled out a	
			suspected diagnosis for the remaining 52%.	
			H&P, and initial ECG provided sufficient information     The partition of the provided for 37/48 ptg (77%)	
			to permit a diagnosis to be made for 37/48 pts (77%) for whom a cause of syncope was ultimately	
			determined.	
Sarasin FP, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
2002	Prospective	Adult pts (≥18 y of age) presenting	To study the role of echocardiography in the stepwise	Relatively small sample size of pts with SUO
<u>12231593</u>	observational	with chief complaint of syncope	evaluation of syncope	and/or arrhythmias.
(66)	Size: n=650 pts	Exclusion criteria:	Paguita	EPS not performed.
	<u>312e</u> . 11-030 pts	None specified	Results: Severe AS suspected in 20/61 pts with systolic	Conclusions:
		Trong specimen	murmur was suspected in 20 of these, confirmed in 8.	Echocardiography is useful for risk
			<ul> <li>In pts with unexplained syncope (n=155),</li> </ul>	stratification—by measuring LVEF, a predictor of
			echocardiography showed no abnormalities that	arrhythmias—only in pts with SUO and with a
			established cause of the syncope.	positive cardiac history, or abnormal ECG.
			Echocardiography was normal or non-relevant in all pts with a negative cardiac Hx and a normal ECG	
			(n=67).	
			● In pts with positive cardiac history or an abnormal	
			ECG (n=88), echocardiography showed LVEF ≤40%)	
			in 24 (27%) and minor non-relevant findings in	
			remaining 64.	
			<ul> <li>Arrhythmias were diagnosed in 12/24 pts with</li> </ul>	

			systolic dysfunction and in 12/64 remaining pts (19%)	
			(p<0.01).	
Probst MA, et al. 2015 25943042 (67)	Study type: Observational cohort  Size: n=3,500 pts	Inclusion criteria: ED visits where any of the 3 pts "reasons for visit" included: fainting (syncope); includes blacking out, passing out, fainting spells; excludes unconsciousness" from the ED portion of the National Hospital Ambulatory Medical Care Survey, 2001–2010  Exclusion criteria: None	1º endpoint: To identify temporal trends in syncope-related ED visits and associated trends in imaging, hospital admissions, and diagnostic frequencies.  Results: Admission rates for syncope pts ranged from 27%—35% and showed no significant downward trend (p=0.1). Advanced imaging rates increased from about 21% to 45% and showed a significant upward trend (p<0.001).	Limitations:  Registry study, potential for residual confounding, miscoding syncope diagnoses  Conclusions: Resource utilization associated with ED visits for syncope appears to have increased, with no apparent improvements in diagnostic yield for admissions
Mendu ML, et al. 2009 19636031 (68)	Study type: Observational cohort study  Size: n=2106 pts	Inclusion criteria: Pts ≥65 y of age admitted to an acute care hospital through ED (2002–2006), with an admission or discharge diagnosis of syncope.  Exclusion criteria: Pts in whom absence of loss of consciousness (e.g. near syncope) was documented were excluded.	1º endpoint:  To determine the frequency, yield, and costs of tests obtained to evaluate older persons with syncope; to calculate the cost per test yield and determined whether the SFSR improved test yield.  Results:  ECG (99%), telemetry (95%), cardiac enzymes (95%), and head CT (63%) were the most frequently obtained tests.  Cardiac enzymes, CTs, echocardiograms, carotid ultrasounds, and electroencephalography all affected diagnosis or management in <5% of cases and helped determine etiology of syncope <2% of the time.  Postural BP, performed in only 38% of episodes, had highest yield in affecting diagnosis (18–26%) or management (25–30%) and determining etiology of the syncopal episode (15–21%).  The cost per test affecting diagnosis or management was highest for electroencephalography (\$32,973), CT (\$24,881), and cardiac enzymes (\$22,397) and lowest for postural BP (\$17–\$20).  The yields and costs for cardiac tests were better among pts meeting, than not meeting, SFSR.	Limitations:  Retrospective diagnosis of database of a single-center, with potential for misclassification of diagnosis by ICD codes  No capturing of testing performed in pts not admitted through ED, or after hospitalization.  Conclusions:  Many unnecessary tests are obtained to evaluate syncope. Selecting tests based on Hx and examination and prioritizing less expensive and higher yield tests would ensure a more informed and cost-effective approach to evaluating older pts with syncope

## Data Supplement 10. Nonrandomized Trials, Observational Studies, and/or Registries of Stress Testing – (Section 3.2.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Woelfel, et al. 1983 <u>6875122</u> (69)	Study type: Small case series  Size: n=3 pts	Inclusion criteria:  1:1 AV conduction at rest developed fixed 2:1 or 3:1 AV block during treadmill exercise testing  Exclusion criteria: N/A	1º endpoint: Determine mechanism of high grade block during exertion.  Results: ■ 3 pts with 1:1 AV conduction at rest developed fixed 2:1 or 3:1 AV block during treadmill exercise testing. EPS documented block distal to the AV node in all 3 pts, and suggested that the exercise-induced block occurred because of increased atrial rate and abnormal refractoriness of the His-Purkinje conduction system.	Limitations:  • Small case series  Conclusions:  • High grade AV block appearing during exercise reflects conduction disease of the His-Purkinje system rather than of the AV node, even in the absence of BBB. Pts with this diagnosis should be considered for permanent cardiac pacing.
Kapoor WN , et al. 1983 <u>6866032</u> (70)	Study type: Prospective cohort  Size: n=204 pts	Inclusion criteria: Symptoms "comparable with syncope"  Exclusion criteria: Tonic-clonic movements; post-ictal state; aura	1° endpoint:  To determine how often a cause of syncope could be established and to define the prognosis of such pts.  Results:  A CV cause was established in 53 pts and a nonCV cause in 54. The cause remained unknown in 97 pts.  At 12 mo, the overall mortality was 14±2.5%.  The mortality rate (30±6.7%) in pts with a CV cause of syncope was significantly higher than the rate (12±4.4%) in pts with a nonCV cause (p=0.02) and the rate (6.4±2.8 %) in pts with syncope of unknown origin (p<0.0001).  The incidence of sudden death was 24±6.6 % in pts with a CV cause, as compared with 4±2.7 % in pts with a nonCV cause (p=0.005) and 3 ±1.8 % in pts with syncope of unknown origin (p=0.0002).	Limitations:  Descriptive study.  Conclusions: Cause of syncope is frequently not established. Pts with a CV cause have a higher incidence of sudden death than pts with a non-CV or unknown cause (VT and SSS most common).

## Data Supplement 11. RCTs Comparing Cardiac Monitoring – (Section 3.2.3)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Krahn, et al. 2001 <u>11435336</u> (71)	Aim: To compare ILR to conventional monitoring in SUO.  Study type: RCT, cross-over  Size: n=60 pts	Inclusion criteria: Recurrent SUO or a single episode of syncope associated with injury that warranted CV investigation.  Exclusion criteria: LVEF <35%; unlikely to survive for 1 y; unable to provide follow-up or give informed consent.	Intervention: ILR with one y of monitoring (n=30).  Comparator: "Conventional testing" with a 2 to 4 wk period of monitoring with an ELR, followed by tilt table, and EPS (n=30).	1º endpoint:  ■ Diagnosis achieved in 14/27 pts randomized to prolonged monitoring compared with 6/30 undergoing conventional testing (52% vs. 20%, p=0.012).  ■ Prolonged monitoring more likely to result in diagnosis than conventional testing (55% vs. 19%, p=0.0014).  ■ Bradycardia (sinus and AVB) detected in 14 pts undergoing monitoring compared with 3 pts undergoing conventional testing (40% vs. 8%, p=0.005).	Limitations:  Single center study, with possible selection bias due to assessment of older population without structural heart disease (excluding pts with a high pretest probability of neurally mediated syncope or VA).  Conclusions:  A prolonged monitoring strategy is more likely to provide a diagnosis than conventional testing in pts with unexplained syncope.  Bradyarrhythmias are a frequent cause of syncope
Krahn AD, et al. 2003 12906979 (72)	Aim: To compare costeffectiveness of ILR to conventional testing.  Study type: RCT, crossover  Size: n=60 pts	Inclusion criteria: Recurrent SUO or a single episode of syncope associated with injury that warranted CV investigation.  Exclusion criteria: LVEF <35%; unlikely to survive for 1y; unable to provide follow-up or give informed consent.	Intervention: ILR with one y of monitoring (n=30).  Comparator: "Conventional testing" with a 2 to 4 wk period of monitoring with ELR, followed by tilt table, and EPS (n=30).	1º endpoint:  ●14/30 pts monitored diagnosed at \$2,731±\$285/pts, \$5,852±\$610/diagnosis, compared with 6/30 conventional pts diagnosed (20% vs. 47%, p=0.029), at a \$1,683±\$505/pts (p<0.0001) and \$8,414±\$2,527/diagnosis (p<0.0001).	Limitations:  ■ Single center study, with possible selection bias due to assessment of older population without structural heart disease (excluding pts with a high pretest probability of neurally mediated syncope or ventricular arrhythmia). Canadian dollars used.  Summary:  ■ A strategy of primary monitoring is more cost-effective than conventional testing in establishing a diagnosis in recurrent SUO.
Farwell, et al. 2006 <u>16314338</u> (73)	Aim: To investigate the impact of ILR on pts with recurrent SUO.	Inclusion criteria: Consecutive pts presenting to single center, ≥16 y of age; acute syncope presentation; ≥2 SUO in 12 mo; no	Intervention: ILR (n=103) Comparator: Conventional (n=98)	1° endpoint: ■ Time to diagnosis: 43% vs. 6% HR: 6.5; 95% CI: 3.7–11.4; p<0.001.	Limitations:  • Single center, non-blinded trial.  Summary:  • ILR significantly increases diagnostic

	Study type: RCT, 18 mo follow-up of previous study which reported 6 mo follow-up did not demonstrate a reduction in syncopal events or an improvement in QoL with ILR.  Size: n=201 pts	indication for pacing; basic workup including Holter, tilt-table unrevealing.  Exclusion criteria: None stated		2º endpoint:  ■ Time to first recurrence: HR: 1.03: (0.67–1.6), p=0.9. Time to second recurrence longer with ILR, p=0.04.  ■ Improved QoL in ILR group (p=0.03) for general wellbeing.  ■ Overall mortality was 12%, p=NS.	rate and ECG directed treatments in a typical unselected syncopal population.  • Long-term follow-up demonstrated a significant subsequent reduction in syncopal events with improved QoL.
Da Costa A, et al. 2013 23582676 (74)	Aim: To compare ILR and conventional follow-up to estimate prevalence of arrhythmia (pause >5 s, 3 <sup>rd</sup> degree AV block, heart rate <30 bpm for 10 m while awake, >10 beats VT, SVT >165 bpm).  Study type: Multicenter RCT  Size: n=78 pts (11 right BBB, 34 left BBB, 33 bifascicular)	Inclusion criteria: S/P single syncopal episode with BBB (QRS≥120 ms); negative workup (including EPS).  Exclusion criteria: 2nd or 3rd degree AV block; LVEF ≤35%; poor prognosis (<1 y); inability to follow-up; HV interval ≥70 m; inducible VT/SVT; carotid sinus hypersensitivity; subclavian steal; OH.	Intervention: ILR (n=41)  Comparator: Conventional (n=37) (Outpatient visits every 3 mo for 36 mo, diary, 12-lead ECG, 7 d event recorder)	1º endpoint:  • 21/78 developed significant arrhythmia: AV block (14), sick sinus syndrome (4), VT (1), SCD (2).  • Events detectable in 19 pts, with a statistically significant difference found between the ILR and conventional follow-up groups (36.6% vs. 10.8%; p=0.01).  • 18 pts received pacemakers; 1 received ICD.  • No predictors of AV block identified in the ILR group.	Limitations:  Highly-specific subset of pts Small sample size (unavoidable) <3 y of follow-up Not designed to test impact of cost  Summary: ILR superior to conventional follow-up in detecting recurrent syncope in pts with isolated syncope, BBB, and negative EPS. Supports early monitoring after first event.

Sivakumaran, et al. 2003 12867227 (75)	Aim: To compare diagnostic utility of ELR to Holter in determining arrhythmic cause of syncope.  Study type: RCT Size: n=100 pts	Inclusion criteria: SUO: index symptoms of syncope, presyncope, or both, referred for ambulatory ECG monitoring.  Exclusion criteria: None stated	Intervention: Initial 48 H Holter (n=51)  Comparator: Initial 30 d ELR (n=49)	1° endpoint:  •63% ELR vs. 24% Holter had arrhythmia identified or excluded, p<0.0001.  • Arrhythmia identified as cause of syncope in 1 patient with ELR (p=0.3).  • Probability of obtaining symptomrhythm correlation 56% for ELR, 22% for Holter (p<0.00001).	Limitations:  Non-blinding; pre-enrollment evaluation not standardized.  Conclusions: ELRs have a much higher diagnostic yield for pts with syncope or presyncope as compared with Holter monitors. Utility of loop recorders is limited by some pts' inability to operate them correctly.
Rothman SA, et al. 2007 17318994 (76)	Aim: To compare the relative value of a MCOT c/w ELR.  Study type: Multicenter RCT  Size: n=266 pts, 17 centers	Inclusion criteria:  A high clinical suspicion of a malignant arrhythmia; symptoms of syncope, presyncope, or severe palpitations occurring less frequently than once per 24 H; nondiagnostic 24 H Holter or telemetry monitor within 45 d prior to enrollment.  Exclusion criteria: NYHA Class IV HF; MI within prior 3 mo; unstable angina; candidate for or recent valvular cardiac surgery; history of sustained VT/VF; frequent PVCs; documented LVEF ≤35%; pts <18 y of age, condition prohibiting completion of or compliance with protocol.	Intervention: MCOT (n=134)  Comparator: Loop (n=132)	1° endpoint:  • Diagnosis made in 88% of MCOT pts compared with 75% of ELR pts (p=0.008).  • MCOT superior in confirming diagnosis of clinically significant arrhythmias 41% vs. 15%, p<0.001.	■ Neither patient nor investigator blinded (although independent strip review). Patient compliance not 100%.      Conclusions:     ■ In diagnosis of pts with symptoms of a cardiac arrhythmia, MCOT provides a significantly higher yield than standard ELR.     ■ MCOT superior to ELR for detection of clinically significant arrhythmias, with shorter time to diagnosis.

## Data Supplement 12. Nonrandomized Trials, Observational Studies, and/or Registries of Cardiac Monitoring – (Section 3.2.3)

Study Acronym;	Study Type/Design;	Patient Population	Primary Endpoint and Results	Summary/Conclusion

Author; Year Published	Study Size		(P values; OR or RR; & 95% CI)	Comment(s)
Krahn AD, et al. 1995 <u>7671366</u> (77)	Study type: Prospective observational Size: n=16 pts	Inclusion criteria: SUO with resting ECG; ambulatory monitoring; myocardial imaging; and TTT. If noninvasive investigations were negative, EPS performed. ILR implanted with negative EPS.  Exclusion criteria: Pre syncope	1° endpoint: Long-term findings in pts with unexplained syncope and negative laboratory investigations.  Results:  16 pts implanted, and 15 pts (94%) had recurrent syncope 4.4±4.2 mo after implantation.  Syncope was secondary to sinus arrest in 5, AV block in 2, VT in 1, SVT in 1, nonarrhythmic in 6.  Successful therapy in all 15 pts, without recurrence of syncope during 13.0±8.4 mo of follow-up.	Limitations:  Small number of implants, and authors comment on minimal incidence of morbidity and mortality.  Conclusions:  ILR useful for establishing a diagnosis when symptoms are recurrent but too infrequent for conventional monitoring techniques.
Krahn AD, et al. 1999 <u>9918528</u> (78)	Study type: Prospective observational Size: n=85 pts	Inclusion criteria:  2 syncopal episodes within the previous 12 mo or a single episode with a Hx of presyncope as well.  Exclusion criteria: Unlikely to survive 1 y; unable to give informed consent; had a previously implanted programmable medical device; were pregnant; or were women of childbearing potential not on a reliable form of contraception	1º endpoint:  Determine cause of syncope in pts with SUO and recurrent undiagnosed syncope with an ILR  Results:  • During a mean of 10.5±4.0 mo of follow-up, symptoms recurred in 58 pts (68%) 71±79 days (2.3±2.6 mo) after ILR insertion.  • Arrhythmia detected in 42% of pts who recorded a rhythm during recurrent symptoms, with bradycardia present in 18 and tachycardia in 3.  • 5/18 bradycardic pts and 2 additional sinus rhythm pts received a clinical diagnosis of neurally mediated syncope.  • Pts who experienced presyncope much less likely to record an arrhythmia during symptoms compared with recurrence of syncope (24% vs. 70%, p=0.0005).	Limitations:  Select population and a small proportion of pts were unable to activate the device after a spontaneous event.  Conclusions: The strategy of prolonged monitoring is effective and safe in pts with SUO.
Moya A, et al. 2001 <u>11551877</u> (79)	Study type: Prospective observational Size: n=111 pts	Inclusion criteria:  Syncope, absence of significant structural heart disease, and a normal ECG; tilt-testing was negative in 82 (isolated syncope) and positive in 29 (tilt-positive); ≥3 episodes of syncope in the previous 2 ys	ILR in pts with isolated syncope and in pts with tilt-positive syncope to obtain further information on the mechanism of syncope and to evaluate the natural Hx of these pts.  Results: Syncope recurred in 28 (34%) and 10 pts (34%), respectively, and ECG correlation was found in 24 (23%) and 8 (28%) pts, respectively.	Limitations:  • Although documentation of bradyarrhythmia concurrent with a syncopal episode is considered diagnostic, unable to discriminate between an intrinsic cardiogenic abnormality and a neurogenic mechanism.  Conclusions:  • In most pts, the likely cause was neurally

		Evaluaion evitoria:		modiated and the most fragiliant reaches the
		Exclusion criteria: None specified	The most frequent finding, which was recorded in 46%	mediated, and the most frequent mechanism was a bradycardic reflex. In the other cases, a
		None specified	and 62% of pts, respectively, was one or more prolonged	normal sinus rhythm was frequently recorded.
			asystolic pauses, mainly due to sinus arrest.	
Brignole M , et al.	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
2001	Prospective	All pts with any type of BBB with	ILR in pts with BBB and negative EPS to evaluate the	The results of the present study cannot be
11673344 (80)	observational	QRS >100 ms, no documentation of	natural history of these pts and obtain additional information	generalized to all syncope pts with BBB but
(00)	<b>Size</b> : n=52 pts	2 <sup>nd</sup> or 3 <sup>rd</sup> degree AV block, and a negative EPS, and SUO	on the mechanism of syncope.	apply only to the minority of those with a negative conventional workup that includes
	<u>0126</u> . 11–32 pts	Theyative Li 3, and 300	Results:	electrophysiological study.
		Exclusion criteria:	During a follow-up of 3–15 mo, syncope recurred in 22	
		None specified	pts (42%).	Conclusions:
			<ul> <li>The most frequent finding, recorded in 17 pts, was</li> </ul>	<ul> <li>In pts with BBB and negative EPS, most</li> </ul>
			≥prolonged asystolic pause mainly attributable to AV block.	syncopal recurrences have a homogeneous
			• The median duration of the arrhythmic event was 47 s. An additional 3 pts developed nonsyncopal persistent 3 <sup>rd</sup>	mechanism that is characterized by prolonged asystolic pauses, mainly attributable to
			degree AVB, and 2 pts had presyncope attributable to AVB	sudden-onset paroxysmal AV block.
			with asystole.	Saddon onest paroxyoniar / tv sicoli.
			No pts suffered injury attributable to syncopal relapse.	
Garcia-Civera R, et	Study type:	Inclusion criteria: 184 pts with	1° endpoint:	<u>Limitations</u> :
al.	Prospective	SUO.	Diagnostic yield of a protocol in which EPS, TTTs, and ILR	Authors feel no ATP testing was a limitation
2003 12628723	observational	EPS: Any of presence of structural	are selectively used in SUO.	No follow-up of all pts with ILR to confirm
(81)	Size: n=184 pts	heart disease or family Hx of SCD;	Results:	diagnosis
(01)	<u>0126.</u> 11-104 pts	abnormal ECG; significant non-	• 32/72 with inclusion criteria had positive EPS. 80/112 had	Conclusions:
		symptomatic arrhythmia on Holter	positive TTT. 23/40 with negative EPS had positive TTT.	In SUO, selective use of EPS or TTT leads
		monitoring; paroxysmal palpitations	ILR implanted in 15/17 pts with negative EPS who	to positive diagnosis in >70% of cases. ILR
		immediately before or after syncope.	subsequently had negative TTT, with diagnostic activation	can be useful in non-diagnosed cases.
		If these pts (defined as Group A)	in 7. Overall, 143/184 pts with positive diagnosis.	
		had negative EPS, they underwent TTT. 112 pts with initial TTT were		
		defined as Group B.		
i e				
		defined as Group B.		
		Exclusion criteria:		
		Exclusion criteria: None		
Ermis C, et al.	Study type:	Exclusion criteria: None Inclusion criteria:	1° endpoint:	Limitations:
Ermis C, et al. 2003 14516882	Study type: Prospective observational	Exclusion criteria: None	1° endpoint:  Evaluate relative utility of auto-activate ILR based on a arrhythmia grading system in terms of the likelihood that	Limitations:  • Small sample, unclear how generalizable scoring system is.

(82)	Size: n=50 pts	Exclusion criteria: None	they provide a diagnostic basis for syncope.  Results:  Of 529 recordings, auto activation accounted for 86.9% of all the documented arrhythmia episodes (194/223 episodes from 30 pts).  Auto activation provided 90.6% (68 of 75 episodes) of all highly likely diagnoses (i.e., grades 0 and I), and 87.1% of all arrhythmia diagnoses (196 of 225 episodes) (i.e., grades 0 to III).	Conclusions:  ■ Study offers strong support for the value of auto-activation ILR systems, as well as a basis for encouraging further development of arrhythmia scoring.
Boersma L, et al. 2004 14697729 (83)	Study type: Prospective observational Size: n=43 pts	Inclusion criteria: SUO, ≥3 episodes of syncope within 6 mo  Exclusion criteria: None	1° endpoint: Diagnosis of arrhythmia by ILR  Results:  ILR able to record arrhythmic event in 12/43. 10 with bradycardia→, 1 PAF→ medication, and polymorphic VT→ ICD.  3/12 had normal workup. Others had abnormal HUTT, EPS, echo, ECG, Holter.	Limitations:  Not all had full diagnostic workup 18 mo follow-up somewhat limited. Small sample  Conclusions:  ILR is a valuable and effective tool to establish an arrhythmic cause for SUO. The results of HUTT and EPS are neither sufficiently sensitive nor specific enough in this pts group.
Solano A, et al. 2004 <u>15231369</u> (84)	Study type: Observational, prospective, 2-hospital  Size: n=2057, 103 ILR	Inclusion criteria: High-risk syncope: (1) were very frequent, or (2) were recurrent and unpredictable or (3) occurred during the prosecution of a `high risk' activity  Exclusion criteria: Presyncope	1° endpoint: ECG diagnosis made by analysis of the ECG tracing obtained during the first syncopal episode that was correctly recorded by the device.  Results: During a median follow-up of 13 mo, syncope recorded in 52 pts. Pts with SHD more frequently had paroxysmal AV block and tachyarrhythmias and pts without SHD more frequently had sinus bradycardia/sinus arrest or no arrhythmia.	Limitations:  • Limited to high-risk group  Conclusions:  • Mechanism of SUO is different in pts with and without SHD, though diagnostic yield and safety are similar in both groups.
Krahn, et al. 2004 <u>15309004</u> (85)	Study type: Prospective observational Size: n=60 pts	Inclusion criteria: ≥30 y, with LVEF ≥35% and SUO (negative 24 h ambulatory/inpatient monitor, echocardiogram) had ILR	1° endpoint: Prespecified arrhythmias: pause >5 seconds; 3rd degree AVB >10 seconds; Heart rate <30 beats/min for >10 seconds while awake;	Limitations:  ■ Population likely to have recurrence and arrhythmias. Asymptomatic arrhythmias considered diagnostic.

		Exclusion criteria: LVEF <35%; limited survival; neurally mediated syncope	wide complex tachycardia >10 beats; narrow complex tachycardia >180 beats/min for >30 beats.  Results: Recurrent symptoms developed in 30 pts during the 1 y follow-up period (47%), with arrhythmias detected in 14 pts. Pre-specified significant asymptomatic arrhythmias developed in 9 pts with bradycardia in 7 pts who underwent pacemaker implantation. 20 pts had borderline asymptomatic arrhythmias. 5 of these pts went on to have more pronounced diagnostic arrhythmias of same mechanism during further follow-up, including pauses of 6–17 s duration in 3 pts.	Conclusions:  ■ Long-term monitoring of pts with unexplained syncope with automatic arrhythmia detection demonstrated that significant asymptomatic arrhythmias were seen more frequently than anticipated, leading to a change in patient treatment.  ■ Automatic arrhythmia detection provides incremental diagnostic usefulness in long-term monitoring of pts with syncope.
Pierre B, et al. 2008 <u>18325892</u> (86)	Study type: Prospective observational  Size: n=95 pts	Inclusion criteria: SUO: ≥3 episodes of syncope, normal workup including EPS, CSM  Exclusion criteria: LVEF ≤30–35%, candidates for primary ICD	To determine influence of cardiac conduction abnormalities that turn up on resting ECG and the impact of underlying cardiac disease on developments during follow-up.  Results:  During an average follow-up period of 10.2±5.2 mo, 27/43 pts developed a new syncope associated with an arrhythmic event.  Syncope no more frequent in subgroup of pts with cardiac conduction abnormalities on resting ECG, while the frequency of arrhythmic events was similar whether or not the ECG was normal.	Limitations:  ■ Relatively small size with extensive negative workup.  Conclusions:  ■ ILR useful diagnostic tool for recurrent syncope of unknown etiology in pts with or without cardiac conduction abnormalities or cardiac disease.  ■ The absence of arrhythmic events was frequently reported in all patient subgroups. This argues against an empirical pacing strategy in pts with cardiac conduction abnormalities on resting ECG suffering from recurrent syncope, but normal EPS.
Pezawas T, et al. 2008 <u>17947364</u> (87)	Study type: Prospective observational	Inclusion criteria: SUO (ISSUE classification) with ≥2 episodes, then ILR implanted	1º endpoint: Stratify mechanisms and predictors of SUO documented by an ILR in pts with and without SHD.	Limitations:  ■ Not necessarily generalizable—referral center.
	Size: n=70 pts	Exclusion criteria: None	Results:  Syncopal recurrence occurred during 16 mo in 30 pts (91%) with SHD and in 30 pts (81%) without SHD.  45% vs. 51%, respectively, had an ILR documented arrhythmia at time of recurrence which led to specific	Conclusions:  ● Presence of SHD has little predictive value for the occurrence or type of arrhythmia in pts with SUO.

			treatment.  • The remaining 45% with SHD and 30% without SHD had normal sinus rhythm at the time of the recurrence.  • Major depressive disorder predictive for early recurrence during ILR follow-up (p=0.01, HR: 3.35; 95% CI: 1.1–7.1).  • 57% of pts with major depressive disorder had sinus rhythm during recurrence compared with 31% of pts without the disorder (p=0.01).  • Conversely, no patient with major depressive disorder had asystole compared with 33% without (p<0.001).	Pts with major depressive disorder are prone to early recurrence of symptoms and have no evidence of arrhythmia in most cases.
Edvardsson N, et al. 2011 21097478 (88)	Study type: Multicenter prospective observational  Size: n=570 pts	Inclusion criteria: Recurrent SUO or pre-syncope  Exclusion criteria: None specified	1° endpoint:  To collect information on the use of ILR in the patient care pathway and to investigate its effectiveness in diagnosis of SUO in everyday clinical practice.  Results:  Pts evaluated by an average of 3 different specialists for management of their syncope and underwent a median of 13 tests (range 9–20).  The percentages of pts with recurrence of syncope were 19, 26, and 36% after 3, 6, and 12 mo, respectively. Of 218 events within the study, ILR-guided diagnosis was obtained in 170 cases (78%), of which 128 (75%) were cardiac.	Limitations:  12% of implanted pts did not have follow-up visit data.  Pts with pre-syncope only were admitted into the registry, and they have been analyzed and reported together with pts with syncope, since the subgroup was small.  Conclusions: A large number of diagnostic tests were undertaken in pts with unexplained syncope without providing conclusive data. In contrast, the ILR revealed or contributed to establishing the mechanism of syncope in the vast majority of pts. The findings support the recommendation in current guidelines that an ILR should be implanted early rather than late in the evaluation of unexplained syncope.
Linker NJ, et al. 2013 <u>24182906</u> (89)	Study type: Multicenter observational registry (PICTURE)  Size: n=514 pts with ILR (25% implanted during initial work-up, 75%	Inclusion criteria: Recurrent SUO or pre-syncope  Exclusion criteria: No evidence of "unexplained syncope," no follow-up data, ILR implanted for another reason	1º endpoint: First recurrence of syncope leading to a diagnosis or for at least 1 y after implant  Results: Initial (8 tests [IQR 6-14]) vs. Full (14 tests [IQR 10-21]), p<0.0001. Hospitalization and injury before implant less common in	Limitations:  • "Unexplained," "initial workup," or "full evaluation" not defined in protocol  Conclusions:  • Diagnostic yield of ILR high in both protocols.  • High number of testing in both protocols may have been mitigated by earlier ILR.

	after "full evaluation"		pts with "initial work-up": 53 vs. 75%, p<0001, and 23% vs. 39%, p<0.001, as were visits to specialists, p<0.001.  ● Recurrence rate:32 initial vs. 36% full at 12 mo Recurrence with ILR diagnosis: 52 vs. 75% at 12 mo;	
Palmisano P, et al. 2013 23701932 (90)	Study type: observational; 2 center study Size: n=56 pts	Inclusion criteria: History of syncope of suspected arrhythmic nature, negative cardiac and neurological workup, who underwent ILR.  Exclusion criteria: None	cardiac dx: 90 vs. 79%  1º endpoint: Identify predictive factors for pacemaker implantation in pts receiving an ILR  Results: ● Clinically significant bradyarrhythmia was detected in 11 pts (20%), of which 9 cases related to syncopal relapses: predictive factors: >75 y of age (OR: 29.9; p=0.035); a Hx of trauma secondary to syncope (OR: 26.8; p=0.039); and the detection of periods of asymptomatic bradycardia, performed before ILR implantation (OR: 24.7; p=0.045).	Limitations:  Non-blinded, clear selection bias  Conclusions: An advanced age, a history of trauma secondary to syncope, and the detection of periods of asymptomatic bradycardia during conventional ECG monitoring were independent predictive factors for bradyarrhythmias requiring pacemaker implantation in pts receiving an ILR for unexplained syncope.
Gibson TC, et al. 1984 <u>6702676</u> (91)	Study type: Retrospective observational  Size: n=1,512 pts with syncope (of 7,364 total)	Inclusion criteria: Pts underwent 24 H Holter monitoring  Exclusion criteria: None	1º endpoint: Diagnostic yield of Holter for syncope diagnosis  Results:  • 31/1512 (2%) of pts had "arrhythmia-related symptom" that could be diagnostic  • 15 pts had syncope and 7 of the episodes were related to an arrhythmia, usually VT  • Presyncope was reported in 241 pts, with a related arrhythmia in 24	Limitations:  Large sample (registry), many confounders Percentages likely low due to sample  Conclusions:  24 H ambulatory monitoring service rarely results in identifying relevant symptom-related arrhythmias in pts with syncope
Linzer M, et al. 1990 <u>2371954</u> (92)	Study type: Prospective observational Size: n=57 pts	Inclusion criteria: ≥1 episode of SUO  Exclusion criteria: Prior EPS	1º endpoint: Utility of ELR after indeterminate Holter recording  Results: In 14 pts, loop recording definitively determined whether an arrhythmia was cause of symptoms (diagnostic yield 25%; 95% CI: 14–38%). Diagnoses included VT (1 patient), high grade AV block (2 pts), SVT (1 patient), asystole or junctional bradycardia from neurally mediated syncope (3 pts) and normal cardiac rhythms (the remaining 7 pts).	Limitations:  Referral bias, small sample.  Conclusions: Early study of external LR, shows utility in SUO.

Locati ET, et al.	Study type:	Inclusion criteria: Recent (within 1	1° endpoint: To evaluate the role of external 4 wk ECG	The 4 wk external ECG monitoring can be
2016	Prospective	mo) episode of syncope or	monitoring in clinical work-up of unexplained syncope	considered as first-line tool in the diagnostic
<u>26519025</u>	observational,	sustained palpitations (index	and/or sustained palpitations of suspected arrhythmic origin	work-up of syncope and palpitation. Early
(93)	multicenter	event), after being discharged from		recorder use, history of supraventricular
		emergency room or hospitalization	Results: For syncope, the 4 wk diagnostic yield was	arrhythmia, and frequent previous events
	Size: 392 pts; 282 pts	without a conclusive diagnosis, and	24.5%, and predictors of diagnostic events were early start	increased the likelihood of diagnostic events
	(71.9%) enrolled for	a suspected arrhythmic origin	of recording (0–15 vs. >15 days after index event) (OR: 6.2,	during the 4 wk external ECG monitoring.
	palpitations and 110		95% CI: 1.3–29.6, p=0.021) and previous Hx of	<ul> <li>Diary-reported symptoms/events, true</li> </ul>
	(28.1%) for syncope.	Exclusion criteria:	supraventricular arrhythmias (OR 3.6, 95% CI:1.4–9.7,	etiology of event unknown (despite
		None specified	p=0.018).	documented arrhythmia). Authors note the
			<ul> <li>For palpitations, the 4 wk diagnostic yield was 71.6% and</li> </ul>	cumulative diagnostic yield observed may be
			predictors of diagnostic events were Hx of recurrent	an overestimation of the true clinical benefit.
			palpitations (p<0.001) and early start of recording	
			(p=0.001).	

### Data Supplement 13. Nonrandomized Trials, Observational Studies, and/or Registries of In-Hospital Telemetry – (Section 3.2.4)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Benezet-Mazuecos, et al. 2007 <u>17965013</u> (94)	Study type: Prospective cohort study  Size: n=122 pts	Inclusion criteria: Presumptive diagnosis of unexplained, likely cardiogenic, syncope.  Exclusion criteria: Syncope and a documented medical condition actually or potentially responsible for the syncope.	To determine the diagnostic value of cardiac remote telemetry in the setting of unexplained syncope is unknown.  Results:  There were no deaths during the time of monitoring (4.8±2.7 days). Events requiring transfer to the coronary care units occurred in 15 pts (14.7%), principally due to AV block and extreme bradycardia.  Cardiac remote telemetry was diagnostic in 18 pts (17.6%) in whom the arrhythmic event occurred simultaneously with the syncopal episode.  ≥86 y of age (p<0.01) and HF on admission (p<0.04) were the strongest predictors of events.  The best cut-off point as a threshold for	Limitations:  ■ Single center study, and CCU protocols not generalizable.  Conclusions:  ■ Cardiac remote telemetry appears to be a useful tool in the management of pts with unexplained syncope, especially in those older and presenting HF on admission.

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			monitoring time was 72 H (sensitivity 73%,	
5			specificity 86%).	
Lipskis DJ, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
1984	Prospective observational	Pts admitted to telemetry	To determine the benefits of telemetry in terms	Older data, not limited to syncope
<u>6711429</u>			of arrhythmia diagnosis and therapy administered.	
(95)	Size: n=205 pts	Exclusion criteria:		Conclusions:
		None specified	Results:	The diagnostic yield of ECG monitoring in pts
			<ul> <li>14 episodes of significant arrhythmias in 12 pts</li> </ul>	with syncope may be low in the absence of a
			who required specific intervention were detected	high amount of suspicion about an arrhythmic
			over 608 patient-days of monitoring.	cause.
			<ul> <li>Significant arrhythmias occurred only in pts with</li> </ul>	
			known or suspected CAD or in those with	
			previously documented arrhythmias.	
Gibson TC, et al.	Study type: Retrospective	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
1984	observational	Pts underwent 24 H Holter monitoring	Diagnostic yield of Holter for syncope diagnosis	<ul> <li>Large sample (registry), many confounders</li> </ul>
<u>6702676</u>				Percentages likely low due to sample
(91)	Size: n=1,512 pts with	Exclusion criteria:	Results:	
	syncope (of 7,364 total)	None specified	• 31/1512 (2%) of pts had "arrhythmia-related	Conclusions:
			symptom" that could be diagnostic	<ul> <li>24 H ambulatory monitoring service rarely</li> </ul>
			<ul> <li>15 pts had syncope and 7 of the episodes were</li> </ul>	results in identifying relevant symptom-related
			related to an arrhythmia, usually VT	arrhythmias in pts with syncope
			<ul> <li>Presyncope was reported in 241 pts, with a</li> </ul>	
			related arrhythmia in 24 pts	
Linzer M, et al.	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
1990	Prospective observational	≥ 1 episode of SUO	Utility of ELR after indeterminate Holter recording	<ul> <li>Referral bias, small sample.</li> </ul>
<u>2371954</u>				
(92)	Size: n=57 pts	Exclusion criteria:	Results:	Conclusions:
		Prior EPS	<ul> <li>In 14 pts, loop recording definitively determined</li> </ul>	<ul> <li>Early study of ELR, shows utility in SUO.</li> </ul>
			whether an arrhythmia was cause of symptoms	
			(diagnostic yield 25%; 95% CI: 14-38%).	
			<ul> <li>Diagnoses included VT (1 patient), high grade</li> </ul>	
			AVB (2 pts), SVT (1 patient), asystole or	
			junctional bradycardia from neurally mediated	
			syncope (3 pts) and normal cardiac rhythms (the	
			remaining 7 pts).	
Schuchert A, et al.	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
2003	Prospective observational	≥2 SUO within 6 mo, negative TTT,	Assess diagnostic yield of ELR in pts with	<ul> <li>Low sample size, all ELR patient triggered.</li> </ul>

<u>12930497</u>		no SHD, no VVS trigger	negative TTT and recurrent syncope.	
(96)	<b>Size</b> : n=24 pts			Conclusions:
	•	Exclusion criteria:	Results:	<ul> <li>Reasons for ELR were infrequent syncopal</li> </ul>
		None specified	<ul> <li>ELR was not useful for arrhythmia detection in</li> </ul>	events after baseline evaluation, with rare
			pts with syncopal events, no overt heart disease,	events during the limited monitoring period in
			and a negative tilt table test because the cardiac	particular, and premature termination or
			rhythm was stored in only 1 of 8 (13%) pts with	unsuccessful recording in 21% of pts.
			recurrent syncope	

### Data Supplement 14. Nonrandomized Trials, Observational Studies, and/or Registries of Electrophysiology Testing – (Section 3.2.5)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and 95% CI)	Summary/Conclusion Comment(s)
Linzer M, et al. 1997 <u>9214258</u> (97)	Study type: Literature review (population studies, referral studies, or case series)  Size: N/A	Inclusion criteria: Published papers were selected if they addressed diagnostic testing in syncope, near syncope, or dizziness  Exclusion criteria: N/A	1º endpoint: To review the literature on diagnostic testing in syncope that remains unexplained after initial clinical assessment.  Results: After a thorough H&P, and electrocardiography, the cause of syncope remains undiagnosed in 50% of pts. In such pts, information may be derived from the results of carefully selected diagnostic tests, especially 1) EPS in pts with organic heart disease, 2) Holter monitoring or telemetry in pts known to have or suspected of having heart disease, 3) loop monitoring in pts with frequent events and normal hearts, 4) psychiatric evaluation in pts with frequent events and no injury, and 5) TTT in pts who have infrequent events or in whom VVS is suspected. Hospitalization is indicated for high-risk pts, especially those with known heart disease and elderly pts.	Limitations:  Older data, methods unclear.  Conclusions: After a thorough H&P, and ECG, the cause of syncope remains undiagnosed in 50% of pts. Stepwise testing may be helpful in elucidating cause of syncope.
Lacroix D, et al. 1991 1950999 (98)	Study type: Prospective cohort  Size: n=100 pts	Inclusion criteria: Pts with syncope of unclear etiology who underwent EPS.  Exclusion criteria: Documented arrhythmia at	1° endpoint: To compare the results of 24 H monitoring and EPS in the evaluation of pts with recurrent syncope, and additionally to analyze the usefulness of the signal-averaged ECG and of body surface potential mapping in predicting the inducibility of VT.  Results:	Limitations:  Neurologic and TTT not performed.  Conclusions: EPS had a higher diagnostic yield than Holter monitoring regardless of cardiac pathology. ECG signal-averaging was useful in predicting VT only in pts

		presentation and those with Wolff-Parkinson-White syndrome	<ul> <li>CAD was found in 46 pts and other heart disease was found in 19. EPS was diagnostic in 44 pts, while Holter monitoring suggested a diagnosis in only 21 pts.</li> <li>Abnormal body surface potential mapping was frequently seen (56%), especially in CAD (70%), or with inducible VT (87%).</li> <li>Late potentials were recorded in 13 pts with CAD; 5 had inducible VT. In 7 other pts with VT, they were either absent or BBB was found.</li> <li>Thirteen deaths occurred, and EPS guided therapy resulted in a low rate of total cardiac death.</li> </ul>	with CAD without BBB. Body surface potential mapping was abnormal in most pts with cardiac disease, but poorly predicted VT.
Click RL, et al. 1987 <u>3825942</u> (99)	Study type: Prospective cohort  Size: n=112 pts	Inclusion criteria: Syncope/near syncope, symptomatic pts with BBB undergoing EPS  Exclusion criteria: CV collapse, or requiring resuscitation	1º endpoint: To determine the role of invasive EP testing in pts with symptomatic BBB.  Results: Cumulative 4 y survival rate and recurrent syncope, respectively:  83% in 16 pts with no therapy (normal study results); 19%  84% in 34 pts with permanent pacing alone; 6%  63% in 39 pts with antiarrhythmic therapy alone; 33%  84% in 21 pts with both antiarrhythmic therapy; 19%	Limitations:  Older data, limited and specific population  Conclusions: In symptomatic pts with BBB and normal EP test results, prognosis is good without treatment. In pts undergoing permanent pacing based on EP testing, survival is good and rate of symptom recurrence is low. EP testing identifies pts with inducible VT for whom antiarrhythmic therapy is indicated but who nevertheless have a poor prognosis.
Reiffel JA, et al. 1985 <u>4072872</u> (100)	Study type: Prospective cohort  Size: n=59 pts	Inclusion criteria: 24 H ambulatory ECG monitoring and then EP testing for unexplained syncope.  Exclusion criteria: None specified	To assess whether findings on ambulatory monitoring not obtained during syncope can be used to indicate the results which are found on EP testing in pts with recurrent syncope.  Results:  Although 29 pts had abnormalities on EP testing, 13 of which were severe, in only 6 were the findings suggested by the abnormalities recorded during ambulatory monitoring.  21 pts had concordance between EP testing and ambulatory monitoring results, but in 15 of the 21 results of both tests were normal.	Limitations:  Not a prospective comparison of ambulatory ECG monitoring and EP testing in all pts with syncope, since pts whose workup stopped after ambulatory ECG monitoring were not enrolled in the study. It is, however, a study of EP results as compared to ambulatory ECG monitoring in pts who do undergo EP testing following non diagnostic ambulatory ECG monitoring -a population frequently encountered in clinical EP laboratories. Thus it biases the results toward the detection of abnormalities by EP tests  Conclusions: Severe abnormalities were more frequently detected in our patient population by EP testing than by ambulatory monitoring, especially if pts had organic heart disease.

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Gulamhusein S, et	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
al.	Prospective	Unexplained syncope/near	To assess the value of clinical EPS using intracardiac recording	<ul> <li>Empirical permanent pacing in pts with symptoms</li> </ul>
1982	cohort	syncope who underwent	and PES in 34 pts who had unexplained syncope and/or	appeared to be beneficial, but this result is difficult to
<u>7137203</u>		PES	presyncope.	evaluate because of the high incidence of spontaneous
(101)	<b>Size:</b> n=34 pts			remission in this group.
		Exclusion criteria:	Results:	
		None specified.	EPS diagnostic in 4 pts (11.8 percent) and led to appropriate	Conclusions:
			therapy that totally relieved symptoms.	The diagnostic yield of EP testing is low in a patient
			• Results were abnormal but not diagnostic in 2 pts (5.8%) and	population that has no ECG abnormality or clinical
			normal in the remaining 28 pts (82.4%).	evidence of cardiac disease.
			Over mean follow up of 15 mo, 16 pts (47%) had no further	
			episodes in the absence of any intervention. In 4 pts (11.8%), a	
			definitive diagnosis was made. In 7 pts, permanent pacing was	
			instituted empirically with relief of syncope.	
Sagrista-Sauleda J,	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
et al.	Retrospective	Syncope of unknown	To assess the diagnostic yield of the head-up tilt test (n=600) and	Retrospective study in very specific population of pts
2001	cohort	etiology who underwent TTT,	electrophysiology (n=247/600) in pts with syncope of unknown	undergoing TTT.
11350095		after H&P, ECG, CSM,	origin established according to simple clinical criteria.	
(102)	Size: n=600 pts	Holter monitoring,	9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	Conclusions:
,	'	echocardiogram (in selected	Results:	The diagnostic yield of the TTT and
		pts), exercise stress testing	Positive responses to the tilt test were more common in pts who	electrophysiology differs in groups of pts with syncope
		(in selected pts),	had suffered their first syncope at an age ≤ 65 y (group I) than in	of unknown origin, established according to simple
		neurological evaluation. EPS	older pts (group II) (47% vs. 33%, p<0.05; OR: 1.8; 95% CI: 1.2–	clinical criteria. These findings have a bearing on
		was performed if clinically	2.78), and in pts with a normal ECG and without organic heart	selecting the most appropriate test in a particular
		indicated, mostly in pts with	disease than in the other subgroups of pts (47% vs. 37%,	patient.
		organic heart disease, an	p<0.008, OR: 1.6).	Factor
		intraventricular conduction	The lowest rate of positive response was observed in older pts	
		defect or a suspicion of	with an abnormal ECG and organic heart disease.	
		arrhythmia-related syncope.	Electrophysiology disclosed abnormal findings in group II more	
			often than in group I (23% vs 7%, p<0.001, OR 3.7, 95% CI: 1.7–	
		Exclusion criteria:	9.2).	
		None specified.	The diagnostic yield from electrophysiology was higher in pts	
			with an abnormal ECG than in those with a normal ECG (22% vs.	
			3.7%, p<0.0005, OR: 7.1), and it was especially low in pts with a	
			normal ECG and without organic heart disease (2.6%).	
Gatzoulis KA, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
19419396	Prospective	Syncope of unknown	To assess the utility of noninvasive electrocardiographic	Specific population, unclear generalizability.
(103)	cohort	etiology who had an ECG,	evaluation (12-lead ECG and 24 H ambulatory	apatima papaiation, antition gariational
()	33.1011		Total action (12 load 200 and 27 Hambulator)	

	<u>Size:</u> n=421 pts	an EPS, and 24 H ambulatory monitoring  Exclusion criteria: None specified	electrocardiographic recordings) to predict electrophysiology study results in pts with undiagnosed syncope.  Results:  Pts were divided into 4 groups: group 1, abnormal ECG and ambulatory monitor; group 2, abnormal ECG only; group 3, abnormal ambulatory monitor; and group 4, normal ECG and ambulatory monitor. The likelihood of finding at least one abnormality during EP testing among the 4 groups was highest in group 1 (82.2%) and lower in groups 2 and 3 (68.1% and 33.7%, respectively). In group 4, any EPS abnormality was low (9.1%). ORs were 35.9 (p<0.001), 17.8 (p<0.001), and 3.5 (p=0.064) for abnormal findings on EPS, respectively (first 3 groups vs. the 4th one).	Conclusions:  • Abnormal ECG findings on noninvasive testing are well correlated with potential brady- or/and tachyarrhythmic causes of syncope, in EPS of pts with undiagnosed syncope.
Hess DS , et al. 1982 <u>7148707</u> (104)	Study type: Prospective observational Size: n=32 pts	Inclusion criteria: Syncope of unclear etiology  Exclusion criteria: None specified	1º endpoint: Detection of brady and tachyarrhythmias in EPS to elucidate cause of SUO.  Results: ■ 18/32 pts had definitive EPS diagnosis; 11 pts with inducible VT 5 pts with SND; 1 patient with infra-His AVB; 1 patient with quinidine-related VT	Limitations:  ■ Small study and most pts had organic heart disease (more inducible VT).  ■ Older data, medical therapy changed now.  Conclusions:  ■ The study shows some value in EPS in elucidating cause of syncope, in selected population with SUO.
Gulamshusein S, et al. 1982 7137203 (101)	Study type: Prospective observational Size: n=34 pts	Inclusion criteria: SUO; all undergoing EPS; ≥1 syncopal or ≥2 presyncopal episodes; no cause of syncope on exam; normal ECG and 48 H Holter, normal neurologic testing (including EEG and CT-head); normal echo and CXR  Exclusion criteria: None specified	1° endpoint: Assess diagnostic yield of EPS in SUO.  Results: • EPS diagnostic in 4 pts and led to therapy.  • During mean 15 mo f/u, 16 pts had no further episodes in absence of any intervention	Limitations:  ■ EPS less diagnostic than predicted: some pts required pacing despite normal or nondiagnostic EPS.  Conclusions:  ■ Diagnostic yield of EPS testing is low in a patient population that has no ECG abnormality or clinical evidence of cardiac disease.
Akhtar M , et al.	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations:</u>

1983	Prospective	SUO (≥ 2 episodes in	To assess results of EPS with PES in pts with recurrent syncope	All pts received EPS, high risk group.
<u>6189057</u>	observational	preceding y); negative		
(105)		evaluation	Results:	Conclusions:
	Size: n=30 pts		Sustained or nonsustained VT and/or VF induced in 11/30; SND	EPS with PES can uncover type of arrhythmic
		Exclusion criteria: None specified	in 4/30; Intra-His AVB in remaining 1/30.	disturbance in a significant number of cases.
			• 14/16 remained free of symptoms following therapy based on	
			results of EPS during a mean 16 mo f/u.	
			<ul> <li>In 2/16 syncope recurred (one arrhythmic and one non-</li> </ul>	
			arrhythmic) despite pacemaker therapy for SND detected during EPS.	
			<ul> <li>In remaining 14/30 pts, EPS and PES did not induce arrhythmia</li> </ul>	
			which could account for patient symptomatology. However, 11/14	
			pts experienced a recurrence of symptoms within a 6–25 mo	
			period (mean 16.2±6.8).	
			Of 15/16 pts with inducible arrhythmias considered clinically in its factor to be determined to a discount for the second factor of the second factor o	
			significant had structural heart disease.  • 3/14 pts without clinically significant arrhythmias had structural	
			heart disease.	
Morady F, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
1984	Prospective	SUO undergoing EPS	Diagnostic yield of EPS with PES in pts with SUO	In treated pts who did not have recurrence of
6475778	observational		Stagnostic yield of Er o marr 20 in plo mar occ	syncope, it is presumed that syncope did not recur
(106)		Exclusion criteria:	Results:	because the cause of syncope was correctly identified
	Size: n=32 pts	2 <sup>nd</sup> or 3 <sup>rd</sup> degree AV block;	<ul> <li>HV interval ≥70 ms or greater in 12 pts</li> </ul>	and effectively treated.
		symptomatic SVT; VT;	<ul> <li>Pathologic infranodal AVB during atrial pacing occurred in 2</li> </ul>	In some pts, the decision to implant PPM was due to
		evidence of SND; carotid	pts→PPM	patient preference, not EPS testing.
		sinus hypersensitivity; or a	Monomorphic VT induced in 9 pts and polymorphic VT in	O I I
		history consistent with	5→AAD	Conclusions:
		classic vasovagal or vasodepressor syncope	Actuarial incidence of sudden death was 10% at 45 mo of follow up.	<ul> <li>Approximately 50% of pts with BBB and unexplained syncope who undergo EPS are found to have a</li> </ul>
		vasouchicssol syllope	follow-up  ■ Only 2 pts had recurrent syncope; both had normal EPS	clinically significant abnormality.
			Only 2 pto had recuirent syncope, bott had nothfal EFS	Long-term management guided by the results of
				ESP generally is successful in preventing recurrent
				syncope.
Doherty JU, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
4005	Decompositive	SUO undergoing EPS	EPS findings of pts with SUO	EPS negative group differed in the frequency of
1985	Prospective	300 undergoing LF3	EF3 illidings of pts with 300	, , ,
3976512 (107)	observational	Exclusion criteria:	Results:	heart disease.

	<u>Size</u> : n=119 pts	Known cause of syncope	<ul> <li>Presence of structural heart disease (p=0.0033) and previous MI (p=0.05) were the only clinical or ECG predictors of a positive EPS.</li> <li>Therapy guided by EPS and pts followed for 27±20 mo. In pts with negative EPS results, 76%±11% symptom free at follow-up, compared to 68%±10% in positive EPS group.</li> <li>No clinical variables helped to predict remission in absence of therapy.</li> <li>One patient in negative EPS response group and 2 pts in EPS positive group died suddenly.</li> <li>Total CV mortality 13% in positive EPS response group, and 4% in negative EPS response group.</li> </ul>	Conclusions:  ■ EPS can identify a subgroup of pts at low risk of recurrence and sudden death in the absence of therapy.
Olshansky B, et al. 1985 <u>3968306</u> (108)	Study type: Prospective observational  Size: n=105 pts	Inclusion criteria: SUO undergoing EPS  Exclusion criteria: None specified	1º endpoint: To determine the significance of inducible tachycardia in SUO  Results:  ● 65% did not have inducible tachycardia. 12/60 pts followed had recurrent syncope.  ● VT or SVT inducible in 35%, and inducible tachycardia common in pts both with and without heart disease.  ● 7/13 pts receiving ineffective therapy had recurrence of syncope or cardiac arrest (p<0.05).  ● On resumption of effective therapy, no syncope recurred for 15.6 mo (p<0.025).	Limitations:  ■ Small number lost to follow up  ■ Does not factor in that some pts have remission spontaneously  ■ Nontrivial number of pts receiving ineffective therapy had high percentage of recurrence.  Conclusions:  ■ Inducible tachycardias found in approximately 30% of pts with SUO, common in pts both with and without heart disease.  ■ Adherence to AAD therapy guided by EPS may prevent recurrence.
Teichman SL, et al. 1985 4025122 (109)  Krol RB, et al. 1987	Study type: Prospective observational Size: n=150 pts  Study type:	Inclusion criteria: SUO undergoing EPS  Exclusion criteria: None specified  Inclusion criteria:	1º endpoint: Diagnostic yield and therapeutic efficacy of EPS in pts with SUO  Results:  162 abnormal EPS findings that could explain SUO in 112 pts His-Purkinje disease in 49 pts (30%), inducible ventricular arrhythmias in 36 (22%), AVB in 20 (12%), SND in 19 (12%), inducible supraventricular arrhythmias in 18 (11%), carotid sinus hypersensitivity in 15 (9%), and hypervagotonia in 5 (3%). Follow up data in 137 pts (91%) (mean 31 mo) showed recurrences in 16/34 pts (47%) without and 15/103 pts (15%) with EP findings despite therapy directed by EPS (p<0.0005).	Limitations:  Observational data, limited sample, no control.  Conclusions:  This study and a review of the literature indicate that EPS useful in elucidating causes of SUO and directing therapy  A significant number of pts benefit from EPS, even when only clearly abnormal findings are considered diagnostic, when only a single syncopal event has occurred, or whether or not organic heart disease or an abnormal ECG is present.  Limitations:

<u>3598006</u> (110)	Prospective observational  Size: n=104 pts	Exclusion criteria: Sustained VT; high grade AV block; CSH; vasovagal/vasodepressor syncope; QT prolongation; AS; HCM; symptomatic postural hypotension; brady/tachyarrhythmia known to cause syncope	To evaluate whether clinical variables enable stratification of pts with SUO into low and high probability of having abnormal EPS (SNRT ≥3 seconds; HV interval ≥100 ms; infranodal block during atrial pacing; monomorphic VT; and SVT associated with hypotension)  **Results:  • 31 pts had positive EPS, inducible VT most common finding (71% of positive studies).  • LVEF ≤40% most powerful predictor of a positive EPS (p<0.00001), followed by the presence of BBB (p<0.00003), CAD (p<0.0003), remote MI (p<0.00006), use of type 1 AAD (p<0.00003), injury related to LOC (p<0.01) and male sex (p<0.01).  • A negative EPS associated with LVEF >40% (p<0.00001), absence of structural heart disease (p<0.00001), normal ECG (p<0.0001) and normal ambulatory ECG monitoring (p<0.0001).  • Probability of a negative study increased as number and duration of syncopal episodes increased.	No episodes of syncope with ECG recorded.      Conclusions:     On the basis of clinical variables, majority of pts with SUO can be predicted to have normal or abnormal EPS. This may lead to cost-effective use of EPS.
Fujimura O, et al. 1989 <u>2594030</u> (111)	Study type: Prospective observational Size: n=21 pts	Inclusion criteria: ECG evidence of intermittent AV block (n=13) or sinus pauses (n=8) causing syncope, but whose cardiac rhythm had reverted to normal by the time of referral  Exclusion criteria: None specified	1º endpoint:     Sensitivity of EPS in detection of transient bradycardia in pts in normal sinus rhythm referred for pacemaker implantation after ECG documentation of transient bradycardia resulting in syncope.      Results:     3/8 with documented sinus pauses had abnormal EPS including a prolonged SNRT in 1 and carotid-sinus hypersensitivity in 2 pts.     3/8 pts had abnormalities unrelated to syncope.     2/13 with documented AVB had abnormalities suggesting correct diagnosis.	Limitations:  Small study limited to pts with transient ECG findings.  Conclusions: Negative EPS in a patient with a normal cardiac rhythm who has experienced syncope does not exclude a transient bradyarrhythmia as a cause of the syncope.
Moazez F, et al. 1991 <u>1985382</u> (26)	Study type: Prospective observational Size: n=91 pts	Inclusion criteria: SUO undergoing EPS  Exclusion criteria: BBB, unknown data on LVEF or SAECG	1° endpoint: To examine usefulness of clinical and noninvasive variables to predict EPS, and to compare EPS results and therapy with syncope recurrence  Results:  • Multivariate analysis identified +SAECG, LVEF, and history of sustained monomorphic VT as risk factors for induction of	Limitations:  ■ BBB pts excluded.  ■ No TTT or isoproterenol infusion performed.  Conclusions:  ■ Pts who have inducible sustained monomorphic VT at EPS can be identified using certain clinical and noninvasive variables.

			sustained monomorphic VT at EPS.	When these pts undergo EP-guided therapy, their
			●17 pts had recurrence of syncope over 19.0±8.3 mo of follow-	rate of recurrence of syncope similar to pts who had no
			up.	arrhythmia induced at EPS.
			<ul> <li>Recurrence rates among empiric, EP-guided (sustained</li> </ul>	<ul> <li>Empiric therapy does not offer any benefit over no</li> </ul>
			monomorphic VT), and no therapy groups were similar.	therapy in reducing the rate of recurrent of scope.
Sra JS, et al.	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations:</u>
1991	Retrospective	SUO undergoing EPS, and	To determine the clinical characteristics of subgroups of pts with	Retrospective evaluation
2029096	observational	HUTT if negative.	SUO having EPS and HUTT and to assess efficacy of various	
(112)		-	therapies.	Conclusions:
, ,	Size: n=86 pts	Exclusion criteria:		<ul> <li>The combination of EPS and HUTT can identify the</li> </ul>
		None specified	Results:	underlying cause of syncope in as many as 74% of pts
		·	• 34% had abnormal EPS, with sustained monomorphic VT	presenting with SUO.
			induced in 72%, with 76% of these pts with structural heart	
			disease.	
			• 40% had syncope provoked by HUTT, with 6% of these pts with	
			structural heart disease.	
			<ul> <li>The cause of syncope remained unexplained in 26%, with 30%</li> </ul>	
			of these pts with structural heart disease.	
			<ul> <li>During a median follow-up period of 18.5 mo, syncope recurred</li> </ul>	
			in 9 (10%) pts.	
Muller T, et al. 1991	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
<u>2044546</u>	Prospective	SUO	EPS findings of pts with SUO.	<ul> <li>Small sample, moderate follow-up</li> </ul>
(113)	observational			
		Exclusion criteria:	Results:	Conclusions:
	Size: n=134 pts	None specified	<ul> <li>Conduction abnormalities and tachyarrhythmia could account</li> </ul>	<ul> <li>19% of pts will have a recurrent event.</li> </ul>
			for syncope in 40 pts (30%).	Female gender may be an independent predictor of
			<ul> <li>37/40 received pacing or antiarrhythmic therapy c/w 23/94 who</li> </ul>	favorable outcome.
			had a negative study and received empiric therapy (p<0.0001).	
			<ul> <li>During a mean follow-up of 22±17 mo, 22 pts had recurrent</li> </ul>	
			syncope and 4 died suddenly	
			<ul> <li>Men had a higher incidence of recurrent syncope than women</li> </ul>	
			(26% vs. 6%, P<0.005).	
Denniss AR , et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
1992	Prospective	SUO undergoing EPS	Compare incidence of EPS abnormalities in pts with and without	Failure to demonstrate mortality reduction may be
<u>1572741</u>	observational		heart disease, and the effect of treatment of these abnormalities	due to high-risk group, refractory to treatment.
(114)			on recurrence of syncope.	
	<u>Size</u> : n=111 pts	Exclusion criteria:		Conclusions:
		None specified	Results:	Syncope pts with heart disease more likely to have a

			<ul> <li>Abnormalities detected in 31/73 with heart disease but in only 6/38 with no heart disease (p&lt;0.01).</li> <li>During follow-up, syncope recurred in 2/37 treated because of abnormal findings, compared with a recurrence rate of 18/74 in untreated group (p&lt;0.05).</li> <li>Probability of remaining free from syncope at 2 y was 0.94 in the treated group and 0.72 in the untreated group (p&lt;0.05).</li> <li>Mortality during follow-up was only in heart disease group with 5/30 treated dying compared with 3/43 untreated pts (p=NS).</li> </ul>	diagnostically useful study than pts with normal hearts.  • Treatment directed at correction of abnormalities detected at EPS reduced recurrence of syncope but did not significantly affect mortality.
Link MS, et al. 1999 10235091 (115)	Study type: Retrospective observational	Inclusion criteria: Syncope or presyncope and CAD, with unclear etiology	1° endpoint: Long-term outcome of pts with CAD and non-diagnostic work-up, including EPS	Limitations:  • Retrospective, HV ≥90 ms excluded.  Conclusions:
(113)	<u>Size</u> : n=68 pts	Exclusion criteria: Sudden cardiac death; spontaneous sustained VT;	Results:  • At a mean follow-up of 30±18 mo, 17 pts had recurrence.  • All 4 arrhythmias occurred in pts with LVEF ≤25%.	Conclusions:  ■ In pts with CAD and syncope, noninducibility at EPS predicts a lower risk of SCD and VT/VF.
		noninvasive testing explained syncope	<ul> <li>Predictors of all-cause mortality: age (p=0.05) and reduced LVEF (p=0.02).</li> <li>Predictors of ventricular arrhythmias: BBB (p=0.07), longer runs of NSVT (p=0.08), lower LVEF (22.5±3% vs. 43±16%), p=0.09).</li> </ul>	<ul> <li>In pts with a reduced LVEF, the risk remains up to 10%/y; these pts may warrant treatment with ICDs.</li> </ul>
Knight BP, et al. 1999 10362200 (116)	Study type: Prospective observational	Inclusion criteria: "Syncope Group": NICM, SUO, and negative EPS who underwent ICD (n=14);	1° endpoint: Determine outcome of pts with NICM, negative EPS, and SUO treated with ICD	Limitations:  ■ Small size, unclear "appropriate" shocks in devices without stored EGM.
(110)	Size: n=33 pts	"Arrest Group": NICM with cardiac arrest and ICD (n=33)	Results:  ■ 50% in Syncope Group vs. 42% in Arrest Group received appropriate shocks (p=0.1).	Conclusions:  ■ The high incidence of appropriate ICD shocks and the association of recurrent syncope with ventricular arrhythmias support treatment of pts with nonischemic
		Exclusion criteria: None specified.	<ul> <li>Mean duration from device implant to first appropriate shock in Syncope Group 32±7 mo (95% CI: 18–45) compared to 72±12 mo (95% CI: 48–96, p=0.1).</li> </ul>	cardiomyopathy, SUO and a negative EPS with an ICD.
Sagristà-Sauleda J, et al. 2001 11350095	Study type: Observational cohort	Inclusion criteria: Group I: first syncope at age ≤65 y (n=464 pts) Group II: first syncope at	1° endpoint: To assess diagnostic yield of TTT and EPS in different groups of pts with SUO established according to simple clinical criteria.	Limitations:  Retrospective, and only TTT pts studied. EPS done at physician discretion.
(102)	<u>Size</u> : n=600 pts	age >65 y (n=136 pts) 4 subgroups in both:	Results:  Positive TTT-more common in (group I) than group II (47% vs.	Conclusions:  • The rate of positive responses to the head-up tilt test

		A: pts who no organic heart disease and a normal ECG (n=359 pts) B: pts with no organic heart disease (n=122 pts) and an abnormal ECG; C: pts with organic heart disease and a normal ECG (n=44 pts) D: pts with organic heart disease and an abnormal ECG (n=75 pts)  Exclusion criteria: None specified	33%, p<0.05; OR: 1.8, 95% CI: 1.2–2.78), and subgroup A (49% vs. 37%, p<0.008, OR:1.6).  • EPS disclosed abnormal findings in group II more than in group I (23% vs. 7%; p<0.001, OR: 3.7; CI: 1.7–9.2).  • Diagnostic yield from EPS was higher in pts with an abnormal ECG (subgroups B and D) than in those with a normal ECG (22% vs. 3.7%,p<0.0005, OR: 7.1), and it was low in pts with a normal ECG and without organic heart disease (2.6%).	was higher in younger pts and in pts with a normal ECG and without organic heart disease (49%), while older pts with an abnormal ECG and with organic heart disease had the lowest rate of positive responses (18%).  • The diagnostic yield of EPS was higher in older pts, in pts with organic heart disease and with an abnormal ECG (26%); it was lowest in pts without organic heart disease and with a normal ECG (2.6%).
Mittal S, et al. 2001 11499726 (117)	Study type: Prospective observational Size: n=118 pts	Inclusion criteria: CAD and unexplained syncope who underwent EPS  Exclusion criteria: Pts with a documented sustained ventricular arrhythmia or those resuscitated from sudden cardiac death.	1º endpoint: To determine the incidence and prognostic significance of inducible VF in pts with CAD and unexplained syncope.  Results: Sustained monomorphic VT was inducible in 53 (45%) pts; in 20 (17%) pts, VF was the only inducible arrhythmia; and no sustained ventricular arrhythmia was inducible in the remaining 45 (38%) pts. There were 16 deaths among during a follow-up period of 25.3±19.6 mo. The overall one and 2 y survival in these pts was 89% and 81%, respectively. No significant difference in survival was observed between pts with and without inducible VF.	Limitations:  All pts had CAD (limited generalizability)  VF rarely induced with 2 extrastimuli  Small sample size  Conclusions:  Induction of VF in pts with CAD and unexplained syncope may be of limited prognostic significance. VF was the only inducible ventricular arrhythmia at EP testing (using up to triple ventricular extrastimuli) in 17% of these pts. ICD implantation in pts with syncope of undetermined origin in whom only sustained VF is induced during EP testing, especially with triple ventricular extrastimuli, may merit reconsideration.

## Data Supplement 15. Nonrandomized Trials, Observational Studies, and/or Registries of Tilt Table Testing – (Section 3.2.6.)

Study Acronym;	Study Type/Design;	Patient Population	Primary Endpoint and Results	Summary/Conclusion
Author;	Study Size		(include P value; OR or RR;	Comment(s)
Year Published	·		and 95% CI)	. ,

Kanny DA at al	Chudu human	Inclusion evitorio:	40 and relati	Limitational
Kenny RA, et al. 1986	Study type:	Inclusion criteria:	1º endpoint:	Limitations:
	Case-control study	Syncope of unclear	To investigate the utility of syncope that remained unexplained despite	Small sample, all with EPS
<u>2872472</u>	C: OF	etiology	full clinical and electrophysiological assessment.	O-malusiana.
(118)	Size: n=25 pts	F .1		Conclusions:
	(15 test, 10 control)	Exclusion criteria:	Results:	Reproduction of symptoms during tilt allows
		None specified	• In 10 pts and one control VVS developed after 29±19 min (p<0.001).	Identification of the contribution to syncope
			In symptomatic pts SBP fell from $150 \pm 32$ to $56 \pm 9$ mm Hg (p<0.001)	made by changes in heart rate and BP and
			and heart rate from $62 \pm 9$ to $38 \pm 12$ bpm (p<0.01). In each case	therefore permits the selection of pts in whom
			symptoms during the test reproduced those previously experienced. No	cardiac pacing may be beneficial.
			clinical findings predicted development of syncope during tilt. Baseline	
			SBP and heart rate did not differ significantly between pts and controls.	
			Pacemakers were implanted in 7 pts who have remained symptom free	
			since implant (follow-up 10 ± 3 mo).	
Fitzpatrick A, et	Study type:	Inclusion criteria:	1º endpoint:	Limitations:
al.	Retrospective cohort	Recurrent syncope	To utilize TTT to discover the incidence of malignant VVS in pts with	All pts underwent EPS, with a large
1991	<b>6</b> : 000 t	, .	recurrent syncope.	percentage (70%) of abnormal findings (limited
<u>2040321</u>	Size: n=322 pts	Exclusion criteria:		generalizability).
(119)		None	Results:	
			Prolonged 60 degrees head-up tilt was performed in 71/93 pts with	Conclusions:
			unexplained syncope, and reproduced VVS and presenting symptoms in	TTT is a valuable provocative tool for VVS and
			53 (75%), or 16% of the whole population reported.	may reduce the number of syncopal pts that
			Positive tilts were significantly less common in a group of 27 pts of	remain undiagnosed, although these early
			similar age without a Hx of syncope (7%), and a random sample of 37	observations do not allow an exact appraisal of
			pts with AVB (n=16), sick sinus syndrome (n=18) and inducible	the sensitivity and specificity of the TTT.
D D I . I	0( 1 (	Landard State Dis	tachyarrhythmia (n=3), (19%, 11% and 0% respectively, p<0.01).	12.26.0
Passman R, et al.	Study type:	Inclusion criteria: Pts	1º endpoint:	Limitations:
2003	Retrospective cohort	with syncope	To assess the prevalence and type of apparent neurologic events	The retrospective nature of this study may
<u>12963568</u>	<b>C:</b>	Fredrick seitenier	associated with tilt table testing.	have resulted in inadequate documentation of all
(120)	<u>Size</u> : n=694 pts	Exclusion criteria:	Decelle	potential seizure-like or atypical neurologic
		None	Results:	events at the time of TTT
			• 222/694 with positive TTT. 18 pts (8%) had neurologic events during	Canalysians
			TTT. 11 pts (5%) had apparent tonic-clonic seizure-like activity and 7 pts	Conclusions:  Neurologic events are common during
			(3%) had non-tonic-clonic neurologic events.	episodes of neurocardiogenic syncope, and this
			The pts with tonic-clonic seizure-like activity had a significantly lower      The pts with tonic-clonic seizure-like activity had a significantly lower      The pts with tonic-clonic seizure-like activity had a significantly lower	diagnosis should be considered in the evaluation
			SBP reading at the termination of tilt table testing than all other pts whose TTT results were positive (p=0.04).	of unexplained seizure-like activity.
				or unexplained seizure-like activity.
			• The heart rate at the time of test termination was significantly lower in	
			the pts with tonic-clonic seizure-like activity and non-tonic-clonic	

Grubb BP, et al. 1991 <u>1952474</u> (121)	Study type: Prospective cohort Size: n=15 pts	Inclusion criteria: Recurrent unexplained seizure-like episodes, unresponsive to antiseizure medication.  Exclusion criteria: None	neurologic events (p<0.01) than in those with positive test results and no provoked neurologic events, and asystole was provoked more frequently in these 2 patient populations (p=0.03).  1º endpoint: To evaluate the usefulness of head-upright TTT in the differential diagnosis of convulsive syncope from epileptic seizures in pts with recurrent idiopathic seizure-like episodes.  Results: Syncope associated with tonic-clonic seizure-like activity occurred in 6/15 (40%) during the baseline tilt and in 4/15 during isoproterenol infusion (total positive tests, 67%). The EEG showed diffuse brain wave slowing (not typical of epileptic seizures) in 5/5 pts during the convulsive episode. All pts who had positive test results eventually become tilt table negative after therapy, and over a mean follow-up period of 21 ± 2 mo, no further seizure-like episodes have occurred.	Limitations:  • Small sample, single center study  Conclusions:  • Upright TTT combined with isoproterenol infusion may be useful to distinguish convulsive syncope from epileptic seizures
Song PS, et al. 2010 20046517 (122)	Study type: Retrospective cohort  Size: n=226 pts	Inclusion criteria: Syncope during HUTT without any other cause of syncope  Exclusion criteria: None	1º endpoint: To assess the incidence and characteristics of seizure-like activities during HUTT-induced syncope in pts with neurally mediated reflex syncope.  Results: 13/226 pts showed seizure-like activities, with 5/226 having multifocal myoclonic jerky movements, 5/226 (2.21%) having focal seizure-like activity involving one extremity, and 3/226 having upward deviation of eye ball. Comparison of pts with and without seizure-like activities revealed no significant differences in terms of clinical variables and hemodynamic parameters during HUTT.	Limitations:  ■ Retrospective in design. Of 1,383 pts with positive HUTT, 1,157 pts were excluded from the study because they did not lose consciousness during HUTT.  Conclusions:  ■ Seizure-like activities occurred occasionally during HUTT-induced syncope in pts with neurally mediated reflex syncope. The seizure-like activities during HUTT might not be related to the severity of the syncopal episodes or hemodynamic changes during HUTT.
Zaidi A, et al. 2000 <u>10898432</u> (123)	Study type: Prospective cohort Size: n=74 pts	Inclusion criteria: Diagnosis of epilepsy, with continued attacks despite adequate anticonvulsant drug treatment (n=36 pts) or uncertainty about the	1º endpoint: To investigate the value of CV tests to diagnose convulsive syncope in pts with apparent treatment-resistant epilepsy.  Results: An alternative diagnosis was found in 31 pts (41.9%), including 13 (36.1%) of 36 pts taking an anticonvulsant medication.	Limitations:  • Small sample, single center; highly unique population  Conclusions:  • A simple, noninvasive CV evaluation may identify an alternative diagnosis in many pts with

		diagnosis of epilepsy, on the basis of the clinical description of the seizures (n=38 pts)  Exclusion criteria: Suspected psychogenic nonepileptic attack disorder	<ul> <li>19 pts (25.7%) developed profound hypotension or bradycardia during the HUTT, confirming the diagnosis of VVS.</li> <li>1 patient had a typical vasovagal reaction during intravenous cannulation. 2 pts developed psychogenic symptoms during the HUTT.</li> <li>7 pts had significant ECG pauses during CSM. In 2 pts, episodes of prolonged bradycardia correlated precisely with seizures according to the insertable ECG recorder.</li> </ul>	apparent epilepsy and should be considered early in the management of pts with convulsive blackouts.
Zaidi A, et al. 1999 <u>10512777</u> (124)	Study type: Prospective cohort  Size: n=21 pts	Inclusion criteria: Recurrent seizure-like episodes and a clinical diagnosis of nonepileptic attack disorder.  Exclusion criteria: None	1° endpoint: To assess the value of HUTT as a provocative test for non-epileptic attack disorder  Results:  17 pts (81%) experienced typical symptoms (non-epileptiform limb shaking in 15 pts, absence in one patient, myoclonic jerking in one patient) during head-up tilt without significant EEG abnormalities or hemodynamic changes.	Limitations:  Small sample, select population.  Conclusions: HUT with suggestion is a safe, well tolerated, sensitive, provocative EEG test for dissociative seizure-like attacks and should be considered in pts with suspected non-epileptic attack disorder.
Luzza F, et al. 2003 <u>12846340</u> (125)	Study type: Retrospective cohort  Size: n=986 pts	Inclusion criteria: Unexplained syncope  Exclusion criteria: None	1º endpoint:  To assess the ability of HUTT in recognizing a psychiatric disorder in some pts affected by unexplained syncope.  Results:  ■ In 266 pts the test induced bradycardia and/or hypotension resulting in syncope or presyncope, allowing a diagnosis of neurally mediated syncope.  ■ In 3 other pts (0.3% of the entire population and 1% of the all positive tests) HUTT provoked LOC despite no significant change in heart rate and/or BP. In all 3 cases unconsciousness was prolonged and no pathological finding was present except lack of response. This phenomenon has been defined as 'pseudosyncope' and related to psychiatric illness	Limitations:  ■ Retrospective design, limited number of pts with pseudosyncope, lack of followup.  Conclusions:  ■ HUTT may contribute to the recognition of psychiatric disorder in some pts affected by unexplained syncope.
Tannemaat MR, et al. 2013 23873974 (126)	Study type: Prospective cohort  Size: n=800 pts	Inclusion criteria: Episode of apparent TLOC during tilt-table testing without EEG changes and without decreases in heart rate	1° endpoint: To provide a detailed semiology to aid the clinical recognition of psychogenic pseudosyncope which concerns episodes of apparent TLOC that mimic syncope.  Results:	Limitations:  Referral bias.  A clinical suspicion of PNES was not a formal exclusion criterion for tilt-table testing, but referral selection will have excluded the majority of these pts nonetheless. This may have

		or BP. The event had to be recognized by the	Of 800 tilt-table tests, 43 (5.4%) resulted in psychogenic pseudosyncope.	affected the prevalence of jerking movements.
		patient or a relative	The median duration of apparent TLOC was longer in psychogenic	Conclusions:
		(present during the test)	pseudosyncope (44 s) than in VVS (20 s, p<0.05). During the event, the	<ul> <li>Psychogenic pseudosyncope is clinically</li> </ul>
		as typical of the	eyes were closed in 97% in psychogenic pseudosyncope but in only 7%	distinct from VVS and can be diagnosed
		patient's episodes.	in VVS (p<0.0001).	accurately with tilt-table testing and
			<ul> <li>A sudden head drop or moving down the tilt table was more common</li> </ul>	simultaneous EEG monitoring.
		Exclusion criteria:	in psychogenic pseudosyncope than in VVS (p<0.01), but jerking	
		None specified.	movements occurred more frequently in VVS (p<0.0001).	
			<ul> <li>In psychogenic pseudosyncope, both heart rate and BP increased</li> </ul>	
			before and during apparent TLOC (p<0.0001).	
Moya A, et al.	Study type:	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
1995	Randomized double-	Syncope and a baseline	To assess the efficacy of oral etilefrine in preventing a positive response	Small sample, drug not used clinically in most
<u>7798528</u>	blind crossover study	positive HUTT.	to HUTT.	centers. The statistical power of the study was
(127)				only 10%.
	<u>Size</u> : n=30 pts	Exclusion criteria:	Results:	
		Previous hypertension	HUTT results were negative in 13 (43%) pts with etilefrine and 15	Conclusions:
		and 11 (11%) because	(50%) with placebo (p=NS). The rate of positive responses decreased	Oral etilefrine (10 mg 3x a day) was not
		of a cardioinhibitory	with repeated testing irrespective of the assigned treatment	superior to placebo in preventing a positive
		response to	<ul> <li>A positive response was obtained during the second HUTT in 20 pts</li> </ul>	response to HUTT. Despite a low statistical
		HUTT.	(10 with placebo, 10 with etilefrine) but in only 12 during the third (7 with	power, the high rate of negative response with
			etilefrine, 5 with placebo) (p<0.05)	placebo (50%) suggests that controlled trials are
				needed to assess the real efficacy of any
				treatment in pts with VVS.
Morillo CA, et al.	Study type:	Inclusion criteria:	1° endpoint:	Limitations:
1993	Double-blind	Recurrent neurally	To determine the efficacy of intravenous and oral disopyramide	Only pts who had a positive response were
<u>8245337</u>	randomized trial	mediated syncope and	phosphate in preventing neurally mediated syncope induced by a HUTT.	crossed over to alternative therapy.
(128)	<b>a.</b> aa .	2 or more successive		
	Size: n=22 pts,	positive HUTT	Results:	Conclusions:
	randomly allocated to	responses	HUTT results were positive for syncope in 12 (75%) of 16 pts	Intravenous disopyramide was ineffective for
	receive either	Fuelusian sultanias	receiving intravenous placebo and in 12 (60%) of 20 pts receiving	the prevention of neurally mediated syncope
	intravenous	Exclusion criteria:	disopyramide (p=0.55, 95% CI: -14%–40%).	provoked by HUTT. No significant effect was
	disopyramide or	Failure to produce	• In the intravenous phase, complete crossover was achieved in 15 pts.	observed after oral therapy with disopyramide.
	placebo	syncope or presyncope	HUTT results during this phase were positive in 13 pts (87%) receiving	
		during testing	placebo and in 12 pts (80%) receiving disopyramide (p=0.50, 95% CI: -	
			19%–32%) and were positive in all pts receiving their initially	
			randomized drug or placebo.	
	1		● In the oral phase, HUTT results were positive in only 2 pts (18%)	

Gibbons, et al. 2006 16832073 (129)	Aims: To investigate the prevalence, symptoms, and neurophysiologic features of delayed OH  Study type: Retrospective, observational, mechanistic  Size: n=230 pts	"Inclusion criteria": OH or delayed during a 60° head-up tilt performed for 45 min  Exclusion criteria: None specified	assigned to placebo and in 3 pts (27%) receiving disopyramide (p=0.54, 95% CI: -42%–24%).  • Syncope recurred in 3 (27%) of the 11 pts receiving disopyramide and 3 (30%) of the 10 pts not treated pharmacologically (p>0.05).  1° endpoint: OH or delayed OH  Results:  • Of 108 pts with OH, 46% had OH within 3 min of HUTT; 15% had OH between 3 and 10 min; and 39% had OH after 10 min of HUTT.  • Delayed OH was associated with mild sympathetic adrenergic dysfunction evident of autonomic testing	Limitations:  • Laboratory study • Referral population  Conclusions: • Delayed OH occurred in 54% of tested population • TTT duration should be extended • Underlying mechanism possibly early or mild sympathetic adrenergic failure
Podoleanu, et al. 2009 19669396 (130)	Aim: To investigated the hemodynamic mechanisms that underlie delayed OH  Study type: Prospective, casecontrol, mechanistic study in human pts  Size: n=13 pts and 9 controls	Inclusion criteria: Pts with delayed OH and (1) symptoms and signs of orthostatic intolerance after 3 mins; and (2) documentation of a delayed decrease in BP pattern during diagnostic tilt testing  Exclusion criteria: The inability of the patient to collaborate and to perform tilt testing.	1° endpoint: The changes in the SBP, heart rate, cardiac output, SV and TPR (in pts with delayed OH compared to age- and sex-matched controls during a modified version of the Italian tilt protocol.  Results: At the end of the test, in pts compared to controls, SBP was significant lower; TPR progressively decreased in pts but not in controls; SV and CO did not change in pts or in controls. Heart rate increased progressively in pts until the end of the test and remained unchanged in controls Administration of elastic compression to the legs counteracts the decrease in SBP and TPR.	Limitations:  Laboratory study Small number of pts Blinding – not stated  Conclusions: In pts with delayed OH, the progressive decrease in SBP is associated with progressive decrease in TPR, while CO and SV show little change. The compensatory increase in HR is insufficient to compensate the decline in BP Administration of elastic compression to the legs counteracts decrease in SBP and decrease in TPR.
Gurevich T, et al. 2014 <u>25531748</u> (131)	Aim: (1) To assess time- related patterns of SBP and DBP responses in pts referred for suspected OH to tilt testing	"Inclusion criteria": Syncope during angioplasty  Exclusion criteria: None specified	1° endpoint: OH or delayed OH  Results:  7% had OH within 3 min, 35% within 30 min, and 40% within 40 min. 270 OH pts, 43 and 91% were identified within 3 and 30 min, respectively	Limitations:  Referral population.  Laboratory study  Conclusions:  Tilt table testing to 30 minus identifies most but no all pts with delayed OH.

Gibbons, et al. 2015 <u>26400576</u> (132)	(2) To assess the percent of delayed OH and factors associated with it.  Study type: Prospective, observational, mechanistic,  Size: n=692 pts; 270 with OH or delayed OH  Aims: To define the long-term outcome of delayed OH  Study type: Prospective, longitudinal follow up, observational, mechanistic  Size: n=108 pts with OH, 75 age- and sexmatched controls	"Inclusion criteria": OH during a 60° head- up tilt performed for 45 mins  Exclusion criteria: None	1° endpoint: OH, delayed OH and clinical outcome including mortality  Results:  • 54% of individuals with delayed OH progressed to OH.  • 31% with delayed OH developed an α-synucleinopathy  • 10-y mortality rate in individuals with delayed OH was 29%; with baseline OH was 64% and in controls was 9%.  • 10-y mortality of individuals who progressed to OH was 50%.	Limitations:  Laboratory study Referral population  Conclusions: Delayed OH frequently progresses to OH Delayed OH frequently progresses to an alpha-synucleinopathy (multiple system atrophy, Parkinson's disease, dementia with Lewy bodies) Delayed OH has a high associated mortality particularly when it progresses to OH
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## Data Supplement 16. Nonrandomized Trials, Observational Studies, and/or Registries of Neurologic Investigation – (Section 3.3)

Study Acronym; Author;	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results	Summary/ Conclusion
Year Published				Comment(s)
Abubakr A, et al.	Study type: Retrospective	Inclusion criteria: Syncope pts	1° endpoint: Classification of EEG findings	Very few abnormal EEGs, but the larger
2005	chart review	selected from a larger population of	including variants of normal.	population of syncope pts is not reported.
<u>15820355</u>		EEG reports		Rare EEG abnormalities. No epileptiform
(133)	Size: n=1,094 syncope pts		Results: 2 (1.5%) abnormal EEGs: one focal	features
			slowing, one diffuse slowing	

Al-Nsoor, et al. 2010	Study type: Perhaps prospective cohort	Inclusion criteria: Syncope in ED seen by a neurologist	1° endpoint: Abnormality contributing to diagnosis	Very high use of CT scans, and firmness of attribution not clear
<u>20672498</u> (134)	<u>Size</u> : n=292 pts	, ,	Results: 254 CT scans (87%); 10 (3.9% of ordered) helped.	
Giglio P, et al. 2005 16292675	Study type: Retrospective chart review	Inclusion criteria: Syncope pts in ED	1° endpoint: Proportion with CT scans; proportion abnormal related to syncope	Fully 34% had CT, but only 1 (3% of ordered) had diagnostic utility relevance
(135)	<u>Size</u> : n=128 pts		Results: 44 had CT; 1 showed old posterior infarction.	
Goyal N, et al. 2006 <u>17111790</u> (136)	Study type: Retrospective chart review  Size: n=117 pts with syncope	Inclusion criteria: Syncope diagnosis by ED MD	1° endpoint: Any clinically significant finding  Results: 117 had CT; 0 (0% of ordered) helped.	Inclusion criteria based on CT use, but the larger population of syncope pts is not reported. CT had no diagnostic utility
	and head CT			
Johnson PC, et al. 2014 25365440	Study type: Retrospective chart review	Inclusion criteria: Syncope coded in billing records, and after non-syncopal diagnoses excluded on chart review	1º endpoint: Test contributed to alleged diagnosis	CT and MRI performed moderately frequently and of no diagnostic utility. Carotid ultrasound less frequently and of no
(137)	<u>Size</u> : n=167 syncope pts of 1,038 adult Texan in-pts with "syncope" screened		Results: 131 CT scans (78.4%); 0% helped. 18 brain MRI (10.7%); 0% helped. 52 carotid ultrasounds (31.1%); 0% helped.	diagnostic utility.
Kapoor WN, et al. 1983 6866032	Study type: Prospective cohort	Inclusion criteria: Diagnosis of syncope after inclusion for TLOC	1º endpoint: Diagnosis of cause of syncope  Results: 65 CT scans (32%); 0% helped. 101	The population was accumulated nearly 40 y ago. Tests are of minimal diagnostic utility.
(70)	<u>Size</u> : n=204 pts in global population		EEGs (49.5%); 1 (1% of ordered) helped.	
Mecarelli O, et al. 2004 <u>15639129</u> (138)	Study type: prospective observational controlled cohort	Inclusion criteria: recurrent syncope, positive tilt test, negative brain MRI	1º endpoint: Abnormal EEG  Results: 0 (0%) abnormal findings on routine EEG but increased slow wave activity during	The report is restricted to VVS pts, and is only one of several. Maybe should delete it, or include them all.
,	Size: 43 pts with vasovagal syncope; 32 controls		hyperventilation	
Mendu ML, et al. 2009 19636031	Study type: Retrospective chart review	Inclusion criteria: ICD 9 in-hospital primary or secondary syncope diagnosis	1° endpoint: Chart documentation that the finding contributed to the diagnosis	One of the largest, but retrospective, firmness of attribution not clear. CT, EEG, carotid ultrasound of minimal
(68)	<u>Size</u> : n=1,920 pts		Results: 1327 CT scans (63%); 35 (2.6% of ordered) helped. 154 brain MRI (19%); 23 (15%)	diagnostic utility. MRI provided some diagnostic utility

Pires LA, et al. 2001 11493131	Study type: Retrospective chart review	Inclusion criteria: ICD 9 syncope in in-patients	of ordered) helped. 267 carotid ultrasounds (20%); 3 (1.1% of ordered) helped. 174 EEG (13%); 3 (1.7%) helped  1º endpoint: Apparently contributed to diagnosis of etiology.	Weak methodology. All investigations of low diagnostic utility
(139)	<u>Size</u> : n=649 pts		Results: 283 CT scans (41%); 5 (1.8% of ordered) helped. 10 brain MRI (1.3%); 3 (30% of ordered) helped. 185 carotid ultrasounds (29%); 0 (0% of ordered) helped. 253 EEG (39%); 6 (2.4%) helped	
Poliquin-Lasnierl L, et al. 2009 19960758 (140)	Study type: Retrospective chart review  Size: n=517 pts	Inclusion criteria: Syncope or falls and EEG ordered	1° endpoint: "Yield" of EEGs  Results: 0 (0%) EEGs showed epileptiform activity	EEG use an inclusion criterion, so studied population. does not represent the large syncope population
Scalfani JJ, et al. 2010 <u>20625024</u> (141)	Study type: Part A retrospective chart review: Part B prospective post-CME cohort  Size: Part A 721; Part B 371	Inclusion criteria: ICD primary or secondary diagnosis of syncope	1° endpoint: Causative finding defined as probably contributing to syncope, OR identifying a high risk subject for arrhythmic death  Results: 583 CT scans (53%); 14 (2.4% of	Pooled sequential 2-stage study
	pts; pooled 1092 because CME had no effect on test ordering		ordered) helped. 208 brain MRI (19%); 12 (5.8% of ordered) helped. 57 carotid ultrasounds (0%); 0 (0% of ordered) helped.	
Sheldon, et al. 1982 <u>9676166</u> (142)	Study type: Prospective observational  Size: n=18 pts	"Inclusion criteria": Syncope or presyncope during head up tilt with isoproterenol provocation  Exclusion criteria: None specified	1º endpoint: EEG changes during syncopal episodes  Results: ■ No pts developed EEG abnormalities before the onset of presyncope,	Limitations:  • Laboratory study  • Unblinded  • Small number of pts  Conclusions:
			<ul> <li>During presyncope, theta wave slowing (8/14) and delta wave slowing (9/14), and background suppression (1/14) were noted During syncope, theta wave slowing (9/18) and delta wave slowing (11/18), and background suppression (6/18) were noted</li> <li>Abrupt changes in the EEG rhythm occurred</li> </ul>	Presyncope and syncope are associated with EEG abnormalities     No single EEG pattern is pathognomonic of presyncope of syncope     The transition from presyncope to syncope is marked by abrupt EEG changes.

		T		
L. DA ()	A*	Last de la desta	within 15 s of the transition to syncope (14/18)	11.4.6.
Low PA, et al.	Aims:	Inclusion criteria:	"1° endpoint":	<u>Limitations</u> :
2004	To estimate autonomic	Known diabetes and willingness to	Autonomic symptoms and test results	<ul> <li>Single region and demographic</li> </ul>
<u>15562211</u>	symptoms and deficits using a	complete general medical and		
(143)	laboratory evaluation of	neurological evaluations, and a full	Results:	
	autonomic function and a	autonomic reflex laboratory evaluation	• OH in 8.4 and 7.4% of type 1 and type 2	Conclusions:
	validated self-report measure	annually	diabetes, respectively (using the criterion of 30	<ul> <li>Autonomic symptoms and deficits are</li> </ul>
	of autonomic symptoms in pts		mmHg SBP)	common in diabetes, but mild in severity
	and matched control pts from	Exclusion criteria:	• OH in 22.9 and 16.2% of type 1 and type 2	The correlation between symptom scores
	the population	None specified	diabetes, respectively (using the criterion of 20	and deficits is overall weak in mild diabetic
			mmHg SBP).	neuropathy, emphasizing the need to
	Study type:		Autonomic neuropathy, defined using a	separately evaluate autonomic symptoms
	Cross-sectional; population		composite testing score, was present in 54% of	and objective tests.
	based, observational		type 1 and 73% of type 2 pts	
	Since m=224 ata with DM			
	Size: n=231 pts with DM (type 1, n=83; type 2, n=148)			
	and n=245 control pts			
Kim, et al.	Aims:	Inclusion criteria:	(40	Limitations:
2009	To assesses the value of	A diagnosis of transthyretin amyloid	"1° endpoint":	• Laboratory study
19618439	standard quantitative	polyneuropathy	Autonomic and sensory test results	Referral population
(144)	autonomic and sensation		Results:	Small number of pts
(144)	tests in detecting,	Exclusion criteria:	Abnormal postganglionic sympathetic	Sitial fluttiber of pts
	characterizing, and	None specified	sudomotor dysfunction was found in 74%	
	quantitating the severity of	Trone specified	• The HRdb was abnormal in 25 (69%)	Conclusions:
	transthyretin amyloid		OH present in 13 pts (36%)	This study provides a rationale for the use
	polyneuropathy		Median SBP fall of 36 mmHg at 1 min (range)	of quantitative autonomic and sensory
	polynouropauty		32–80 mm Hg	testing as standard, objective, and
	Study type:		52 55 mm rig	quantitative measures for assessing the
	Retrospective, observational			severity of TTR-A-PN
	' '			,
	Size: n=36 pts			
lodice V, et al.	Aim:	Inclusion criteria:	1° endpoint:	<u>Limitations</u> :
2012	To evaluate the autonomic	Autopsy confirmed cases of MSA who	Autonomic test results, clinical features	None
<u>22228725</u>	characterization of MSA in	had undergone formal autonomic		
(145)	autopsy confirmed cases	testing, including adrenergic, sudomotor	Results:	Conclusions:
		and cardiovagal functions and	OH was present in 21 pts and symptomatic in	Severe and progressive generalized
	Study type:	Thermoregulatory Sweat Test	19 pts	autonomic failure with severe adrenergic

Thaisetthawatkul P, et al.	Retrospective, observational, autopsy study in human pts  Size: n=29 pts  Aim: To assess autonomic function	Exclusion criteria: None listed.  Inclusion criteria: Clinically probable dementia with Lewy	Norepinephrine normal supine (203.6±112.7pg/ml). Orthostatic increment of was reduced (33.5±23.2%) Severe generalized autonomic failure in most pts 20/22 had anhidrosis and 18 had thermoregulatory sweat test % anhidrosis >30%  1º endpoint: Autonomic test results, clinical features	and sudomotor failure combined with the clinical phenotype is highly predictive of MSA.  Limitations: Referral bias
2004 <u>15159482</u> (146)	in pts with dementia with Lewy bodies  Study type: Retrospective, observational study in human pts  Size: n=20 DLB pts, 20 age- matched MSA and PD pts	bodies and MSA pts and clinically definite PD pts  Exclusion criteria: Coexistent conditions, such as diabetes, that account for the symptoms of dysautonomia.	Results:  OH present in 10/20 dementia with Lewy bodies, 17/20 MSA, and 1/20 PD pts  Most common abnormal TST pattern in dementia with Lewy bodies was distal pattern, found in 54% of pts; while in MSA the most common pattern was global pattern, found in 41% of pts	Clinical diagnoses Autonomic testing in demented pts  Conclusions: Autonomic dysfunction is frequent in dementia with Lewy bodies and the severity is intermediate between that of multiple system atrophy and Parkinson disease.
Thieben MJ, et al. 2007 17352367 (147)	Aim: To evaluate the prevalence and pathogenetic mechanisms of POTS  Study type: Observational, retrospective, mechanistic  Size: n=152 pts	Inclusion criteria: Baseline sinus rhythm with no evidence of arrhythmia or cardiac disease, sustained heart rate increment of 30 beats/min or greater in response to 10 mins of head-up tilt, and symptoms of orthostatic intolerance Symptoms present for more than 3 mo.  Exclusion criteria: (1) OH defined as a decline of 30 mm Hg or more in SBP or 20 mm Hg or more in mean BP within 3 mins of standing or HUTT; (2) pregnancy or lactation; (3) presence of another cause of autonomic failure	1° endpoint: Autonomic test results, clinical features  Results:  • Mean orthostatic heart rate increment was 44 beats/min.  • 50% of pts had sudomotor abnormalities (apparent on both the quantitative sudomotor axon reflex test and TST),  • 34.9% had significant adrenergic impairment	Limitations:  Referral population. Laboratory study  Conclusions: Findings suggest a neuropathic basis for at least half the cases of POTS

Gibbons C, et al. 2013 24386408 (148)	Aim: To define the neuropathology, clinical phenotype, autonomic physiology and differentiating features in individuals with neuropathic and nonneuropathic POTS.  Study type: Observational, mechanistic  Size: n=24 pts and 10 controls	Inclusion criteria: POTS was defined as an increase in heart rate of >30 beats per min upon standing with symptoms of orthostatic intolerance, without any known medical condition or medication causing the tachycardia  Exclusion criteria: DM, impaired glucose tolerance, vitamin deficiencies, heavily metal toxicity, thyroid disorders, pheochromocytoma, hypoadrenalism, anxiety, cardiac disease, volume depletion, drug abuse and medication side effect	1° endpoint: Autonomic test results, clinical features, nerve density from skin biopsy  Results: • Pts with neuropathic POTS and had significantly lower resting and tilted heart rates; reduced parasympathetic function; and lower phase 4 Valsalva maneuver overshoot compared with those with non-neuropathic POTS	Limitations:  ■ Referral population.  ■ Laboratory study  Conclusions:  ■ POTS subtypes may be distinguished using small fiber and autonomic structural and functional criteria.
Martinez-Fernandez, et al. 2008 17974603 (149)	Study type: Prospective Registry  Size: n=359 pts	Inclusion criteria: Symptomatic pts with TIA or non- invalidating stroke, asymptomatic pts. with 85% stenosis, TCD detected microemboli/ exhausted CVR or silent lesions  Exclusion criteria: N/A	1° endpoint: Occurrence of CSR and/or syncope during internal CAA  Results: CSR and syncope occurred in 62.7 % and 18.6% of pts. EEG changes more prominent in pts. with cardio-inhibitory syncope, Syncope is more frequent in cardio-inhibitory CSR (p<0.001), Risk of syncope during CAA in pts with CSR (OR: 4.2; 95% CI:1.9–9.1) Risk of syncope in pts. with cardio-inhibitory CSR and vasodepressor/mixed CSR (OR: 6.9; 95% CI: 3.2–15.0 and OR: 1.4; 95% CI: 0.6–3.7) respectively.	Syncope is common in pts undergoing CAA and can be misdiagnosed as frontal seizures, cardio-inhibitory response most frequent mechanism of syncope.     Limitations: Beat to beat analysis of BP was not performed.
Gibbons, et al. 2015 <u>26400576</u> (132)	Aims: To define the long-term outcome of delayed OH  Study type: Prospective, longitudinal follow up, observational, mechanistic  Size: n=108 pts with OH, 75	"Inclusion criteria": OH during a 60° HUTT performed for 45 mins  Exclusion criteria: None specified	1º endpoint: OH, delayed OH and clinical outcome including mortality  Results:  • 54% of individuals with delayed OH progressed to OH.  • 31% with delayed OH developed an α-synucleinopathy	Limitations:  Laboratory study Referral population  Conclusions: Delayed OH frequently progresses to OH Delayed OH frequently progresses to an alpha-synucleinopathy (multiple system

age- and sex-matched controls	<ul> <li>10 y mortality rate in individuals with delayed OH was 29%; with baseline OH was 64% and in controls was 9%.</li> <li>10 y mortality of individuals who progressed to OH was 50%.</li> </ul>	atrophy, Parkinson's disease, dementia with Lewy bodies)  • Delayed OH has a high associated mortality particularly when it progresses to OH

## Data Supplement 17. Nonrandomized Trials, Observational Studies, and/or Registries of ARVCD – (Section 4.2.4)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Corrado D, et al. 2003 14638546 (150)	Study type: Retrospective  Size: n=132 pts	Inclusion criteria: ARVC pts treated with ICD  Exclusion criteria: ARVC with only minor criteria, idiopathic RV VT, myocarditis, IDCM, Uhl's anomaly	1° endpoint: ICD treated arrhythmia  Results: of 132 pts, 64 (48%) had appropriate ICD intervention in FU of 39 mo. Of 21 pts with syncope 8 (38%) had appropriate ICD therapy including 5 with VFL/VF.	Unexplained syncope had an OR of 7.5 for appropriate ICD interventions (p=0.07; 95% CI: 0.84–1.81)
Corrado D, et al. 2010 <u>20823389</u> (151)	Study type: Retrospective Size: n=106 pts	Inclusion criteria: ARVC pts receiving ICDs  Exclusion criteria: Prior sustained VT or VF	1° endpoint: Appropriate ICD interventions.  Results: Of 106 pts 25 (24%) had appropriate ICD interventions in f/u of 58 mo. Pts presenting with syncope had a 9%/y incidence of appropriate ICD intervention.	Syncope independently predicted for an appropriate ICD shock (HR: 2.94; 95% CI: 1.83 to 4.67; p=0.013) and shocks for VF/VFL (HR: 3.16; 95% CI: 1.39–5.63; p=0.005).
Bhonsale A, et al. 2011 21939834 (152)	Study type: Retrospective Size: n=84 pts	Inclusion criteria: ARVD/C pts receiving ICDs  Exclusion criteria: Prior sustained VT or VF	1° endpoint: Appropriate ICD interventions  Results: Appropriate ICD therapy in 40 (48%) in f/u of 4.7 y. Of 23 pts presenting with syncope 10 (25%) had appropriate ICD interventions	Syncope was not a predictor of appropriate ICD intervention
Bhonsale A, et al. 2013 23671136 (153)	Study type: Retrospective Size: n=215 pts	Inclusion criteria: Diagnosed with ARVD/C  Exclusion criteria: None	1° endpoint: SCD, sustained arrhythmia, appropriate ICD intervention  Results: 86 (40%) had primary endpoint in mean f/u of 7 y. Of 41 pts with syncope, the primary endpoint was met in 30 (73%).	Symptomatic pts (syncope, presyncope and palpitation) predicted for ventricular arrhythmias (p<0.001).
Link MS, et al. 2014	Study type: Prospective observational	Inclusion criteria: ARVD/C	<u>1° endpoint</u> : Sustained ventricular arrhythmias	Syncope was not a predictor of VA.

<u>25011714</u>		Exclusion criteria: Sarcoid cardiac	Results: 48 pts with VA. Of 28 pts with	
(154)	Size: n=137 pts; 108 with ICDs	disease	syncope 14 (50%) met primary endpoint	
Corrado D, et al.	Study type: Consensus	Inclusion criteria: None	1° endpoint: None	• In ARVC pts with syncope an ICD should be
2015	statement			considered
<u>26216213</u>		Exclusion criteria: None	Results: None	
(155)	Size: None			

## Data Supplement 18. Nonrandomized Trials, Observational Studies, and/or Registries of Sarcoid Heart Disease – (Section 4.2.5)

Study Acronym Author, Year	Study Type/Design*; Study Size	Patient Population	Primary Endpoint and Results	Summary/ Conclusion Comment(s)
Winters SL, et al. 1991 <u>1894867</u> (156)	Study type: Retrospective Size: n=7 pts	Inclusion criteria: Documented (n=6) or highly suspected (n=1) Sarcoidosis with ECG abnormalities	1° endpoint: Findings during EPS  Results: Sustained VT was easily inducible in all pts. Steroid therapy did not prevent spontaneous VT. Despite anti-arrhythmic therapy, 2 pts had SCD and an additional 4 recurrent VT.4 pts received an ICD and all 4 received appropriate therapy.	Poor response to ant-arrhythmic drug therapy     ICD therapy is recommended as primary therapy in pts with sarcoidosis and VT
Koplan, et al. 2006 <u>16876741</u> (157)	Study type: Retrospective Size: n=8 pts	Inclusion criteria: Cardiac sarcoidosis with recurrent VT	1° endpoint: To define the clinical characteristics of pts with CS and the EP findings during EPS.  Results: All pts had a reduced LVEF except for 1 pts (Mean 34% ± 15%) and had failed previous anti-arrhythmic drug therapy.  EPS revealed evidence of scar-related reentry with multiple morphologies. Areas of low-voltage scar were present in the RV in all 8 pts. Ablation was only partially helpful. 5 out of 8 pts eventually required cardiac transplantation.	<ul> <li>Sarcoidosis can be misdiagnosed as idiopathic VT or ARVD.</li> <li>Catheter ablation is only partially successful.</li> </ul>
Jefic, et al. 2009 <u>19187909</u> (158)	Study type: Retrospective Size: n=42 pts	Inclusion criteria: CS	1° endpoint: To determine response to medical therapy and radiofrequency ablation  Results: In 9 out of 21 pts with VT/VF recurrence post-ICD implant, drug therapy was ineffective requiring radiofrequency ablation. The most frequent VT circuit was reentry in the pertricuspid area. All pts had either a decrease (n-=4) or complete elimination (n=5) during follow up (19.8 ± 19.6 mo).	In pts with CS and refractory VT, catheter ablation is effective in eliminating or reducing the VT burden.

Furushima, et al. 2004 <u>15119697</u> (159)	Study type: Retrospective Size: n=8 pts	Inclusion criteria: CS and sustained monomorphic VT	1º endpoint: Mechanism and outcome of VT associated with cardiac sarcoidosis  Results: Most VT is due to reentry. The inducibility rate depends on the presence or absence of an active phase. ICD therapy is effective.	While most VT is due to reentry, inducibility depends on the disease state including response to immunosuppressive therapy.
Hiramitsu S, et al. 2005 <u>16315784</u> (160)	Study type: Questionnaire survey  Size: n=49 pts	Inclusion criteria: CS treated with steroid therapy	<u>Results:</u> The most common initial steroid dose used was 30 mg/day or 60 mg on alternate days. This dose was continued for 1 mo followed by tapering by 5mg every 2 to 4 wk until reaching the maintenance dose of 5–10 mg/d.  Steroid therapy was reported to result in improvement in 54%, no change in 40%, and deterioration in 6% of cases.	There is a fairly uniform use of steroid therapy in the management of CS with clinical improvement in over one-half of the cases.
Kandolin R, et al. 2011 <u>21427276</u> (161)	Study type: Retrospective study  Size: n=72 pts	Inclusion criteria: Unexplained AV block	1° endpoint: To determine the prevalence of CS and giant cell myocarditis in young and middle-aged adults undergoing pacemaker implantation for AV block  Results: CS and giant cell myocarditis were found in 14 (19%) and 4 (6%) pts, respectively. The majority (16/18, 89%) were women.  Over an average of 48 mo of follow-up, 7 (39%) of 18 pts with CS or giant cell myocarditis vs. 1 of the 54 pts in whom AV block remained idiopathic, experienced either cardiac death, cardiac transplantation, VF, or treated sustained VT (p<0.001).	<ul> <li>CS and giant cell myocarditis account for &gt;25% of young and middle-aged adults presenting with AV block.</li> <li>These pts are at high risk of having major adverse events.</li> </ul>
Chapelon-Abric C, et al. 2004 15525844 (162)	Study type: Retrospective Size: n=41 pts	Inclusion criteria: CS	1º endpoint: Clinical characteristics and response to therapy  Results: Cardiac signs were clinical in 63% of cases and electrical in 22%.  During an average follow up of 58 m, 87% of pts showed improvement on immunosuppressive therapy and 54% were cured from a clinical and laboratory point of view.	Most pts with CS respond to immunosuppressive therapy.
Yodogawa K, et al. 2011	Study type: Retrospective	Inclusion criteria: CS and VA	1º endpoint: Efficacy of corticosteroid therapy in the treatment of VA	Corticosteroid therapy may be effective for VA in the early stage, but

21496164	Size: n=31 pts	T		is loss affective in the late stage
(163)	<u>σιze</u> : π–στ μιs		Results: Overall, there were no significant differences in the number of PVCs and in the prevalence of NSVT before and after steroid therapy. However, in pts with LVEF ≥ 35% (n=17), there was a significant reduction in the number of PVCs (from 1820 ± 2969 to 742 ± 1425, p=0.048) and in the prevalence of NSVT (from 41 to 6%, p=0.039).  The less advanced LV dysfunction group showed a significantly higher prevalence of gallium-67 uptake compared with the advanced LV dysfunction group (LVEF <35 %, n=14). In the	is less effective in the late stage.
			advanced LV dysfunction group (LVEF \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	
			differences in these parameters.	
Schuller JL, et al. 2012	Study type: Retrospective	Inclusion criteria: CS and ICD for primary or secondary	1° endpoint: ICD therapy in pts with CS	• Almost one-third of pts with CS and ICD receive appropriate therapies.
<u>22812589</u>	<u>Size:</u> n=112 pts	prevention of sudden death	<b>Results:</b> Over a mean follow up period of 29.2 mo, 32.1% of pts	<ul> <li>Adjusted predictors for ICD therapies</li> </ul>
(164)			received appropriate therapies. VT storms and inappropriate therapies occurred in 14.2 % and 11.6% of pts respectively.	included left or right ventricular dysfunction.
			Covariates associated with appropriate ICD therapies included LVEF <55% (OR: 6.52; 95% CI: 2.43–17.5), right ventricular dysfunction (OR: 6.73; 95% CI: 2.69–16.8), and symptomatic HF (OR: 4.33 95% CI: 1.86–10.1).	
Betensky BP, et al. 2012 22338670	Study type: Retrospective  Size: n=45 pts	Inclusion criteria: CS and ICD for primary or secondary prevention of sudden death	<u>1° endpoint:</u> To determine the prevalence and incidence of ventricular tachy-arrhythmias in pts with CS and to identify predictors of appropriate therapy	The annual incidence rate for appropriate ICD therapy is 15%.
(165)			Results: Appropriate and inappropriate ICD therapies were observed in 37.8% (15% per y) and 13.3% of pts, respectively.	Longer follow-up, left ventricular systolic dysfunction, and complete heart block were associated with appropriate ICD therapy.
			Longer ICD follow-up $(4.5 \pm 3.1 \text{y vs. } 1.5 \pm 1.5 \text{y; p=0.001})$ , depressed left ventricular EF $(35.5\% \pm 15.5\% \text{ vs. } 50.9\% \pm 15.5\%; \text{p=0.002})$ , and complete heart block $(47.1\% \text{ vs. } 17.9\%; \text{p=0.048})$ were associated with appropriate ICD therapy.	
Kron J, et al. 2013 23002195	Study type: Retrospective  Size: n=235 pts	Inclusion criteria: Consecutive pts with CS and ICD	1° endpoint: To evaluate the efficacy and safety of ICD therapy in pts with CS	Almost a third of pts with CS and ICD receive appropriate ICD therapy over a mean follow-up of 4.2 ± 4.0 y.
(166)	<u>σίες</u> . 11-200 μισ		Results: Over a mean follow-up of 4.2 ± 4.0 y, 36.2% pts	0 0 0 1 1 0 1 0 1 0 1 0 1 0 1 0 1 4.2 ± 4.0 y.

			received an appropriate ICD therapy and 24.3% received inappropriate shocks.  Pts who received appropriate ICD therapies were more likely to be male (73.8 vs. 59.6%, p=0.0330), have a history of syncope (40.5 vs. 22.5%, p=0.0044), lower LVEF (38.1 ± 15.2 vs. 48.8 ± 14.7%, p≤0.0001), ventricular pacing on baseline ECG (16.1 vs. 2.1%, p=0.0002), and a secondary prevention indication (60.7 vs. 24.5%, p<0.0001) compared with those who did not receive appropriate ICD therapies.	Predictors of appropriate ICD therapies include a history of syncope, depressed LV function and ventricular pacing.
Mehta D, et al. 2011 <u>21193539</u> (167)	Study type: Retrospective Size: n=76 pts	Inclusion criteria: Evidence of CS but without symptoms	1º endpoint: To assess the role of programmed electrical stimulation study in risk assessment in pts with sarcoidosis  Results: 11% of pts were inducible and received an ICD (LVEF 36.4±4.2% vs. 55.8±1.5%, p<0.05).  Over a median follow-up of 5 y, 6 of 8 pts in the group with inducible VA had ventricular arrhythmia or died, compared with 1 death in the negative group (p<0.0001).	Programmed electrical stimulation may help identify pts with CS who are at risk of having ventricular arrhythmias.

# Data Supplement 19. Nonrandomized Trials, Observational Studies, and/or Registries of Brugada Syndrome – (4.3.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Morita H, et al. 2008 18838563	Study type: Retrospective Size: n=115 pts	Inclusion criteria: Symptomatic and asymptomatic BS	<u>1° endpoint</u> : Prevalence of fragmented QRS and its prognostic value	<ul> <li>Fragmented QRS appears to be a marker for spontaneous VF and syncope</li> </ul>
(168)		Exclusion criteria: N/A	Results: Fragmented QRS was more prevalent in pts with VF (85%) and syncope (50%) when compared to asymptomatic pts (34%).	5,
Gehi, et al. 2006 <u>16836701</u> (169)	Study type: Meta-analysis assessing predictors of cardiac events  Size: n=1,545 pts	Inclusion criteria: Studies were included if they m, et al.I of the following criteria: 1) prospective cohort studies of the natural history of pts with Brugada-type ECG, 2) studies included >10 pts, 3) primary data on cardiac events was provided	1° endpoint: SCD, syncope and ICD shock  Results: The overall rate was 10% over an average of 32 mo. Predictors of adverse events included  • Syncope and SCD (RR: 3.24; 95% CI: 2.13–4.93)  • Men compared with women (RR: 3.47; 95% CI: 1.58–7.63), and	Male sex, spontaneous Type I ECG pattern and Hx of SCD and syncope are good predictors of future cardiac events

Benito B, et al. 2008 19007594 (170)	Study type: Prospective follow up study Size: n=384 pts	and 4) stated clearly that structural heart disease was ruled out.  Exclusion criteria: If not all inclusion criteria are met  Inclusion criteria: Pts with BS  Exclusion criteria: N/A	Spontaneous compared with drug-induced Type I ECG (RR: 4.65; 95% CI: 2.25–9.58)      1º endpoint: To assess phenotype and prognosis differences between men and women      Results: Men had greater rates of spontaneous Type 1 ECG, ST elevation and VF inducibility (p<0.001), syncope (18% vs. 14%) and aborted SCD (6% vs. 1%).      Conversely, conduction parameters and QTc increased more in women in response to Na channel	<ul> <li>Men with BS present with a greater risk clinical profile than women and have a worse prognosis.</li> <li>Conduction disturbances may be a marker of risk in the female population</li> </ul>
Morita H, et al. 2008 <u>18838563</u> (168)	Study type: Retrospective Size: n=115 pts	Inclusion criteria: Symptomatic and asymptomatic BS  Exclusion criteria: N/A	blocker.  1º endpoint: Prevalence of fragmented QRS and its prognostic value  Results: Fragmented QRS was more prevalent in pts with VF (85%) and syncope (50%) when compared to asymptomatic pts (34%).	Fragmented QRS appears to be a marker for spontaneous VF and syncope
Sarkozy, et al. 2011 <u>21727093</u> (171)	Study type: Registry  Size: n=280 consecutive pts	Inclusion criteria: Type 1 ECG pattern  Exclusion criteria: N/A	1º endpoint: Prevalence of family history of SD and its prognostic value  Results: SD was present in 69 out of 157 families (43%). During follow-up VF or SD-free survival rate was not different between pts with or without a family Hx of SD of a first-degree relative, between pts with or without a family Hx of multiple SD of a first-degree relative at any age and between pts with or without a family Hx of SD in first-degree relatives' ≤35 y of age.	Family Hx of SD is not predictive for future arrhythmic events even if considering only SD in first-degree relatives or SD in first-degree relatives at a young age.
PRELUDE Registry. Priori SG, et al. 2012 22192666 (172)	Study type: Registry Size: n=308 pts	Inclusion criteria: Spontaneous or drug-induced type 1 ECG  Exclusion criteria: Hx of cardiac arrest	1º endpoint: Arrhythmic events in pts with and without inducible VT/VF      Results: During a median follow up of 34 mo, there were 14 arrhythmic events. 9/14 occurred in non-inducible pts.      Arrhythmia inducibility was not a predictor of	VT/VF inducibility is unable to identify high-risk pts, whereas the presence of a spontaneous type I ECG, Hx of syncope, ventricular effective refractory period <200 ms, and QRS fragmentation seem useful to identify candidates for primary

			T	
			arrhythmic events.	prevention ICD implants.
			Syncope and spontaneous Type I ECG (HR: 4.20)	
			and VERP<200ms (HR:3.91), fragmented QRS (HR:	
			4.94) were significant predictors of arrhythmias.	
Sacher F, et al.	Study type: Multicenter	Inclusion criteria: BS with ICD	1° endpoint: Appropriate shocks and ICD	The annual rate of appropriate ICD
2006	outcome report	implant	complications including inappropriate shocks	therapy is low. Appropriate ICD shocks
<u>17116772</u>				are more frequent in symptomatic than
(173)	Size: n=220 pts including	Exclusion criteria: N/A	Results: During a mean follow-up of 38±27 mo, no	in asymptomatic pts (12% vs. 4%;
	88 with syncope		pts died and 18 pts (8%) had appropriate device	p=0.05).
			therapy. The annual event rate was 2.6% with an	
			annual complication rate of 8.9%.	<ul> <li>Not all syncope in pts with BS is</li> </ul>
			·	arrhythmic.
			In pts with syncope, 10% received an appropriate	,
			shock during a 19.5–59 mo FU period. 7% had	
			syncope recurrence without any documented	
			arrhythmia. The HR for asymptomatic vs. syncope pts	
			was 0.43 (CI: 0.24–0.74).	
Sarkozy, et al.	Study type: Retrospective	Inclusion criteria: Spontaneous or	1° endpoint: Appropriate and inappropriate ICD	The authors could not confirm that
2007	single center study	drug induced Type 1 ECG pattern	shocks.	syncope was an independent predictor
<u>17251258</u>		BrS with syncope (n=26) and/or +		of appropriate ICD shocks.
(174)	Size: n=47 pts	family Hx (n=26) who underwent ICD	<b>Results:</b> During a median follow up of 47.5 mo, 7 pts	4 pts had recurrent syncope with no
		implant for primary prevention	(15%) had appropriate shocks. All were male (3	documented arrhythmia suggesting a
			syncope, 3 + family Hx and 1 had both). 4 pts had	reflex mediated mechanism.
		Exclusion criteria: N/A	recurrent syncope with no documented arrhythmia.	
			Spontaneous Type 1 ECG pattern and NSVT were	
			more frequent among pts with appropriate shocks	
Rosso R, et al.	Study type: Retrospective	Inclusion criteria: BS pts with ICD	1° endpoint: Efficacy and complications of ICD	Appropriate device therapy was
2008	multicenter study (12 Israeli	implants: Cardiac arrest (18.6%),	therapy	limited to CA survivors while none of
<u>18669142</u>	centers)	syncope (52.5%), inducible VF in		the other pts including those with
(175)	,	asymptomatic (23.7%), and positive	Results: During FU (4–160 mo), 5/11 pts with CA	syncope and/or inducible VF suffered
	Size: n=59 pts	family Hx of SD (0.5%)	had appropriate device therapy. None of the pts	an arrhythmic event.
		, ,	without prior CA had appropriate device therapy.	,
		Exclusion criteria: N/A		
FINGER Brugada	Study type: Registry from	Inclusion criteria: Pts with	1° endpoint: SCD	Low event rate even in pts with
Syndrome Registry	11 tertiary centers in 4	spontaneous or drug-induced Type 1		syncope
Probst V, et al.	European countries:	ECG pattern	Results: The cardiac event rate per y was 7.7% in	• Family Hx, inducibility of VT/VF and
2010	France, Italy, Netherlands,		pts with aborted SCD, 1.9% in pts with syncope, and	the presence of SCN5A mutation were
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20100972	Germany Registry	Exclusion criteria: Diseases that	0.5% in asymptomatic pts.	not predictive of arrhythmic events.
(176)	(FINGER)	mimic BS	Symptoms and spontaneous type 1 ECG were	
			predictors of arrhythmic events, whereas sex, familial	
	Size: n=1029 consecutive		Hx of SCD, inducibility of VT during EPS, and the	
	pts		presence of an SCN5A mutation were not predictive	
			of arrhythmic events.	
Conte, et al.	Study type: Retrospective	Inclusion criteria: Pts with	1° endpoint: Appropriate and inappropriate shocks	<ul> <li>ICD therapy was an effective</li> </ul>
2015	single center	spontaneous or drug-induced Type 1	and device complications	strategy in BS, treating potentially
<u>25744005</u>		ECG pattern who underwent ICD		lethal arrhythmias in 17% of pts during
(177)	<u>Size</u> : n=176 pts	implantation.	Results: During a mean follow-up period of 83.8 ±	long-term follow-up. Risk stratification
			57.3 mo, spontaneous sustained VAs occurred in 30	by EPS may identify asymptomatic pts
		Exclusion criteria: N/A	pts (17%). 8 pts (4.5%) died.	at risk for arrhythmic events and could
			<ul> <li>Appropriate ICD shocks occurred in 28 pts (15.9%),</li> </ul>	be helpful in investigating syncope not
			and 33 pts (18.7%) had inappropriate shocks.	related to VAs.
			Electrical storm occurred in 4 pts (2.3%).	
			28 p pts (15.9%) experienced device-related	
			complications.	
			• 105 (59.7%) pts had syncope with 53 (50.4%)	
			having a family Hx of SD. Spontaneous Type 1	
			pattern was present in 18.1%. Appropriate and	
			inappropriate shocks occurred in 10.5% and 17.1% of	
			cases.	
			<ul> <li>In multivariate Cox regression analysis, aborted</li> </ul>	
			SCD and VA inducibility on EP studies were	
			independent predictors of appropriate shock	
			occurrence.	
Hiraoka, et al.	Study type: Retrospective	Inclusion criteria: BS with age 35 y	1° endpoint: Cardiac events (VF or SCD)	The presence of SCD or syncope is
2013	analysis of the Japan	of age or younger	, , ,	a risk factor for cardiac events in pts
<u>23702150</u>	Idiopathic Ventricular		Results: During a mean follow-up period of	with BS
(178)	Fibrillation registry	Exclusion criteria: N/A	43±27mo, cardiac events (VF and/or SCD) developed	
			in 8 cases, with 5 of 12 cases in the VF (41.7%), 2 of	
	<u>Size</u> : n=69 pts		17 cases in the Syncope (11.8%) and 1 of 40 cases in	
			the asymptomatic group (2.5%).	
			The VF group had a worse prognosis for cardiac	
			events than the Syncope and Asymptomatic group.	
			Multivariate analysis revealed symptoms as a risk	
			factor for predicting cardiac events.	
Sacher F, et al.	Study type: Prospective	Inclusion criteria:	1° endpoint: Cardiac events including syncope	VA occurred only in pts with syncope

2012	registry	Pts diagnosed with BS between 1999		suspected to be arrhythmic in origin at
<u>22504046</u>		and 2010	Results:	a rate of 5.5% per y. No sudden death
(179)	Size: n=203 pts		<ul> <li>Of 203 pts, 57 (28%) experienced syncope.</li> </ul>	occurred in pts with nonarrhythmic
		Exclusion criteria: N/A	23 pts with suspected arrhythmic syncope (Group 1),	syncope or with syncope of doubtful
			17 pts with non-arrhythmic syncope (Group 2) and 17	origin.
			with syncope of doubtful origin (Group 3).	
			• After mean follow-up of 65 ± 42 mo, 14 pts in Group	
			1 remained asymptomatic, 4 had recurrent syncope,	
			and 6 had appropriate ICD therapy. In Group 2, 9 pts	
			remained asymptomatic and 7 had recurrent	
			neurocardiogenic syncope. In Group 3, 7 remained	
			asymptomatic and 9 had recurrent syncope.	

## Data Supplement 20. Nonrandomized Trials, Observational Studies, and/or Registries of Short-QT Pattern and Syncope – (Section 4.3.2)

Study	Study Type/Design;	Patient Population	Primary Endpoint and Results	Summary/Conclusion
Acronym; Author;	Study Size		(P values; OR or RR; & 95% CI)	Comment(s)
Year Published				
Gollob, et al. 2011 21310316	Study type: Retrospective review of reported cases of SQTS.	Inclusion criteria: Reported cases of SQTS in English	1° endpoint: The creation of formal diagnostic criteria to facilitate the diagnostic evaluation of suspected cases of SQTS	Diagnostic criteria may lead to a greater recognition of this condition and provoke screening of at-risk family members.
(180)	Size: n=15 articles described unique	Exclusion criteria: N/A		,
	cases of SQTS		Results: A total of 61 cases were identified with a mean QTc value of 307 ms (range 248–381 ms). Short QT syndrome criteria were developed and consisted of 4 components including ECG, clinical Hx, family and genotype. An overall score of 4 points or greater indicates a high-probability diagnosis of SQTS, whereas 2 points or less makes a diagnosis of SQTS low probability. Pts with a score of 3 points are considered to have an intermediate probability of having SQTS.	
Gaita, et al. 2003	Study type: Retrospective	Inclusion criteria: Short QT interval with a Hx of syncope,	1° endpoint: Comprehensive EP evaluation	The short QT syndrome is characterized by familial sudden death, short refractory periods,
<u>12925462</u> (181)	<u>Size</u> : n=6 pts belonging to 2 families with idiopathic short QT interval	palpitations or resuscitated SD.	Results: At baseline ECG, all pts exhibited a QT interval ≤280 ms (QTc ≤300 ms). During EPS (n=4),	and inducible VF.

Brugada R, et	Study type: Prospective	Exclusion criteria: N/A Inclusion criteria: Short QT	short atrial and ventricular refractory periods were documented in all and increased ventricular vulnerability to fibrillation in 3 of 4 pts.  1° endpoint: Characterization of the genetic basis for	The authors demonstrated a novel genetic
al. 2004 14676148 (182)	Size: 3 families with hereditary short-QT syndrome and a high incidence of ventricular arrhythmias and SCD.	interval and history of ventricular arrhythmias or SCD  Exclusion criteria: N/A	SQTS  Results: In 2/3 families, the authors identified 2 different missense mutations resulting in the same amino acid change (N588K) in the S5-P loop region of the cardiac I <sub>Kr</sub> channel HERG (KCNH2). The mutations dramatically increase I <sub>Kr</sub> , leading to heterogeneous abbreviation of action potential duration and refractoriness, and reduce the affinity of the channels to I <sub>Kr</sub> blockers.	and biophysical mechanism responsible for SD in infants, children, and young adults caused by mutations in KCNH2.
Gallagher, et al. 2006 16996877 (183)	Study type: Retrospective  Size: n=12,012 pts	Inclusion criteria: Pts who underwent routine medical examination for occupational reasons  Exclusion criteria: N/A	1º endpoint: Survival  Results: The shortest QTc encountered was 335 ms.  Information about subsequent survival was available for 36 of the 60 pts with the lowest 1/2 centile of QTc values.  None of these pts died during the 7.9 ±4.5 y subsequent to the ECG that demonstrated the short QT interval.	<ul> <li>QTc ≤ 330 ms is extremely rare</li> <li>QT interval in the lowest 1/2 centile of the normal range does not imply a significant risk of SD.</li> </ul>
Anttonen O, et al. 2007 17679619 (184)	Study type: Retrospective Size: n=10,822 pts	Inclusion criteria: Randomly selected middle-aged pts enrolled in a population study and followed up for 29±10 y  Exclusion criteria: N/A	1º endpoint: All cause and CV mortality  Results: 10,822 randomly selected and followed for 29±10 y. The prevalence of SQTS (<340ms) was 0.4% and (<320ms) 0.1%. There were no SD or aborted CA or documented VA during follow up.	A short QT interval does not appear to indicate an increased risk for all-cause or CV mortality
Funada A, et al. 2008 18543308 (185)	Study type: Retrospective Size: n=10,984 pts	Inclusion criteria: Pts who had an ECG between February 2003 and May 2004  Exclusion criteria: Irregular rhythms, conduction disturbances and wide QRS	1º endpoint: Prevalence of SQTS (<300ms)  Results: In 10,984 pts, the prevalence of SQTS was 1.25% in males and 1.63% in females (2 SD below the mean). Only 3 pts had QTc<300ms. None were symptomatic.	SQTS is very rare
Kobza, et al. 2009	Study type: Retrospective	Inclusion criteria: Swiss male citizen 18–19 y of age.	1° endpoint: Prevalence of LQTS and SQTS	Short QT syndrome is a very rare entity in the population of young male adults

19303371	<b>Size</b> : n=41,767 ECGs		Results: The prevalence of SQTS (<320ms) was	
(186)		Exclusion criteria: Artifact,	0.02% and none of the pts had a QTc<300ms	
		pre-excitation and BBB.	·	
Giustetto C, et	Study type: Retrospective review	Inclusion criteria:	1° endpoint: Prevalence of arrhythmic events	Symptomatic pts are at high risk
al.	from the European Short QT registry	QTc≤360ms with cardiac		Hydroquinidine is effective in preventing
2011		arrest (n=18) or syncope	Results: The event rate was 3.3% per y and was	arrhythmic events
<u>21798421</u>	Size: n=53 pts	(n=8); Asymptomatic	limited to pts who were not receiving Hydroquinidine.	
(187)		QTc≤340ms and Family	<ul> <li>Of the 12 pts with a previous CA, 11 had an ICD</li> </ul>	
		members of affected pts	with 1 receiving appropriate shocks during follow-up.	
		(n=27)	<ul> <li>Of the 8 pts with syncope, 4 received an ICD and</li> </ul>	
			only 1 received appropriate shock for VF.	
		Exclusion criteria: N/A		

## Data Supplement 21. Nonrandomized Trials, Observational Studies, and/or Registries of Long-QT Syndrome – (Section 4.3.3)

Study Acronym; Author;	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Year Published				
Ouriel K, et al.	Study type:	Inclusion criteria: LQTS	1° endpoint: Cardiac events	LCS is associated with significant clinical benefits in
1995 <u>8574528</u>	Retrospective	refractory (n=9) or intolerant (n=1) to BB therapy	Results: No death. 9/10 developed Horner's	pts with long QT syndrome and the procedure should be considered when symptoms are refractory and
(188)	Size: n=10 pts	to BB therapy	syndrome. The frequency of symptoms	malignant, or when contraindications to β-blockers are
		Exclusion criteria: N/A	decreased from a mean of 7.1/y to	present.
			0.1/y(p<0.001). During a mean follow up of 1.3	
			y. All but 1 pts remained symptom- free. The	
			youngest pts died suddenly 10 mo after surgery.	
Priori SG, et al.	Study type:	Inclusion criteria: 193	1° endpoint: Cumulative probability of cardiac	The probability of having a cardiac event depends on
2003	Retrospective	consecutively genotyped families	event defined as syncope, cardiac arrest or SD	the genotype and sex.
<u>12736279</u>		with LQTS in Pavia, Italy		
(189)	Size: n=674 pts		Results:	
		Exclusion criteria: N/A	The incidence of first cardiac event was 30%	
			(LQT1), 46% (LQT2) and 42% (LQT3).	
			QTc was an independent predictor in LQT1 and	
			LQT2 whereas sex was independent predictor	
			in LQT3.	

	1			
Locati EH, et al.	Study type:	Inclusion criteria: LQTS pts and	1° endpoint: To evaluate age and sex-related	Data derived from a large registry (LQTS International
1998	Retrospective	affected family members	differences	Registry)
<u>9631873</u>				
(190)	Size: n=479 probands	Exclusion criteria: N/A	Results:	<ul> <li>In LQT1, male sex until puberty and female sex</li> </ul>
	and n=1041 affected		Among LQTS pts, the risk of cardiac events	during adulthood increase the risk of cardiac events.
	family members with		was higher in males until puberty and higher in	
	LQTS		females during adulthood. The same pattern	
			was evident among LQT1 gene carriers.	
			No age-sex difference in event rate was	
			detected in LQT2 and LQT3 carriers.	
Jons C et el.	Study type:	Inclusion criteria: LQTS pts with	1° endpoint: To identify risk factors for fatal	Cohort limited to pts with syncope
2010	Retrospective	QTc>450 ms presenting with	arrhythmias (aborted CA, appropriate ICD	ICD should not be the first line therapy in pts with a
20170817	Retrospective	syncope as a first symptom were	therapy and SCD)	
(191)	<b>Size:</b> n=1,059 pts	drawn from the International LQTS	therapy and 30D)	single episode of syncope as they have the lowest risk
(131)	<u>012e.</u> 11–1,000 pts	Registry	Results:	ICD is likely to save lives in pts with syncope despite
		rtegistry		BB therapy.
		Exclusion criteria: N/A	The lowest risk was in pts with 1 syncopal     The lowest risk was in pts with 1 syncopal	
		LACIUSION CITTERIA. IN/A	episode before the start of BB therapy.	
			Pts with syncope after BB or who were not	
			treated with BB therapy had a 3.6 fold increase	
			in risk.	
Zareba W, et al.	Study type: Outcome	Inclusion criteria:	1º endpoint: Death during follow up	ICD therapy saves lives
2003	data	ICD group (n=125): 54 CA, 19		
<u>12741701</u>		syncope despite BB and 52 for	Results: 1 death (1.3%) over 3 y in 73 ICD pts	
(192)	Size: n=286_pts with	other reasons	and 26 deaths (16%) in non-ICD pts over 8-y	
	LQTS; 125 with an ICD	<ul> <li>Non-ICD group (n=161): 89 CA</li> </ul>	follow up.	
	and 161 without an ICD	and 72 syncope despite BB		
		Exclusion criteria: N/A		
Schwartz, et al.	Study type:	Inclusion criteria: LQTS with an	1° endpoint: To determine the characteristics	Cohort limited to pts with an ICD
2010	Retrospective	ICD in the European LQTS	of LQTS pts receiving an ICD, indications and	• Age<20 y, QTc >500ms, prior CA and cardiac events
<u>20837891</u>		Registry	follow up	despite medical therapy were strong predictors of
(193)	<u>Size</u> : n=233 pts		·	appropriate ICD therapy.
			Results: 91% had symptoms including 44%	Absence of these risk factors indicates good
		Exclusion criteria: N/A	with prior CA. 41% had not been on prior drug	prognosis.
			therapy.	
			• During 4.6±3.2 y, at least 1 shock was	
			received by 28% of pts.	
			Predictors of appropriate ICD therapy	

Horner JM, et al. 2010	Study type: Retrospective	Inclusion criteria: Genetically confirmed LQTS including 51 pts	included age <20 y at implantation, QTc >500ms, prior CA and cardiac events despite therapy.  • No appropriate ICD therapy within 7 y in pts with none of these factors.  1° endpoint: Report outcome	• Syncope was a predictor of appropriate therapy (p=0.05)
<u>20816872</u> (194)	<u>Size:</u> n=459 pts	(14 LQT1, 22 LQT2, and 15 LQT3) who received an ICD from 2000 to 2010  Exclusion criteria: N/A	Results: During an average FU of 7.3 y, 12 (24%) of ICD recipients experienced an appropriate shock and none of the no-ICD group died. Predictors of appropriate therapy included secondary prevention indications, non-LQT3 genotype, QTc >500ms, syncope, TDP and negative family Hx.	• In 408 pts with no risk factors, no deaths were reported
Priori SG, et al. 2004	Study type: Retrospective	Inclusion criteria: Genotyped LQTS pts treated with BB	1° endpoint: Incidence of cardiac events	<ul> <li>Response to BB depend on the genotype</li> <li>LQT1 pts are better responders when compared to</li> </ul>
<u>15367556</u> (195)	<b>Size:</b> n=335 pts	Exclusion criteria: N/A	Results: Cardiac events occurred in 10%, 23% and 32% of pts with LQT1, LQT2 and LQT3. Predictors included non-LQT1 and QTc >500ms and first occurrence <7 y of age.	LQT2 and LQT3.  • QTc >500ms and first occurrence <7 y of age are predictors of future cardiac events
Vincent GM, et al. 2009 19118258 (196)	Study type: Retrospective  Size: n=216 pts	Inclusion criteria: Genotyped long-QT1 treated with BB and followed for a median of 10 y  Exclusion criteria: N/A	Pesults: Cardiac events on BB therapy  Results: Cardiac events occurred in 157 pts (73%) at a median age of 9 y, with CA in 26 (12%).  QT-prolonging drugs were used by 17 pts; 9 of 17 (53%) had CA compared with 17 of 199 nonusers (8.5%; OR: 12.0; 95% CI: 4.1–35.3; p<0.001).  The risk for CA/SD in compliant pts not taking QT-prolonging drugs was dramatically less compared with noncompliant pts on QT-prolonging drugs (OR: 0.03; 95% CI: 0.003–0.22; p=0.001). None of the 26 pts with CA before BB had CA/SD on BB.	<ul> <li>BB are extremely effective in long-QT syndrome type 1 and should be administered at diagnosis and ideally before the preteen years.</li> <li>BB noncompliance and use of QT-prolonging drug are responsible for almost all life-threatening "beta-blocker failures."</li> </ul>

Liu JF, et al. 2011 <u>21329841</u> (197)	Study type: International Long QT registry  Size: n=1,648 pts	Inclusion criteria: QTc ≥ 450 ms and/or documented LQTS-causing mutation and enrolled in registry before the 20 y age.  Exclusion criteria: N/A	1º endpoint: Recurrence of syncope after the first event  Results: Multivariate analysis demonstrated that QTc ≥ 500 ms was a significant predictor of a first syncope episode (HR: 2.16).  • Pts who experienced ≥ 1 episodes of syncope had a 6- to 12-fold (p<0.001 for all) increase in the risk of subsequent fatal/near-fatal events independently of QTc duration.  • BB therapy was associated with a significant reduction in the risk of recurrent syncope and subsequent fatal/near-fatal events.	Children and adolescents who present after an episode of syncope should be considered to be at a high risk of the development of subsequent syncope episodes and fatal/near-fatal events regardless of QTc duration.
Chockalingam P, et al. 2012 23083782 (198)	Study type: Retrospective  Size: n=382 (101 symptomatic) pts with LQT1/LQT2	Inclusion criteria: LQT1 and LQT2 pts on BB therapy (Proprnolol, Metoprolol and Nadolol)  Exclusion criteria: Less than 1 y of age at BB initiation	1º endpoint: To compare the efficacy of Propranolol, Metoprolol and Nadolol in pts with LQT1/LQT2      Results: QTc shortening was significantly greater with Propranolol.      None of the asymptomatic pts had cardiac events.      15% of the symptomatic had breakthrough with the greatest risk among those taking Metoprolol.	Not all BB are the same     Propranolol appears to be better than Metoprolol and Nadolol
Schwartz P, et al. 2004 <u>15051644</u> (199)	Study type: Retrospective  Size: n=147 pts	Inclusion criteria: LQTS pts who underwent LCSD (99% symptomatic with 75% of those treated with BB remaining symptomatic  Exclusion criteria: N/A	1° endpoint: Long-term efficacy of LCSD  Results: Post-LCSD, 46% remained symptomatic. The mean yearly number of cardiac events per patient dropped by 91% (P<0.001). Among 74 pts with only syncope before LCSD, all types of cardiac events decreased significantly as in the entire group, and a post-LCSD QTc <500 ms predicted very low risk.	<ul> <li>LCSD is associated with a significant reduction in the incidence of aborted cardiac arrest and syncope in high-risk LQTS pts when compared with pre-LCSD events.</li> <li>However, LCSD is not entirely effective in preventing cardiac events including sudden cardiac death during long-term follow-up.</li> <li>LCSD should be considered in pts with recurrent syncope despite β-blockade and in pts who experience arrhythmia storms with an implanted defibrillator.</li> </ul>
Collura, et al. 2009 <u>19467503</u>	Study type: Retrospective	Inclusion criteria: Secondary prevention in 11 pts including 8 with LQTS and primary prevention	1° endpoint: Outcome with LCSD using video- assisted thoracic surgery	Videoscopic denervation surgery, in addition to traditional LCSD, offers a safe and effective treatment option for the personalized medicine required for pts

(200)	Size: n=20 pts including 12 with LQTS, 2 JLNS, 4	in 9 pts.	Results: There were no perioperative complications. The average length of available	with LQTS/CPVT.
	genotype negative LQTS	Exclusion criteria: N/A	follow-up was 16.6 ± 9.5 mo (range 4–40 mo).	
	and 2 CPVT		Among the 18 pts who underwent VATS-LCSD,	
			the average time from operation to dismissal	
			was 2.6 d (range 1–15 d), the majority being	
			next-day dismissals. Among those receiving	
			LCSD as secondary prevention, there has been	
			a marked reduction in cardiac events.	
Abu-Zeitone A, et al.	Study type:	Inclusion criteria: Pts with LQTS	1° endpoint: Compare efficacy of different BB	<ul> <li>BB efficacy differed by genotype. Nadolol was the</li> </ul>
2014	Retrospective	who were prescribed common BB		only BB associated with a significant risk reduction in
<u>25257637</u>		(atenolol, metoprolol, propranolol,	Results: In LQT1, the risk reduction for first	pts with LQT2.
(201)	Size: n=1,530 pts	or nadolol).	cardiac events was similar among the 4 BB	<ul> <li>Pts experiencing cardiac events during BB therapy</li> </ul>
			(atenolol, metoprolol, propranolol and nadolol),	are at high risk for subsequent cardiac events, and
		Exclusion criteria: Prescribed BB	but in LQT2, nadolol provided the only	propranolol is the least effective drug in this high-risk
		after the age of 40 or have an ICD	significant risk reduction (HR: 0.40 (95%CI: 0.16 to 0.98).	group.
			<ul> <li>Among pts who had a prior cardiac event</li> </ul>	
			while taking BB, efficacy for recurrent events	
			differed by drug (p=0.004), and propranolol was	
			the least effective compared with the other BB.	

#### Data Supplement 22. Nonrandomized Trials, Observational Studies, and/or Registries of CPVT-Medical Therapy – (Section 4.3.4)

Study Acronym; Author;	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR;	Summary/Conclusion Comment(s)
Year Published			& 95% CI)	
Padfield, GJ, et al.	Study type:	Inclusion criteria: CPVT secondary to	1° endpoint: Safety of flecainide as mono-	Flecainide mono-therapy is an option
2016	Retrospective	mutations in the RyR2 gene who refused	therapy in pts with CPVT	in pts with CPVT who are intolerant to
<u>26416620</u>		(n=1) or were intolerant to BB therapy (n=7)		BB therapy.
(202)	Size: n=8 pts		Results: Flecainide mono-therapy was better	
, ,		Exclusion criteria: N/A	than, or at least as effective as, BB mono-	
			therapy in reducing exercise-induced arrhythmia.	
			•No episodes of arrhythmic pre-syncope,	
			syncope, or CA occurred in pts on flecainide	
			mono-therapy during the follow-up period of 37.1	
			mo (range1.4–75.5 mo).	

Loophordt at al	Cturdy tymes	Inclusion evitoria: Cuncono duo to	40 4	First second of advances in demandant
Leenhardt, et al. 1995	Study type: Observational	Inclusion criteria: Syncope due to documented or suspected VA.	1° endpoint: Syncope recurrence and exercise induced VA	First report of adrenergic-dependent
7867192	Observational	documented of suspected VA.	Induced VA	ventricular tachy-arrhythmia in pts with normal QT interval and no structural
(203)	Size: n=21 pts	Exclusion criteria: N/A	Results: On BB therapy, the pts' symptoms and	heart disease.
(203)	<u>5126</u> . 11–2 1 pts	Exclusion chiena. WA	polymorphic tachyarrhythmias disappeared.	
			During a mean follow-up period of 7 y, 3	BB help suppress exercise induced arrhythmias
			syncopal events and 2 sudden deaths occurred,	annyunnas
			probably due to treatment interruption.	
Priori, et al.	Study type:	Inclusion criteria: Exercise or emotion	1° endpoint: Clinical and genetic	CPVT is a clinically and genetically
2002	Retrospective	induced bidirectional VT (n=14), PMVT (n=12)	characterization	heterogeneous disease manifesting
12093772	1 tota oop ootavo	and catecholaminergic idiopathic VF (n=4)	Characterization	beyond pediatric age with a spectrum
(204)	Size: n=30 probands and	and sateshelanimorgic larepatine vi (ii i)	Results: Genotype-phenotype analysis showed	of polymorphic arrhythmias.
(,	118 family members	Exclusion criteria: N/A	that pts with RyR2 CPVT have events at a	BB reduce arrhythmias, but in 30%
	, , , , , , , , , , , , , , , , , , , ,		younger age than do pts with non-genotyped	of pts an implantable defibrillator may
			CPVT and that male sex is a risk factor for	be required.
			syncope in RyR2-CPVT (RR:4.2).	33.34
			All 39 clinically affected pts were treated with	
			BB; however, antiadrenergic drugs provided only	
			incomplete protection from recurrence of	
			sustained VT and VF.	
			<ul> <li>18 of 39 pts treated with β-blockers had</li> </ul>	
			cardiac arrhythmias. An ICD was recommended	
			and implanted in12/18. Over a follow-up of ≈2 y,	
			50% of pts with the ICD received an appropriate	
			shock to terminate ventricular tachyarrhythmias.	
Sumitomo, et al.	Study type:	Inclusion criteria: 1) Exercise or	1° endpoint: Questionnaire responses and ECG	<ul> <li>Pts with CPVT have a poor</li> </ul>
2003	Questionnaires were	catecholamine induced VA (>3 beats) with at	characteristics	prognosis. BB do not always control
<u>12482795</u>	sent to major Japanese	least 2 morphologies 2) absence of known		symptoms thus the need for other
(205)	pediatric centers	secondary causes including electrolyte	Results: The initial CPVT manifestations were	pharmacological and non-
	Size n=20 contars	abnormalities and structural heart disease	syncope (79%), cardiac arrest (7%), and a family	pharmacological therapies.
	Size: n=29 centers	and 3) no evidence of long QT or Brugada.	Hx (14%).	
		Exclusion criteria: N/A	There was 100% inducibility of CPVT by  Oversign 75% by extraphologing influeing and  Oversign 75% by extraphologing influeing and  Oversign 75% by extraphologing influeing and  Oversign 75% by extraphologing influeing and	
		LAGIUSION GIRENA. N/A	exercise, 75% by catecholamine infusion, and none by programmed stimulation.	
			<ul> <li>During a follow up of 6.8 (4.9) y, sudden death</li> </ul>	
			occurred in 24% of the pts. BB completely	
			controlled CPVT in only 31% of cases. Calcium	
			antagonists partially suppressed CPVT in	
			antagonists partially suppressed of virill	

			autosomal dominant cases.	
Hayashi, et al.	Study type: Multicenter	Inclusion criteria: Exercise induced	1° endpoint: Incidence of cardiac events	BB reduce the cardiac event rate in
2009	observational study	polymorphic ventricular arrhythmias or		both CPVT pts and affected families;
19398665	Observational study	identification of a mutation in the RYR2 or	(exertional or stress induced syncope, aborted CA, appropriate ICD shocks or SCD)	however, they are not completely
(206)	Size: n=101 pts	CASQ2 gene	CA, appropriate ICD shocks of SCD)	protective.
(200)	<u>5126</u> . 11–101 pts	OADQ2 gene	Results: During a mean follow-up of 7.9 y,	protective.
		Exclusion criteria: >55 y of age	cardiac events occurred in 27 pts (27%),	
		<u>Exclusion ornaria</u> . * 66 y or ago	including 2 mutation carriers with normal	
			exercise tests.	
			The estimated 8 y event rate was 32% in the	
			total population and 27% and 58% in the pts with	
			and without BB, respectively. Absence of BB HR:	
			5.48; 95% CI: 1.80–16.68) and younger age at	
			diagnosis (HR: 0.54 per decade; 95% CI: 0.33-	
			0.89) were independent predictors.	
			The estimated 8 y event rate for fatal or near	
			fatal events (ACA, SCD) was 13%. Absence of	
			BB (HR: 5.54; 95% CI: 1.17–26.15) and Hx of	
			aborted CA (HR: 13.01; 95% CI: 2.48–68.21)	
			were independent predictors.	
van der Werf, et al.	Study type: Meta-	Inclusion criteria: CPVT pts	1º endpoint: Arrhythmic, non-fatal and fatal	The variability in outcome with BB
2012	analysis including 11		events	therapy is due to multiple factors
<u>21893508</u>	studies using BB and	Exclusion criteria: N/A		including the dose, compliance and
(207)	review of other therapies		Results: Median FU was 20 mo 8 y. 88% of pts	concomitant use of other drugs
	Size: n=403 pts		were given BB.	including flecainide and Verapamil.
	<u>312e</u> . 11-403 pts		•The estimated overall 4- and 8 y arrhythmic	
			event rates were 18.6% (95% CI: 8.3–28.9) and 37.2% (95% CI: 16.6–57.7), respectively.	
			•Estimated 4- and 8 y near-fatal arrhythmic	
			event rates were 7.7% (95% CI: 3.7–11.7) and	
			15.3% (95% Cl: 7.4–23.3), respectively.	
			•Fatal events occurred in 3.2% (95% CI: 1.6–	
			4.8) at 4 y and 6.4% (95% CI: 3.2–9.6) at 8 y	
			follow-up	
van der Werf, et al.	Study type: Chart	Inclusion criteria:	1° endpoint: Reduction of VA during exercise	Flecainide reduced exercise-induced
2011	review from 8 tertiary	1) Exercise induced PMVT or bidirectional VT	testing	VA in pts with CPVT not controlled by
<u>21616285</u>	referral centers	2) Mutation in the gene encoding RyR2 or		conventional drug therapy.
(208)		cardiac Calsequestrin	Results: Exercise tests comparing flecainide in	,

		T	T	
	Size: n=33 pts	Fredrice witoric N/A	addition to conventional therapy with	
		Exclusion criteria: N/A	conventional therapy alone were available for 29	
			<ul><li>pts.</li><li>The median daily flecainide dose in responders</li></ul>	
			was 150 mg (range 100 to 300 mg).	
			• 22 pts (76%) had either partial (n=8) or	
			complete (n=14) suppression of exercise-induced	
			VA with flecainide (p<0.001).	
			No pts experienced worsening of exercise-	
			induced VA.	
Swan, et al.	Study type: Prospective	Inclusion criteria: Pts with clinical diagnosis	1° endpoint: Effect of verapamil and	First study to demonstrate in vivo
2005	physiology study in	of CPVT and carrying a RyR2 mutation	magnesium on exercise induced VA.	that verapamil can suppress
<u>15720454</u>	human pts			premature ventricular complexes and
(209)		Exclusion criteria: N/A	Results: Premature ventricular complexes	non-sustained ventricular salvoes in
	Size: n=6 pts		appeared later and at higher heart rate during	CPVT caused by RyR2 mutations.
			verapamil compared to baseline (119 ± 21 vs.	Physiology study with no long-term
			127 ± 27 min−1, p<0.05). Magnesium did not inhibit the arrhythmias.	follow up
Doggo et al	Childry france	Inclusion evitories CDVT ato with a Uv of	,	The combination of coloring about
Rosso, et al. 2007	Study type: Retrospective 2 center	Inclusion criteria: CPVT pts with a Hx of syncope or CA and exercise induced	1° endpoint: Exercise induced arrhythmias and clinical outcome	The combination of calcium channel  blackers with BB might be better than
17765612	study	ventricular ectopy despite maximally tolerated	Clinical outcome	blockers with BB might be better than BB alone.
(210)	Study	BB therapy	Results: 1) 3 pts had non-sustained VT on	Short-term study
(2.0)	Size: n=5 pts	25 thorapy	blockers, and none of them had VT on	Short-term study
	<u> </u>		combination therapy. 2) The number of	
		Exclusion criteria: N/A	ventricular ectopic beats during the whole	
			exercise test went down from 78 ± 59 beats to 6	
			± 8 beats. 3) 1 pts with recurrent spontaneous	
			VT leading to multiple shocks from her ICD	
			despite maximal blocker therapy remained free	
			of arrhythmias for 7 mo since the addition of	
0.0.1.1	20 1 1	Living the state of the state o	verapamil therapy.	D " DD"
Sy R, et al. 2011	Study type:	Inclusion criteria: Hx of sudden cardiac	1º endpoint: Long-term outcome and relation	Despite BB therapy and selective
21315846	Retrospective	arrest or symptoms occurring in the context of physical activity or acute emotion in	between age and clinical presentation	ICD implantation, breakthrough
(211)	Size: n=27 pts	conjunction with exercise or adrenaline-	Results: Presentation was CA in 33% and	arrhythmias occur and may be associated with adverse outcomes
(211)	<u>0126</u> . 11-21 pto	induced polymorphic or bidirectional VT of ≥ 4	syncope in 56%, and 11% were asymptomatic.	associated with adverse outcomes
		beats.	Polymorphic or bidirectional VT was provoked	
		First-degree relatives of affected individuals	with exercise in 63% and adrenaline in 82%.	
L			1	

		were diagnosed with CPVT if polymorphic or bidirectional  VT was observed during exercise or adrenaline challenge, on Holter monitoring, or if genetic testing was positive for the disease-causing mutation in the family.  Exclusion criteria: N/A	• During follow-up of 6.2±5.7 y, 2 pts died despite having an ICD, 4 pts received ICD therapy for VT, and 5 pts had inappropriate therapy for SVT. Pts presenting with late-onset CPVT (>21 y of age; n=10) were often female (80%) and less likely to have <i>RyR2</i> (Ryanodine receptor type 2) mutations (33%), and fatal events were not observed during follow-up (4.1±3.6 y).	
Roston TM, et al. 2015 25713214 (212)	Study type: Retrospective cohort study  Size: n=226 pts	Inclusion criteria: 170 probands and 56 relatives  Exclusion criteria: N/A	1º endpoint: Treatment outcome  Results: Symptomatic presentation was reported in 176 (78%). Syncope (p<0.001), cardiac arrest (p<0.001), and treatment failure (p=0.008) occurred more often in probands.  ■ BB were prescribed in 205 of 211 pts (97%) on medication, and 25% experienced at least 1 treatment failure event. ICDs were placed in 121 (54%) and were associated with electrical storm in 22 (18%). Flecainide was used in 24% and LCSD in 8%. 6 deaths (3%) occurred during a cumulative follow-up of 788 pts-y.	BB were almost universally initiated; however, treatment failure, noncompliance and sub-therapeutic dosing were often reported.     Treatment failure was rare in the quarter of pts on flecainide.     LCDS was not uncommon although the indication was variable.     ICDs were common despite numerous device-related complications.

#### Data Supplement 23. Nonrandomized Trials, Observational Studies, and/or Registries of CPVT- LSCD and ICD Therapy – (Section 4.3.4)

Study Acronym;	Study Type/Design;	Patient Population	Primary Endpoint and Results	Summary/Conclusion
Author;	Study Size		(P values; OR or RR;	Comment(s)
Year Published			& 95% CI)	
Moray A, et al.	Study type: Retrospective	Inclusion criteria: 10 y of age boy with	1° endpoint: Safety of simultaneous ICD	Simultaneous ICD insertion and
2011	Case report	CPVT	insertion and thoracoscopic sympathectomy	thoracoscopic sympathectomy is feasible and
<u>21478052</u>				safe in pts with CPVT
(213)	Size: n=1 patient	Exclusion criteria: N/A	Results: The procedure was safe suggesting	·
			that it is a better approach than sequential	
			procedures	

Celiker A, et al. 2009 <u>19102802</u> (214)	Study type: Retrospective Size: n=16 children pts	Inclusion criteria: Diagnosis of CPVT  Exclusion criteria: N/A	1° endpoint: Clinical features, treatment and outcome  Results: The mean age of pts at the onset of symptoms and at the time of diagnosis was 7.8 ± 2.5 y, and 10.6 ± 3.5 y, respectively. Syncope was the main complaint in 11.  •Treatment included propranolol plus verapamil if VT was still inducible. ICD was implanted in 4 pts.  Of the 16 pts, 4 died suddenly, giving a rate of mortality of 25%.	CPVT must be considered in the differential diagnosis of syncope in children without heart disease but with a normal QT interval. Medical treatment with propranolol and verapamil may decrease the incidence of arrhythmia. Implantation of an ICD should be considered in those resistant to drug therapy.
Wilde AA, et al. 2008  18463378 (215)  De Ferrari, et al. 2015  26019152 (216)	Study type: Retrospective single center experience  Size: n=3 pts  Study type: Retrospective including pts from 11 centers worldwide  Size: n=63 pts	Inclusion criteria: CPVT with symptoms despite BB therapy 3/3) and mexiletine (1/3)  Exclusion criteria: N/A  Inclusion criteria: Asymptomatic and symptomatic pts  Exclusion criteria: N/A	1° endpoint: Cardiac events  Results: LCSD resulted in marked reduction in cardiac arrhythmias and improvement in QOL.  1° endpoint: Cardiac events  Results: LCSD was performed in 9 asymptomatic and 54 symptomatic pts including 38 pts (25 syncope) with breakthrough events despite optimal medical therapy.  • The 1 and 2 y cumulative event-free survival rates were 87% and 81%. The percentage of pts with major cardiac events despite optimal medical therapy (n=38) was reduced from 100% to 32% (p <0.001) after LCSD.	First study to provide evidence that left cardiac sympathetic denervation may be an effective alternative treatment, especially for pts whose symptoms are not adequately controlled by means of BB therapy.      LCSD is an effective antifibrillatory intervention for pts with CPVT. Whenever syncope occurs despite optimal medical therapy, LCSD could be considered the next step rather than an ICD and could complement ICDs in pts with recurrent shocks.
Waddell-Smith, et al. 2015 26224781 (217)	Study type: Retrospective Survey- based  Size: n=47 pts who underwent LCSD including 40 with LQTS and 7 with CPVT	Inclusion criteria: Underwent video- assisted thoracoscopic LCSD and completion of a telephone survey  Exclusion criteria: N/A	1º endpoint: Physical and psychological effects of LCSD and pts satisfaction.  Results: Side effects were reported by 42 of 44 (95%). 29 (66%) reported left sided dryness, 26 (59%) a Harlequin-type (unilateral) facial flush, 24 (55%) contralateral hyperhidrosis, 17 (39%) differential hand temperatures, 5 (11%) permanent ptosis (4	Despite significant morbidity resulting from LCSD, pts with LQTS and CVPT have high levels of post-operative satisfaction.

			(9%) transient ptosis). 5 (11%) have thermoregulation difficulties, 4 (9%) a sensation of left arm paraesthesia and 3 (7%) lost their sympathetic flight/fright response.  • 38 pts (86%) were happy with procedure, 33 (75%) felt safer and 40 (91%) recommend the procedure. 40 (91%) pts were happy with their scar.	
Marai, et al. 2012 <u>22481011</u> (218)	Study type: Retrospective  Size: n=27 pts	Inclusion criteria: CPVT  Exclusion criteria: N/A	1° endpoint: Death  Results: 27 pts were followed for 1-15 y (median 9). 20 were symptomatic at baseline and 13 remained symptomatic after treatment with high dose BB.  • 8 pts refused ICD with 6 eventually dying. 5 received an ICD with 4/5 experiencing a VT storm not responsive to ICD shocks but with spontaneous termination. No death occurred	ICD should be recommended in pts refractory to BB therapy.     These pts may have recurrent ventricular tachycardia storms treated but not terminated by recurrent ICD shocks, without degeneration to ventricular fibrillation.
Roses-Noguer, et al. 2014 24120999 (219)	Study type: Retrospective Size: n=13 pts	Inclusion criteria: CPVT with an ICD implant for cardiac arrest (7 pts) and syncope (6 pts)  Exclusion criteria: N/A	in the ICD group.  1º endpoint: Effectiveness of ICD shocks  Results: Among appropriate shocks, 20 (32%) were effective in terminating sustained arrhythmia and 43 (68%) were ineffective.  • Shocks delivered to triggered arrhythmias nearly always failed (1 of 40; 3% effective), while shocks delivered to VF were usually successful (19 of 23; 83% effective; p<0.001). No pts died.	The effectiveness of ICD shock therapy in CPVT depends on the mechanism of the rhythm treated. Shocks delivered to initiating triggered arrhythmias nearly always fail, whereas those for subsequent VF are usually effective.

## Data Supplement 24. Nonrandomized Trials, Observational Studies, and/or Registries of Early Repolarization Pattern – (Section 4.3.5)

	Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
M	lahida S, et al.	Study type:	Inclusion criteria: ER syndrome	1° endpoint: Inducibility of VA	<ul> <li>Programmed stimulation</li> </ul>
20	015	Retrospective	with a history of aborted sudden		protocols do not enhance risk

<u>25593056</u>	multicenter study	death due to ventricular fibrillation	Results: VF was inducible in only 18 of 81 (22%) pts. During	stratification in pts with ER
(220)	<u>Size</u> : n=81 pts	Exclusion criteria: Structural heart disease and >60 y of age.	follow-up of 7.0±4.9 y, 6 of 18 (33%) pts with inducible VF during EPS experienced VF recurrences, whereas 21 of 63 (33%) pts who were non-inducible experienced recurrent VF (p=0.93).	syndrome.
Morady F, et al. 1986 <u>3717024</u> (221)	Study type: Retrospective Size: n=109 pts	Inclusion criteria: 52 pts with a Hx of documented, sustained monomorphic VT and inducible VT and 57 pts with non-clinical inducible polymorphic VT or VF.  Exclusion criteria: N/A	1° endpoint: Characteristics of coupling intervals that induce clinical and non-clinical VT and VF  Results: The mean coupling intervals of the first, second and third extra stimuli that induced nonclinical VT/VF were significantly shorter than the corresponding coupling intervals that induced the clinical VTs.  Regardless of the basic drive cycle length, the shortest coupling interval required to induce a clinical VT was 180 ms. Depending on the drive cycle length, 29 to 70% of nonclinical VT/VF induced by 3 extrastimuli required a coupling interval of less than 180 ms to induce.	The results of this study demonstrate that the coupling intervals required to induce non-clinical forms of VT or VF are often shorter than the coupling intervals required to induce clinical VT raising concerns about the specificity of EP studies when aggressive stimulation protocols are used.
Nunn LM, et al. 2011 <u>21737021</u> (222)	Study type: Retrospective Size: n=363 pts	Inclusion criteria: Families of sudden arrhythmic death syndrome probands  Exclusion criteria: N/A	<u>1° endpoint</u> : The prevalence of J-point elevation among the relatives of sudden arrhythmic death syndrome probands <u>Results:</u> A total of 363 first-degree relatives from 144 families were evaluated. J-point elevation in the inferolateral leads was present in 23% of relatives and 11% of control pts (OR: 2.54, 95% CI: 1.66–3.90; p<0.001).	J-point elevation is more prevalent in the relatives of sudden arrhythmic death syndrome probands than in controls. This indicates that ER is an important potentially inheritable pro-arrhythmic trait or marker of pro-arrhythmia in sudden arrhythmic death syndrome.
Haissaguerre M, et al. 2008 18463377 (223)	Study type: Retrospective  Size: n=206 case pts and 412 control pts matched for age, sex, race and level of physical activity	Inclusion criteria: Resuscitated from cardiac arrest due to idiopathic VF  Exclusion criteria: Age >60 y of age	1° endpoint: Prevalence of ER  Results: ER was more frequent in case pts with idiopathic ventricular fibrillation than in control pts (31% vs. 5%, p<0.001).  During a mean follow-up of 61 ± 50 mo, defibrillator monitoring showed a higher incidence of recurrent ventricular fibrillation in case pts with a repolarization abnormality than in those without such an abnormality (HR: 2.1; 95% CI: 1.2–3.5; p=0.008).	Among pts with a Hx of idiopathic VF, there is an increased prevalence of early repolarization.
Rosso, et al. 2008 <u>18926326.</u> (224)	Study type: Case control study  Size: n=45 pts with	Inclusion criteria: Idiopathic VF compared with age and sex matched control pts	1° endpoint: Prevalence of J point and ST elevation  Results: J-point elevation was more common among pts with idiopathic VF than among matched control pts (42% vs. 13%,	J-point elevation is found more frequently among pts with idiopathic VF than among healthy control pts. The frequency of J-

	idiopathic VF and 121	Evaluaian aritaria. The processes of	n=0.001). This was true for I point also ation in the inferior leads	noint alevation among vours
	young athletes	Exclusion criteria: The presence of an etiology for the cardiac arrest	p=0.001). This was true for J-point elevation in the inferior leads (27% vs. 8%, p=0.006) and for J-point elevation in leads I to aVL	point elevation among young athletes is higher than among
	young atmetes	an eliblogy for the cardiac arrest	(13% vs. 1%; p=0.009). J-point elevation in V(4) to V(6) occurred	healthy adults but lower than
			with equal frequency among pts and matched control pts (6.7%	among pts with idiopathic VF.
			vs. 7.3%; p=0.86).	among plo maridiopaulio vi .
			The presence of ST-segment elevation or QRS slurring did not	
			add diagnostic value to the presence of J-point elevation.	
Merchant FM, et al.	Study type:	Inclusion criteria: Idiopathic VF and	1° endpoint: Prevalence of ER and QRS notching	Left precordial terminal QRS
2009	Retrospective	ICD implant		notching is more prevalent in
<u>19892058</u>			Results: ER was present in 9/39 (23%) pts. QRS notching was	malignant variants of ER than in
(225)	Size: n=39 cases of	Exclusion criteria: Structural heart	significantly more prevalent among cases when present in leads	benign cases.
	idiopathic VF	disease, CAD or the presence of an	V4 (44% vs. 5%, p=0.001) and V5 (44% vs. 8%, p=0.006), with a	
		arrhythmia susceptibility syndrome	similar trend in lead V6 (33% vs. 5%, p=0.013).	
T'11 ( )	0, 1,	(LQTS, SQTS, WPW, BS or ARVD)		
Tikkanen, et al. 2009	Study type:	Inclusion criteria: Community	<u>1° endpoint</u> : Prevalence and prognostic significance of ER	An ER pattern in the inferior
19917913	Retrospective	based general population	including death from cardiac cause, death from arrhythmia and	leads of a standard ECG is
(226)	Size: n=10,864 middle-	Exclusion criteria: N/A	death from any causes	associated with an increased risk of death from cardiac causes in
(220)	aged pts	Exclusion chiena. WA	Results: ER was present in 630 pts (5.8%): 384 (3.5%) in	middle-aged pts.
	agou pio		inferior leads and 262 (2.4%) in lateral leads, with elevations in	middic-aged pts.
			both leads in 16 pts (0.1%).	
			J-point elevation of at least 0.1 mV in inferior leads was	
			associated with an increased risk of death from cardiac causes	
			(adjusted RR: 1.28; 95% CI: 1.04–1.59; p=0.03).	
			<ul> <li>J-point elevation of more than 0.2 mV in inferior leads (n=26;</li> </ul>	
			0.3%) had a markedly elevated risk of death from cardiac causes	
			(adjusted RR: 2.98; 95% CI: 1.85–4.92; p<0.001) and from	
D / 1 / 1	<b>2</b> 1 1 0		arrhythmia (adjusted RR: 2.92; 95% CI: 1.45–5.89; p=0.01).	
Patel, et al. 2010	Study type: Case	Inclusion criteria: CAD + ICD	1° endpoint: Prevalence of ER	ER and, in particular, notching  in the piritarian leads in accordant.
20657030	Control design	implant + sustained arrhythmic events	Descritor Overall contraction in 2 or more leaders	in the inferior leads is associated
(227)	Size: n=60 pts (CAD +	GVGIIIS	Results: Overall, early repolarization in 2 or more leads was more common in cases than control pts (32% vs. 8%, P=0.005).	with increased risk of life- threatening VA in pts with CAD,
(221)	ICD + sustained	Exclusion criteria: Pts who had an	Early repolarization was noted more commonly in inferior leads	even after adjustment for LVEF.
	arrhythmic events) and	acute MI during follow up, suspected	(23% vs. 8%, p=0.03), and a trend was noted in leads V4	over and adjustment for EVEL.
	n=60 control pts (CAD +	BS and pts with QRS ≥120 ms	through V6 (12% vs. 3%, p= 0.11).	
	ICD + no arrhythmic	•	<del> </del>	
	events)			
Tikkanen, et al.	Study type:	Inclusion criteria: Pts participating	1° endpoint: Mortality over a 30±11 y follow up period	<ul> <li>ST-segment morphology</li> </ul>

2011 21632493 (228)	Retrospective  Size: n=10,957 pts	in the Finnish Social Insurance Institution's Coronary Heart Disease Study who had undergone clinical baseline examinations between 1966 and 1972.  Exclusion criteria: Pts with missing data	Results: Pts with ER≥ 0.1 mV and horizontal/descending ST variant (n=412) had an increased HR of arrhythmic death (RR: 1.43; 95% CI: 1.05–1.94).  •When modeled for higher amplitude ER (>0.2 mV) in inferior leads and horizontal/descending ST-segment variant, the HR of arrhythmic death increased to HR: 3.14 (95% CI: 1.56–6.30).  • However, in pts with ascending ST variant, the relative RR for arrhythmic death was not increased (RR: 0.89; 95% CI: 0.52–1.55).	variants associated with ER separates pts with and without an increased risk of arrhythmic death in middle-aged pts.  • Rapidly ascending ST segments after the J-point, the dominant ST pattern in healthy athletes, seems to be a benign variant of ER
Sinner, et al. 2010 <u>20668657</u> (229)	Study type: Population based study applying a case-cohort design  Size: n=1,945 pts representing a source population of 6,213 individuals, were analyzed	Inclusion criteria: 25-74 y of age Exclusion criteria: N/A	1° endpoint: Prevalence of ERP and its association with cardiac and all-cause mortality  Results: Prevalence of ERP was 13.1%. ERP was associated with cardiac and all-cause mortality, most pronounced in those of younger age and male sex; a clear ERP-age interaction was detected (p=0.005).  Age-stratified analyses showed HRs for cardiac mortality of 1.96 (95% CI: 1.05–3.68, p=0.035) for both sexes and 2.65 (95% CI: 1.21–5.83, p=0.015) for men between 35–54 y of age. An inferior localization of ERP further increased ERP-attributable cardiac mortality to HRs of 3.15 (95% CI: 1.58–6.28, p=0.001) for both sexes and to 4.27 (95% CI: 1.90–9.61, p<0.001) for men between 35-54 y of age.	ERP was associated with about a 2- to 4-fold increased risk of cardiac mortality in individuals between 35 and 54 y. An inferior localization of ERP was associated with a particularly increased risk.

## Data Supplement 25. RCTs Comparing Vasovagal Syncope – (Section 5.1.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Lu CC, et al.	Aim: Assess whether	Inclusion: Healthy male	Intervention: Ingestion of 10%	1° endpoint: Orthostatic tolerance	Glucose water attenuates
2008	glucose water ingestion will	-	glucose water before 70 degree	(time to presyncope during 70 degree	reflex role of PVR during
<u>18772858</u>	reduce orthostatic tolerance	Exclusion: Hx of syncope,	HUTT	HUTT): 13 of 15 (87%) ingesting pure	orthostatic stress, perhaps by
(230)	in young healthy volunteers	any medications		water were able to complete the full tilt	vasodilatation in splanchnic
			Comparator: Ingestion of pure	without presyncope, but 7 of 15 (47%)	circulation or raising plasma
	Study type: Analytical,		water 5 mins before 70 degree	ingesting glucose water could	osmolality which may enhance

	Randomized controlled crossover, prospective cohort,  Size: n=15 pts		HUTT	complete the full tilt. Test was terminated sooner in glucose water group (40.0 +/- 6.9 min) vs. pure water group (43 +/- 5.6 min), p=0.008. There was no difference in symptom scores (p=0.26) between 2 groups.  Safety endpoint: N/A	barorefex control of SNS.
Schroeder, et al. 2002 12451007 (231)	Aim: To assess water drinking on orthostatic tolerance in healthy pts  Study type: Analytical, randomized controlled, prospective crossover  Size: n=13 pts	Inclusion: Healthy volunteers  Exclusion: Regular medication except oral contraceptives	Intervention: 500 mL nonsparkling mineral water at room temperature  Comparator: 50 mL nonsparkling mineral water • then 60 degree HUTT for 20 min followed by LBNP for 10 min each at -20, then -40, -60 mmHg	1° endpoint: Drinking 500 mL water prolonged time to presyncope in 11 pts from 31 +/-3 min to 36+/-3 min (P<0.001). Supine heart rate, BP, SV, and cardiac output were not significantly different with 500 mLwater drinking. With HUTT, 500 mL water drinking blunted decrease in SV from -45+/-2% to -38+/-3%, p<0.01  Safety endpoint: N/A	Water drinking 500 mL increases orthostatic tolerance, with the effect apparently mediated with factors beyond increasing plasma volume. Increase in peripheral resistance and vasoconstrictor tone may have role.
El-Sayed, et al. 1996 8673750 (232)	Aim: To evaluated salt supplementation in syncope with orthostatic intolerance, Study type: Analytical, Randomized placebo controlled, prospective cohort, Size: n=20 pts; Study type: Analytical, observational, open label, prospective cohort Size: n=11 pts	Inclusion: Recurrent syncope without etiology  Exclusion: None	RDBPCT: Intervention: Sodium chloride 10 mmol Comparator: Placebo 12x daily then 60 degree HUTT with LBNP up to -40 mmHg Open label: Intervention: Slow sodium 10 mmol 12x daily (pts told it was a "mineral dietary supplement") then 60 degree HUTT with LBNP up to -40 mmHg	1° endpoint: RDBPCT: 8 of 10 pts taking salt, vs. 3 of 10 taking placebo showed significant increases in plasma and blood volumes (p<0.05); all pts with increased plasma and blood volumes showed improved tolerance to orthostatic stress (time to presyncope)  Safety endpoint: N/A  Open label: 7 of 11 taking salt had increased plasma and blood volumes, and these pts showed improved symptoms of orthostatic tolerance	Pts with salt supplementation (increasing plasma volume by >90 mL) had significant increase in orthostatic tolerance. Pts with signs of high salt intake at baseline (by 24 H urinary sodium excretion) did not benefit from additional salt loading
Brignole M, et al. 2002 12475469 (233)	Aim: Whether handgrip or arm-tensing would increase BP during impending syncope and avoid LOC  Study type: Randomized;	Inclusion criteria: ≥1 episode of syncope; ≥1 syncopal episodes preceded by prodromal; syncope reproduced during 2 tilt tests performed on different days	Intervention: Hand-grip or arm-tensing  Comparator: Placebo	1° endpoint: Syncope or presyncopal recurrence with maneuver  1° Safety endpoint: N/A	63% in the active arm became asymptomatic vs. 11% in control (p<0.02); 5% vs. 47% developed syncope (p=0.01).     F/U 9±3m 99% performing maneuver prevented syncope

	single blind placeho	<b>∖10</b> ,,			
	single-blind, placebo- controlled; cross-over tilt	≥18 y.			Summary:
	efficacy study	Exclusion criteria: N/A			Isometric arm contraction helps
	enicacy study	Exclusion Citteria. N/A			to abort impending syncope
	Size: n=19 pts				BP increased
Man Dille at al	Aim: Assess effectiveness	Inchesion outtonio December	Internation Converting	40 1 1 0	
Van Dijk, et al. 2006		Inclusion criteria: Recurrent	Intervention: Conventional	1° endpoint: Syncope recurrence	• 32% PCM vs. 51% control
	of PCM in daily life	syncope and prodome (≥3	therapy+ PCM (leg-crossing,		(p=0.005); median yearly
<u>17045903</u>	Charles towns - Donale asiand	syncope episodes in 2 y or	hand grip, arm tensing	1° Safety endpoint: N/A	syncope burden lower in PCM
(234)	Study type: Randomized	(≥1 syncope and ≥3	Commenter Commenteral		group (p<0.004); RRR: 39% in
	(multicenter)	presyncope in 1 y	Comparator: Conventional		PCM group.
	000 1		therapy		
	Size: n=223 pts	Exclusion criteria: Heart			
	standard n=117;	disease, OH, other causes for			Summary:
	standard+PCM n=106	syncope, life-expectancy <1;			PCM effective, safe in VVS
		unable to follow-up			with prodrome.
Foglia-Manzillo, et	Aim: Efficacy of tilt training	Inclusion criteria: Recurrent	Intervention: Tilt-training	1° endpoint: Positive tilt test;	• F/U 1 y; syncope recurrence
al.	in preventing tilt-induced	syncope; 2 consecutive	(30min standing against wall 6	syncope recurrence	28%; presyncope 45%; 17%
2004	syncope	positive nitrate-potentiated	days a 1 wk x 3 wk).		performed tilt-training; of the 5
<u>15121070</u>		head-up tilt test		1° Safety endpoint: N/A	compliant 3 neg tilt table; none
(235)	Study type: Randomized				had recurrence.
	(multicenter)	Exclusion criteria: Other	Comparator: No tilt-training		
		causes of syncope			Summary:
	Size: n=68 pts; tilt-training				Tilt-training not effective in
	n=35; controls n=33				reducing tilt-testing positivity
					because of poor compliance.
On YK, et al.	Aim: Effectiveness of	Inclusion criteria: VVS by	Intervention: Daily sessions x 4	1° endpoint: Tilt response at 1 min;	• 56% positive HUT in training
2007	repeated home orthostatic	positive HUTT	wk. Standing against wall 1–	syncope recurrence	group and 53% in control
<u>17461874</u>	self-training		2 times a day until prodrome of		(p=0.85); syncope or pre-
(236)		Exclusion criteria: Other	for up to 30 min	1° Safety endpoint: N/A	syncope occurred in 42.9% vs.
	<b>Study type</b> : Randomized	causes of syncope after			41.5% controls (p=0.82) during
		comprehensive evaluation,			16.9 m of F/U.
	Size: n=33 pts; tilt-training	structural heart disease.	Comparator: No tilt-training		
	n=16; control n=17				Summary: Tilt-training
					ineffective in reducing positive
					HUT response.
Duygu, et al.	Aim: Effectiveness of	Inclusion criteria: Recurrent	Intervention: Conventional+tilt-	1° endpoint: Syncope recurrence	<ul> <li>Follow up 12+2 m; syncope</li> </ul>
2008	repeated orthostatic self-	syncope ( <u>&gt;</u> 2 events in 6m)	training (Standing against wall 1-		recurrence 56% control and

18439174 (237)	training  Study type: Randomized  Size: n=82 pts; 1:1	and + HUTT  Exclusion criteria: Other causes of syncope after comprehensive evaluation	2X a d until prodrome of for up to 30 min x 1m; then every other day x2 m then 2x a wk)  Comparator: Conventional	1° Safety endpoint: N/A	37% tilt-training (p=0.1); frequency of recurrence similar in all types of VVS; rate of episodes higher in vasodepressor type.  Summary: Tilt-training did not reduce
Salim, et al. 2005 15708690 (238)	Aim: Effectiveness of salt and fludocrotisone in prevention of VVS in children  Study type: Randomized (pediatric)  Size: n=32 pts; florinef 0.1mg/day and salt 1g/d n=18; control n=14	Inclusion criteria: ≥1 syncope or presyncope; +HUTT; <18 y of age; no prior therapy for syncope  Exclusion criteria: No structural heart disease	Intervention: florinef 0.1mg/day and salt 1g/d  Comparator: Placebo	1° endpoint: Syncope or pre-syncope recurrence  1° Safety endpoint: N/A	syncope recurrence  • Follow up 176±117d; recurrence 36% in controls and 55% active arm (p<0.04).  Summary: Symptoms were more frequent in the placebo group.
Romme JJ, et al. 2011 21752826 (239)	Aim: Effectiveness of midodrine in pts not responding to non-pharmacological treatment (STAND-trial)  Study type: randomized, double-blind crossover (3 m then 1 wk washout)  Size: n=23 pts	Inclusion criteria: ≥3 syncope in 2 y; prodrome in 80% episodes; +HUTT  Exclusion criteria: LOC not due to VVS; already using pharmacotherapy for rx VVS	Intervention: Midodrine  Comparator: Placebo	1° endpoint: Recurrence of syncope or presyncope, side effects and QoL  1° Safety endpoint: N/A	Syncope and presyncope recurrence did not differ between treatment (48 vs. 65%, p-0.22); (74 vs. 78%, p=0.90) Side effects and QoL did not differ.  Summary: Addition of midodrine to non-pharmacological therapy not effective
Kaufman H, et al. 2002 12205647 (240)	Aim: Efficacy of midodrine  Study type: Randomized, double-blind cross-over  Size: n=12 (5 mg or placebo day 1 and opposite on day 3)	Inclusion criteria: ≥2 syncope in 1 y; +HUT Exclusion criteria: N/A	Intervention: Midodrine  Comparator: Placebo	1° endpoint: Recurrence of syncope 1° Safety endpoint: N/A	Midodrine produced no significant change in BP or heart rate     Response to HUT: NMS 67% on placebo and 17% on midodrine (p<0.02)

	and 1 h after HUTT				
Perez-Lugones, et al. 2001 11513446	Aim: Efficacy of midodrine  Study type: Randomized	Inclusion criteria: ≥1 syncope per mo and (2) a positive HUTT.	Intervention: Midodrine (5 mg pot id titrated up to 15 tid if required) q 6 daytime	1° endpoint: Syncope recurrence 1° Safety endpoint: N/A	F/u 6m; 81% midodrine and 4% in conventional remained asymptomatic (p<0.001)
(241)	Size: n=61 pts; midodrine n=31; conventional n=30	Exclusion criteria: 1) other causes of syncope; 2)CVD and /or systemic disease; or 3) SBP150 mmHg or dBP 95 mmHg	Comparator: Conventional		<b>Summary</b> : Midodrine provides a significant benefit compared to conventional therapy.
Ward, et al. 1998 <u>9505918</u> (242)	Aim: Benefit of midodrine on symptom frequency and hemodynamic response during HUTT  Study type: Randomized (double-blind placebo controlled cross over)  Size: n=16 pts	Inclusion criteria: >2 presyncope or syncope; no HTN meds; reproducible syncope with GTN on HUTT  Exclusion criteria: Did not meet inclusions	Intervention: midodrine x 1 mo  Comparator: Placebo	1° endpoint: Symptom frequency and hemodynamic response HUTT  1° Safety endpoint: N/A	Midodrine 7.3 symptom free days than placebo (p<0.0001); QoL improved with midodrine; 14 placebo group tilt-induced syncope vs. 6 midodrine (p=0.01)      Summary: Midodrine associated with reduced symptom frequency; symptom HUTT and improved QoL.
Qingyou, et al. 2006 <u>17137891</u> (243)	Aim: Effectiveness of midodrine in prevention of VVS in children  Study type: Randomized (open-label) (pediatric)  Size: n=26 pts; midodrine+ conventional n=13; conventional n=13	Inclusion criteria: ≥3 syncope/y  Exclusion criteria: Other causes of syncope after comprehensive evaluation	Intervention: conventional + midodrine (1.25 mg bid if +HUTT after 1wk then increased 2.5 mg bid then another med added if still +HUTT after 1 wk)  Comparator: Conventional	1° endpoint: Syncope recurrence 1° Safety endpoint: N/A	Follow up 10±8 m; 80% controls vs. 22% midodrine (p=0.023)      Summary: Midrodrine effective in treating VVS in children.

Madrid, et al. 2001	Aim: Efficacy of atenolol	Inclusion criteria: ≥2 syncope 1 y	Intervention: Atenolol 50 mg/d	1° endpoint: Time to syncope recurrence	• ITT, syncope recurrence similar both groups; KM p value
<u>11216978</u>	Study type: Randomized		Comparator: Placebo		0.45 for time to first recurrence
(244)	(double-blind and placebo-	Exclusion criteria: PAD, DM,		1° Safety endpoint: N/A	
	controlled)	AV disease, autonomic			Summary: Recurrence of
		dysfunction, neoplastic or			syncope similar in pts treated
	Size: n=50 pts; atenolol	psych, drug addiction			with atenolol compared to
	n=26; placebo n=24				placebo.
Flevari, et al.	Aim: Efficacy of	Inclusion criteria: ≥2	Intervention: Propranolol,	1° endpoint: Syncope and pre-	Follow up 3m periods
2002 12142117	propranolol, nadolol and placebo in recurrent VVS	syncope 3m; +HUTT	nadolol, placebo 3 mo cross- over	syncope recurrence	syncope and pre-syncope
(245)	placebo in recurrent vv3	Exclusion criteria:	over	1° Safety endpoint: N/A	reduced by all drugs; [ANOVA]: chi-square =67.4; p<0.0001 for
(240)	Study type: Randomized 3	Autonomic failure, HTN,	Comparator: See above	1 Salety enupoint. N/A	syncopal attacks; chi-square
	mo cross-over	COPD, PVD	<u> </u>		=60.1; p<0.0001 for
		,			presyncopal attacks
	<b>Size</b> : n=33				' ' '
					Summary: B-blockers and
					placebo equally effective in
					decreasing syncope and pre-
					syncope
Brignole, et al.	Aim: Efficacy of medical	Inclusion criteria: Frequent,	Intervention: Drugs: atenolol	1° endpoint: Syncope recurrence	• Follow up 10±7m; absence of
1992	treatment in preventing VVS	unexplained syncope or pre-	n=7;dihyroergotamine	40.064	syncope recurrence after 20m
<u>1632399</u> (246)	Study type: Randomized	syncope; 2 +HUTT	n=2;domperidone n=2; cafedrine n=1; stocking ± drug n=3	1° Safety endpoint: N/A	70% treatment and 67%
(240)	<u>Study type</u> . Randomized	Exclusion criteria: Other	11-1, Stocking ± drug 11-3		placebo
	<b>Size:</b> n=30 pts; 1:1	causes of syncope after	Comparator: Placebo		Summary: Outcomes similar in
	<u> </u>	comprehensive evaluation	<u>comparator</u> i lacosc		either medically treated or
					placebo groups.
POST	Aim: Effectiveness of b-	Inclusion criteria: ≥2	Intervention: Metoprolol	1° endpoint: Syncope recurrence	• 36% in control and 36%
Sheldon, et al.	blockers in prevention VVS	syncope over lifetime or ≥1			metoprolol (p=0.99)
2006		syncope 6 mo; +HUTT	Comparator: Placebo	1° Safety endpoint (: N/A	,
<u>16505178</u>	Study type: Randomized				Summary:
(247)	(multicenter)	Exclusion criteria: Oother			Syncope recurrence did not
	<b>6:</b>	cause of syncope; PPM,			differ between metoprolol or
	Size: n=208 pts; metoprolol	contraindication to b-blocker;			placebo groups.
	n=108; placebo n=100	prior trial b-blocker ≥25 mg bid			
		Diu			

Theodorakis, et al. 2006 16627439 (248)	Aim: Effectiveness of placebo, propranolol, fluoxetine in VVS  Study type: Randomized (multicenter)  Size: n=96 pts; placebo n=22; propranolol n=24; fluoxetine n=30	Inclusion criteria: ≥5 syncope lifetime or ≥2 in 1 y, last 1m prior; no drugs  Exclusion criteria: Other cause of syncope; contraindications to study medications	Intervention: Placebo, propranolol, fluoxetine  Comparator: See above	1° endpoint: Syncope or pre-syncope recurrence 1° Safety endpoint: N/A	41% controls, 51% metoprolol, 22% fluoxetine, log rank p>0.05; well-being improved in the fluoxetine group (p<0.01) before and after treatment.      Summary:     Fluoxetine equivalent to propranolol and placebo; effective for reducing presyncope; improves well-being.
Takata TS, et al. 2002 <u>12234955</u> (249)	Aim: Effect of fluoxetine on CV reflexes  Study type: Randomized (double-blind)  Size: n=19; control n=10; fluoxetine n=9	Inclusion criteria: Healthy; +CSM or LBNP (lower body negative pressure)  Exclusion criteria: Psychiatric, neurological or cardiac disease, prior SSRI or MOI	Intervention: Fluoxetine 20 mg daily  Comparator: Placebo	1° endpoint: Syncope  1° Safety endpoint: N/A	Decreases arterial baroreceptor sensitivity but does not prevent presyncope LBNP      Summary: Prevention of presyncope does not occur in LBNP.
Di Girolamo, et al. 1999 <u>10193720</u> (250)	Aim: Effectiveness of paroxetine in VVS resistant to other drugs  Study type: Randomized  Size: n=68;1:1	Inclusion criteria: Recurrent syncope; failed conventional therapy; +HUTT  Exclusion criteria: Other causes of syncope after comprehensive evaluation (EPS); depression or panic disorder	Intervention: Paroxetine 20 mg daily  Comparator: Placebo	1° endpoint: Syncope recurrence  1° Safety endpoint: N/A	•17.6% paroxetine vs. 52.9% placebo (p<0.0001)      Summary: Paroxetine improves recurrence in pts intolerant to conventional therapy.
Gaggioli, et al. 1997 <u>9352988</u> (251)	Aim: To determine the effect of vasodilator therapy on upright tilt testing for syncope  Study type: Case-control randomized study	Inclusion criteria: 1) ≥1 episodes of syncope occurring during chronic (>6 m) vasodilator treatment with angiotensin-converting enzyme inhibitors, long-acting nitrates, or calcium antagonists, or an association	Intervention: Vasodepressor therapy continued  Comparator: Vasodepressor therapy discontinued	1° endpoint: Vasovagal reaction during upright tilt testing 2 wk after randomization  1° safety: N/A	Results: TTT positive in 85% who continued vasodepressor therapy and 52% who discontinued (p=0.02); type of medication did not influence results  Summary: Chronic vasodilator

<b>Size</b> : n=45	of these or with diuretics, all		therapy enhances susceptibility
<u>0126.</u> 11–40	given within the		to VVS during TTT.
	recommended dosage range;		to vvo during 111.
	2) positive response to upright		
	TTT performed during the		
	same treatment which had		
	been administered at the time		
	of the occurrence of the		
	spontaneous syncopal		
	spell(s); and 3) negative work-		
	up for other causes of		
	syncope.		
	Facilities esitente.		
	Exclusion criteria:		
	Identifiable causes of syncope		
	1) OH, which was defined as		
	a decline 220 mm Hg in SBP,		
	or ~10 mm Hg in DBP, within		
	3 min of standing or using a		
	tilt table in the head-up		
	position, at an angle of ~60"		
	presence of important		
	clinical conditions		
	contraindicating the		
	interruption of vasodilator		
	therapy, namely, overt HF,		
	severe hypertension, etc; (3)		
	recent (within the previous 6		
	mo) MI or stroke or other		
	diseases; (4) very severe		
	general diseases; (5)		
	concomitant therapy with BB		
	or any other vasoactive drugs;		
	and (6) intermittent or		
	discontinuous vasodilator		
	administration.		

## Data Supplement 26. Nonrandomized Trials, Observational Studies, and/or Registries of Vasovagal Syncope – (Section 5.1.1)

Study Acronym; Author; Year Published	Study Type/Design*; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Pitt, et al. 2004 <u>15316842</u> (252)	Study type: Observational Determine whether syncope pts and control pts show different responses of BP to postural maneuvers; carbohydrate or water  Size: n=7 pts	Inclusion criteria: syncope or presyncope related to upright posture; ≥episode in prior 6 m; +HUTT: drop in SBP <80 mmHg with symptoms  Exclusion criteria: Evidence of cardiac or neurological etiology on work-up	1° endpoint: BP response  Results: Carbohydrate: 85% meal or 500 ml of tap water alternated 1-2 wk; before and after crouching  • Before meal or water no difference btw groups in BP or in response to maneuvers; in pts standing BP did increase after water; BP after crouch increased largely after meal but smaller after water.	In pts with posturally related syncope unlike in control; carbohydrate ingestion and water result in opposite effects on BP during postural maneuvers.
Krediet, et al. 2002 <u>12270863</u> (253)	Study type: Observational Effects of leg crossing and lower body tensing 30s  Size: n=21 pts	Inclusion criteria: Recurrent VVS syncope; positive tilt table  Exclusion criteria: Other causes of syncope after comprehensive evaluation	1° endpoint: Syncope or presyncope recurrence after tilt test and use of counter-maneuvers  Results: 5/20 (25%) vasovagal reaction averted by maneuver prior to tilt; In follow up (10m) 13 pts used counter-maneuver in daily life and 2 fainted; 10 with presyncope benefited.	Counter-maneuvers can help to alleviate prodromal symptoms and can prevent in some recurrent syncope.      BP increased
Di Girolamo, et al. 1999 10534467 (254)	Study type: Controlled Study, standing against wall up to 40 min  Size: n=47 pts; consent n=24 and refusal (n=23)	Inclusion criteria: Refractory VVS syncope; positive nitrate-potentiated head-up tilt test  Exclusion criteria: Other causes of syncope after comprehensive evaluation	1° endpoint: Syncope recurrence  Results: HUTT response evaluated at 1m: 26.1% of control group and 95.8% of training group became tilt-neg (p<0.0001); syncope recurrence (18.2±5.3 m) 56.3% control vs. 0% in training group (p<0.0001)	Tilt training significantly improves symptoms in those unresponsive or intolerant of medications.
Reybrouck, et al. 2002 12418741 (255)	Study type: Observational (long term f/u); 1-2m against will  Size: n=38	Inclusion criteria: Recurrent VVS syncope and positive tilt without pharmacological provocation  Exclusion criteria: Other causes of syncope after comprehensive evaluation	1° endpoint: Syncope recurrence  Results: Follow up (43±7.8 m); 29/38 abandoned tilt training; 82% free of syncope; 6/7 recurrent syncope discontinued training; 19 compliant for 1 y no syncope recurrence reported	Syncope recurrence may improve symptoms.
Kinay, et al. 2004 <u>15557724</u> (256)	Study type: Observational In-hospital training: 3 consecutive session w/o	Inclusion criteria: Recurrent VVS syncope; positive nitrate-potentiated head-up tilt test	1° endpoint: Syncope recurrence  Results: F/U 356 ±45 d;81% free of recurrent syncope.	Short-term tilt-training is effective.

	syncope; home: 2 session standing against wall 15 min 2m; no activity	Exclusion criteria: Other causes of syncope after comprehensive evaluation		
	<b>Size</b> : n=32			
Samniah, et al. 2001	Study type: Observational	Inclusion criteria: Recurrent VVS syncope ≥1 y; failed ≥2 meds	1° endpoint: Syncope recurrence	Midodrine effective and safe in pts with VVS refractory to standard drug therapy.
<u>11423066</u>	<b>Size</b> : n=20		<b>Results:</b> Follow up 21.9 (15,36); 14/18 resolution	
(257)		Exclusion criteria: BP >160/90; symptomatic IHD; CVA	of symptoms; 4 partial response	
Sheldon, et al.	Study type: Non-	Inclusion criteria: ≥2 VVS syncope or 1	1° endpoint: Syncope recurrence	B-blocker may not have significant
1996	randomized	syncope and ≥4 presyncope; +lso HUTT		effects in preventing syncope recurrence
<u>8806338</u>			Results: Event occurred 17/52 b-blockers; 28/101	after a positive HUT.
(258)	Size n=153; 52 received b-	Exclusion criteria: Other causes of	pt control; actuarial probability of remaining	
	blocker; 101 control	syncope after comprehensive evaluation	syncope similar in both groups	
Sheldon, et al.	Study type: Post-hoc	Inclusion criteria: Obs: ≥2 VVS	1° endpoint: Syncope recurrence	B-blocker prevents syncope recurrence
2012	POST; retrospective	syncope or 1 syncope and ≥4		in middle-aged pts (>42 y of age).
<u>22972872</u>	observational	presyncope or 1 syncope with trauma;	Results: A pooled analysis of both studies yielded	
(259)		+HUTT	an estimate of the HR: 1.58 (CI: 1.00–2.31) for <42	
	<b>Size:</b> n=160; BB=52 in obs; POST n=108; <42 or	Inclusion POST	y, and HR: 0.52 (CI: 0.27–1.01) for ≥42.	
	>42 y of age	Exclusion criteria: Other cause of		
		syncope; PPM, contraindication to b-		
		blocker; prior trial b-blocker <a>25</a> mg bid		

## Data Supplement 27. RCTs Comparing Pacemakers in Vasovagal Syncope – (Section 5.1.2)

Study Acronym Author Year	Aim of Study; Study Type*; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator ( # patients)	Endpoint Results (include Absolute Event Rates, P value; OR or RR; and 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Connolly, et al.	Aim: Effectiveness of PPM	<u>Inclusion criteria</u> : ≥6 lifetime syncope;	Intervention: PPM with	1° endpoint:	• Adjusted RRR 90.8% (CI: 71.0%–
1999	compared with	+HUTT	rate drop	Syncope recurrence	97.1%, p<0.0001); effect on presyncope
<u>9935002</u>	pharmacological therapy in				NS (p=0.56)
(260)	recurrent VVS	Exclusion criteria: Other causes of	Comparator: Placebo		Mean age no PPM 40 and PPM 46 y of
		syncope after comprehensive evaluation			age.

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	Study type: Randomized  Size: n=54;1:1 (terminated early)				Summary: In severely symptomatic, PPM significantly reduces syncope recurrence.
Sutton R, et al. 2000 10899092 (261)	Aim: Effectiveness of DDI pacemaker with rate drop on syncope recurrence  Study type: Randomized (multicenter)  Size: n=42; PPM n=19; no PPM=23	Inclusion criteria: :≥3 syncope 2 y; + cardio inhibitory response (HUTT)  Exclusion criteria: Other causes of syncope after comprehensive evaluation; recent MI, HF (NYHA III-IV), chronic disease	Intervention: DDI + hysteresis  Comparator: placebo	1° endpoint: Syncope recurrence	•1 (5%) PPM vs. 14 (61%) non-PPM, p<0.0006; KM 1,3,5 y 0%,6%, 6% PPM and 39%, 50%, 75% no PPM (p=0.0004) • Mean age no PPM 56 and PPM 64 y of age.  Summary: In those with cardio-inhibitory response, DDI pacing with hysteresis reduces likelihood of syncope.
Ammirati, et al. 2001 11435337 (262)	Aim: Effectiveness of PPM compared with pharmacological therapy in recurrent VVS  Study type: Randomized (multicenter)  Size: n=93; PPM n=46; no PPM n=47 terminated early	Inclusion criteria: >35 y of age; ≥3 syncope 2 y; +HUTT with syncope and bradycardia  Exclusion criteria: Other causes of syncope after comprehensive evaluation	Intervention: PPM with rate drop  Comparator: atenolol	1° endpoint: Syncope recurrence	•2 (4.3%) PPM vs. 12 (25.5%) drug; OR 0.133 (0.028–0.632), p=0.004.  Summary: DDD with rate drop more effective than atenolol for prevention of syncope.
Connolly, et al. 2003 12734133 (263)	Aim: If pacing reduces syncope recurrence  Study type: Randomized (multicenter, double-blind)  Size: n=100 pts; DDD n=48; ODO n=52	Inclusion criteria: ≥6 lifetime syncope;3 in 3 y; +HUTT  Exclusion criteria: Other causes of syncope after comprehensive evaluation, valvular, coronary, myocardial, major non CVD, ECG abnormalities	Intervention: DDD with rate drop  Comparator: ODO	1° endpoint: Syncope recurrence	Cumulative risk at 6m 40% (25%–52%)     ODO and 31% (-33%–63%) DDD, p=0.14.      Summary: Pacing did not reduce risk of recurrent syncope.
Raviele A, et al. 2004 15451153 (264)	Aim: if pacing reduces syncope recurrence  Study type: Randomized (multicenter, double-blind,	Inclusion criteria: ≥6 lifetime syncope; 1 in last y; +HUTT(asystole or mixed)  Exclusion criteria: Other causes of syncope after comprehensive evaluation	Intervention: DDD with rate drop  Comparator: OOO	1° endpoint: Syncope recurrence	Follow up med 715d, 8(50%) on vs. 5(38%) off (p=NS); no difference in the mixed and asystole subgroups.      Summary: Active pacing was not

	placebo-controlled)  Size: n=29 pts; on n=16; off n=13				significant associated with reduction in syncope recurrence compared to inactive pacing.
Brignole, et al. 2012 22565936 (265)	Aim: effectiveness of cardiac pacing in NMS and asystole  Study type: Randomized (multicenter, double-blind, placebo-controlled)  Size: n=77; on n=16; off n=13	Inclusion criteria: :≥40 y of age; ≥3 syncope in 2 y; ILR with ≥3s asystole or ≥6s asystole w/o syncope  Exclusion criteria: ≥1 cardiac abnormalities that suggested cardiac syncopesinus bradycardia <50 bpm or sinoatrial block; Mobitz I second-degree AV block; BBB; rapid paroxysmal SVT or VT; preexcited QRS complexes; prolonged QT interval; BS; ARVC)· nonsyncopal loss of consciousness; CSS	Intervention: DDD with rate drop  Comparator: ODO	1° endpoint: Syncope recurrence	•2 y estimated recurrence 57% (40%-74%) ODO and 25% (13%-45%) DDD, p=0.039. absolute RR 32% and relative RR 57% with DDD  Summary: DDD effective in reducing recurrence of syncope in ≥40 y of age with severe asystolic component.
Flammang, et al. 1999 11228858 (266)	Aim: Effectiveness of pacing in symptom recurrence with abnormal adenosine 5-triphosphate  Study type: Randomized (open label)  Size: n=20; Dual chamber Pacemaker on n=10; Pacemaker off n=10	Inclusion criteria: VVS and abnormal cardioinhibitory (i.e. electrocardiographic) response during ATP test.  Exclusion criteria: Syncope due to neurological, metabolic or arrhythmic etiology	Intervention: Pacemaker on  Comparator: Pacemaker off	1° endpoint: Syncope recurrence	Follow up mean 52m; syncope recurrence PPM 0 (0%); No PPM 6 (60%)     All-cause mortality: Pacemaker 3 (30%); No Pacemaker 1 (10%)      Summary:     PPM in pts with abnormal ATP have fewer syncope recurrences
Flammang, et al. 2012 22086879 (267)	Aim: effectiveness of pacing in unexplained syncope and positive adenosine 5-triphosphate  Study type: Randomized (single blind ,multicenter)  Size: n=80; active n=39; passive n=41	Inclusion criteria: syncope of unknown origin; AV or SA block >10s under ATP administration  Exclusion criteria: ≥1+EPS, carotid sinus hypersensitivity, sustained or episodic atrial or VT documented sinus or AV node conduction disorders (including first-degree AV block), + PPM and ICD ,heart transplant list, pregnancy, asthma or	Intervention: DDD 70 bpm  Comparator: back-up 30 bpm	1° endpoint: Syncope recurrence	• Follow up mean 16m; 8/39 (21%) active vs. 27/41 (66%) HR: 0.25 (0.12–0.56)  Summary: Dual chamber PPM reduces syncope by 75%.

Occhetta, et al. 2004 15519257 (268)	Aim: To determine whether dual-chamber rate adaptive CLS prevents recurrence of VVS CLS – tracks variation of intracardiac impedance during systolic phase of cardia cycle on beat-to-beat basis; activates AV sequential pacing when detecting increased contractility during early phase of VVS.  Study type: Randomized (single blind, multicenter)  Size: n=26 pts; active n=9; control n=17	severe chronic bronchitis, systemic infection, or DM  Inclusion criteria: >5 syncopal episodes and/or >2 in the last y before enrolment; refractoriness to conventional drug therapy and tilt-training+HUTT with cardio inhibition (+2A or 2B VASIS).  Exclusion criteria: previous MI, CHF, severe chronic disease	Intervention: DDD  Comparator: DDI (40 bpm)  Randomization between DDD (9/26) (17/26) and DDI only during 1st y  24 pts recruited in 2nd y programmed to DDD-CLS	1° endpoint: 2 VVS during 1 y follow-up.	Follow up mean 44 m; 7/9 DDI had met primary endpoint; 41 pts programmed to DDD-CLS none had VVS      Summary:     Effectiveness of DDD-CLS in preventing VVS with cardioinhibition
Russo, et al. 2013 <u>23723446</u> (269)	Aim: The effect of dual-chamber CLS in the prevention of syncope recurrence in refractory VVS  Study type: Randomized (single blind, crossover)  Size: n= 50 pts	Inclusion criteria: >40 y of age; sinus rhythm; recurrent unpredictable syncope; no medications that could affect circulatory control; refractoriness to conventional drug therapy and/or tilt-training; +HUTT with cardioinhibition - asystole >3 s ( 2B VASIS)  Exclusion criteria: other causes of syncope after comprehensive evaluation	Intervention: DDD CLS on  Comparator: DDD CLS off	1º endpoint: Syncope recurrence in the CLS on and off phases	Pts with syncope recurrence at 18 mo: Pacemaker CLS ON 1 (2%); Pacemaker CLS OFF 8 (16%) Pts with presyncope at 18 mo: Pacemaker CLS ON 4 (8%); Pacemaker CLS OFF 18 (27.8%)  Summary: Effectiveness of DDD-CLS in preventing VVS with cardioinhibition

## Data Supplement 28. Nonrandomized Trials, Observational Studies, and/or Registries of Pacemakers in Vasovagal Syncope – (Section 5.1.2)

Study Acronym (if applicable) Author; Year	Study Type/Design*; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and 95% CI)	Summary/Conclusion Comment(s)
Deharo, et al. 2013	Study type: Observational	Inclusion criteria: Sudden onset syncope without prodrome and normal	1° endpoint: Pathophysiology of sudden-onset syncope	Low adenosine plasmatic levels defines distinct form syncope from

23810895 (270)	Size: n=15 pts with syncope without prodrome and normal heart and ECG compared to n=31 VVS	heart and ECG; VVS  Exclusion criteria: Other causes of syncope after comprehensive evaluation	Results: Study group- lower median adenosine plasmatic level; ≤0.36 umol/l 73% sensitivity; 93% specificity	VVS.
Brignole, et al. 2011 <u>21570228</u> (271)	Study type: Observational Size: n=18 pts	Inclusion criteria: Syncope; normal ECG, no structural heart disease, paroxysmal 3AVB associated with syncope	1º endpoint: Clinical characteristics unexplained syncope with paroxysmal AVB	Efficacy of PPM in idiopathic AVB.
		Exclusion criteria: Other causes of syncope after comprehensive evaluation	Results: Follow up mean 4±4 y; AVB without P-P cycle or PR interval prolongation; 17 pts had dual-chamber PPM no syncope recurrence.	
Lelonek M, et al. 2007 (272)	Study type: Observational  Size: n=34 pts Pacemaker n=22 (DDI +hysteresis) No pacemaker n=12 (pharmacological: midodrine or b-blocker) -all educated on behavior measures	Inclusion criteria: Tilt-induced cardio depressive syncope with asystole >3 s (2B VASIS)  Exclusion criteria: Other causes of syncope after comprehensive cardiac	1° endpoint: Syncope recurrence  Results: Syncope recurrence at 18 mo: Pacemaker 5 (23%); No pacemaker 3 (25%); p>0.05	Pacemaker or pharmacological treatment effective
	-all educated on behavior measures	and neurological evaluation	No injury in either group	

## Data Supplement 29. RCTs Comparing Carotid Sinus Syndrome – (Section 5.1.3)

Study Acronym Author Year	Aim of Study; Study Type*; Study Size (N)	Patient Population	Study Intervention (include # patients) / Study Comparator (include # patients)	Endpoint Results (include Absolute Event Rates, P value; OR or RR; and 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Brignole, et al.	Aim: Efficacy of permanent	Inclusion criteria: Recurrent	Intervention: Pacing	1° endpoint: Symptom	Syncope recurrence in 57% of the non-
1992	pacing.	syncope or presyncope causing		recurrence	pacing group and 9% of the pacing group
<u>1561975</u>	<b>2</b> 4 1 4 507	trauma or future trauma or	Comparator: No pacing		(p=0.0002); the actuarial rate of absence of
(273)	Study type: RCT	decreased QoL; cardioinhibitory		1° Safety endpoint: N/A	syncopal recurrence after 1,2,3 and 4 y was
	Size: n=60 pts; no	or mixed symptoms reproducible CSM; no other cause (extensive			64%, 54%, 36%, and 38%, respectively, for
	pacing=28; pacing 32	w/u monitoring, neuro, EPS)			the nonpacing group, and 100%, 97%, 93%, and 64%, respectively, for the pacing group
	(VVI=18, DDD=14)	w/a morntoring, nearo, Er 3)			(p=0.0001).
	(*** 10, 555-14)	Exclusion criteria: SN			(ρ-0.0001).
		dysfunction, prolonged HV AV			Summary: Permanent pacing effective in
		block on EPS			CSS

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Ryan, et al. 2010 <u>19933747</u> (277)	Aim: Cardiac pacing for recurrent falls in pts with cardioinhibitory CSH would reduce fall recurrence.  Study type: RCT, open label  Size: n=141; ITT n=129; Pacing on n=68 No pacemaker (ILR) n= 61	Inclusion criteria: ≥ 65 y; symptoms consistent with CSH with a minimum of 2 unexplained falls and/or one syncope in prior 1 y; 3 s of asystole in response to CSM; a MMS >19.  Exclusion criteria: Neoplasm, renal or hepatic failure; and at time of randomization significant HF.	Intervention: Pacing Comparator: No pacing	1° endpoint: Number of falls after implant.  2° endpoint: Time to fall event, presyncope, quality of life and cognitive function  1° Safety endpoint: N/A	<ul> <li>(4.5%)</li> <li>Pts with soft tissue injury due to fall at 12 mo: pacemaker 26 (29.9%); no pacemaker 32 (36.4%)</li> <li>All-cause mortality at 12 mo: pacemaker 5 (5.7%); No pacemaker 3 (3.4%)</li> <li>Summary: Pacing associated with less falls and injury; no reduction in syncope events</li> <li>Pts reporting syncope after pacemaker implant RR: 0.47 (95% CI: 0.26–0.86); The number of syncopal events was also significantly less after implant, 0.52 (95% CI: 0.29–0.95).</li> <li>Syncope recurrent events at 24 mo: Pacemaker 0.42 mean events; No pacemaker 0.66 mean events; RR: 0.87 (95% CI: 0.3–2.48)</li> <li>2° endpoints:</li> <li>Pts with falls at 24 mo: Pacemaker 44 (67%); No pacemaker 33 (53%); RR 1.25 (95% CI: 0.93–1.67)</li> <li>Syncope-related falls at 24 mo: pacemaker 4.33 events; No pacemaker 6.52 events; RR: 0.79 (95% CI: 0.41–1.5)</li> <li>Summary: No difference in falls, syncope and other secondary endpoints between 2 groups.</li> </ul>
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# Data Supplement 30. Nonrandomized Trials, Observational Studies, and/or Registries of Permanent Pacing in Carotid Sinus Hypersensitivity – (Section 5.2.2)

Study Acronym (if applicable) Author, Year	Study Type/Design*; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; and 95% CI)	Summary/Conclusion Comment(s)
Sugrue, et al. 1986	<b>Study type</b> : Retrospective, observational study of untreated	Inclusion criteria: ≥1 episodes of syncope or presyncope;	1° endpoint: Symptom recurrence	PPM effective in CSS predominately cardioinhibitory

3941204	compared to pacing or	cardioinhibitory, vasodepressor or	Results: Incidence of recurrence 27% no treatment,	
(278)	anticholinergic drugs	mixed; no other cause	22% drug group, 9% pacing group; those with	
(210)	antionomicigie drugs	mixed, no other eduse	cardioinhibitory CSS had no recurrence of syncope with	
	<b>Size</b> : n=56	Exclusion criteria: N/A	DVI pacing (9/9) and 8 of 10 were asymptomatic with	
	untreated=13	<u>Exolation officina</u> . 1477	VVI pacing	
	anticholinergic=20		, v v pasg	
	pacing=23			
Blanc, et al.	Study type: Retrospective,	Inclusion criteria: Cardio inhibitory	1° endpoint: Symptom recurrence after pacemaker	PPM effective in CSS
1984	observational	,	implant	
6424619		Exclusion criteria: N/A		
(279)	Size: n=54 pts; no pacing=33		Results: 50% of pts had recurrence of syncope with no	
	pacing=21		pacing vs. 0% in pacing group	
Morley, et al.	Study type: Prospective,	Inclusion criteria: Cardioinhibitory	1º endpoint: Symptom persistence, vasodepressor	PPM effective in CSS; AV
1982	observational	with pacemaker	response, pacemaker effect	sequential pacing preferred
<u>7073901</u>				
(280)	Size: n=70 pts; pacing mode	Exclusion criteria: N/A	Results: Persistence of symptoms with a final pacing	
	(VVI, DVI, DDD, AAI)		mode VVI 11%; 8% DVI and 8% DDD, AV sequential	
			pacing eliminated hypotensive effects of VVI pacing	
Gaggioli, et al.	<u>Study type</u> : Retrospective,	Inclusion criteria: Cardioinhibitory	1° endpoint: Symptom recurrence after pacemaker	PPM effective in CSS;
1995	observational	or mixed; no other cause	implant	recurrence does occur in mixed
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(281)	Size: n=169 pts; VVI n=59 DDD n=110	Exclusion criteria: N/A	Results: Syncope recurrence was 7% at 1 y, 16% at 3	
	DDD 11-1 10		y, and 20% at 5 y; 21% syncope recurrence in pts with vasodepressor response.	
Maggi, et al.	Study type: case-control (age-	Inclusion criteria: Cardio inhibitory	1° endpoint: Syncope recurrence	Cardio inhibitory CSS predicts
2007	sex matched 2:1)	CSM and spontaneous syncope by	1 enapoint. Syncope recurrence	associated with asystole during
17507364	SCA Materieu 2.1)	ILR	Results: Asystole 89% CSS and 50% controls; 14 of	spontaneous syncope benefit from
(282)	Size: n=18 pts	Control group: negative CSM, tilt	CSS with asystole DCH PPM; f/u 35 ±22m syncope	pacing
()	<u> </u>	and ATP	burden decreased 1.68 (1.66–1.70) episodes to 0.04	paomig
			(0.038–0.042) with PPM (98% RR)	
		Exclusion criteria: Structural	, , , , , , , , , , , , , , , , , , , ,	
		cardiac disease, conduction,		
		symptomatic OH, non-syncopal		
		cause of LOC		
Lopes, et al.	Study type: Retrospective	Inclusion criteria: Cardio inhibitory	1° endpoint: Symptom recurrence after pacemaker	Permanent pacing effective in
2011	observational	or mixed in whom pacemaker	implant	CSS; recurrence does occur in
<u>21169606</u>	400 /	implanted		mixed type
(283)	<u>Size</u> : n=138 pts		Results: Syncope recurrence 10.9%; 5.8% minor	

		Exclusion criteria: N/A	symptoms/presyncope; mixed CSS predicted recurrence (HR: 2.84; 1.20–6.71; p=0.017)	
Brignole, et al. 2011 21570228 (271)	Study type: Systematic review  Size: 12 studies; n=601 pts with pacing and 305 untreated	Inclusion criteria: Cardioinhibitory or mixed  Exclusion criteria: Case reports	1° endpoint: Syncope recurrence; up to 5 y follow-up  Results: 0–20%in pacing group and 20-60% in untreated group; 3 studies with control groups RR 0.24 (0.12–0.48)	Benefit of cardiac pacing with significant reduction in recurrence; lead to reduced morbidity     Recurrence 20% of paced pts at 5 y
Menozzi, et al. 1993 <u>8237805</u> (284)	Study type: Prospective observational  Size: n=23 pts	Inclusion criteria: Recurrent or severe episodes of syncope and presyncope causing major trauma or risk of death; asystolic response >3 s with CSM or eyeball compression with and without positive head-up tilt test; VVI pacemakers ability to track asystolic episodes.	1º endpoint: Occurrence of asystolic episodes  Results: Follow up 15 ± 7 mo; asystolic episodes occurred in 74% of pts; actuarial estimate of occurrence of asystolic episodes of >3 and >6 s were 82% and 53% after 2 y. 12 episodes >3–6 s (0.7%) and 20 episodes of >6s (43%)	Asystolic response to vasovagal maneuvers predicts occurrence of spontaneous asystolic episodes. Spontaneous episodes are asymptomatic and incidence is low.
Striyger, et al. 1986 2429277	Study type: Prospective observational	Exclusion criteria: No other identifiable cause  Inclusion criteria: Repeated syncope of unknown cause; CSM asystole of >4 sec; cardioinhibitory	1° endpoint: Efficacy of VVI pacing in preventing recurrence	VVI pacing for isolated form of cardioinihibitory syncope results in complete resolution of symptoms.
(285)	Size: n=20 pts	based on EPS  Exclusion criteria: N/A	Results: Mean 20 mo; no pts had reoccurrence of syncope	ounplace recording to a symptome.
Walter, et al. 1978 356576 (286)	Study type: Prospective observational  Size: n=21 pts	Inclusion criteria: Syncope of unknown cause or pre-syncope; CSM ventricular asystole of >3 sec	1° endpoint: N/A  Results: 17 pts had cardio inhibitory, 2 vasodepressor and 2 mixed. 11 pts PPM of these 9 had no further	PPM in cardio inhibitory syncope is associated with less reoccurrences.
,		Exclusion criteria: N/A	symptoms or rare pre-syncopal events; 2 of the pts with PPM had mixed response on CSM and had pre-syncope or syncope related to drop in BP.	
Crilley, et al. 1997 9338027	Study type: Prospective observational	Inclusion criteria: recurrent falls, pre-syncope or syncope and CSM >3 s ventricular asystole	1º endpoint: Outcomes of DCH PPM on elderly with falls, pre-syncope and syncope associated with cardioinhibitory syncope	DCH PPM is effective for hypersensitive cardioinhibitory syncope.
(287)	Size: n=42 pts	Exclusion criteria: N/A	Results: All pts had DDI pacemaker implant; 84% no longer had further syncope mean follow up 10 mo and	

symptoms unchanged in 22%

## Data Supplement 31. RCTs for Type of Permanent Pacing in Carotid Sinus Hypersensitivity – (Section 5.2.2)

Study Acronym Author Year	Aim of Study; Study Type*; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (include Absolute Event Rates, P value; OR or RR; and 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Brignole, et al. 1988 <u>2463565</u> (288)	Aim: Evaluate importance of atrial synchronism for mixed CSS  Study type: RCT (single blind, crossover)  Size: n=23 pts	Inclusion criteria: Mixed CSS  Exclusion criteria: Isolated cardioinhibitory or vasodepressor	Intervention: DVI/DDD  Comparator: VVI	1° endpoint: Symptom recurrence; VA conduction, OH, pacemaker effect  1° Safety endpoint: N/A	DVI vs. VVI, syncope occurred in 0% vs. 13% (p= 0.25); pre-syncope in 48% vs. 74% (p=0.04); DVI was the modepreferred by 64% of pts, remaining 36% did not express any preference (p=0.001).      Summary: DVI/DDD pacing effective in 61% compared to VVI. When pacemaker effect, ventriculoatrial conduction and OH are present, VVI failure is possible, therefore DVI/DDD stimulation is indicated
McLeod, et al. 2012 <u>22548372</u> (289)	Aim: Investigate impact of pacing modes (DDDR, DDR with sudden brady response and VVI) on syncope recurrence and QoL  Study type: RCT (double-blind, sequential cross over – 6 m)  Size: n=21 pts	Inclusion criteria: Cardioinhibitory/ mixed CSS; symptoms reproducible CSM  Exclusion criteria: Isolated vasodepressor response to CSM; another cause for LOC; structural heart disease, PPM	Intervention: DDDR, DDR with sudden brady response and VVI  Comparator:	1° endpoint: Syncope and pre-syncope recurrence; QoL 9SF-36)  1° Safety endpoint: N/A	Frequency of V pacing in VVI mode marginally less than any DDDR modes (p=0.04)  For any pacing mode syncope recurrence (29–2; p<0.001) and presyncope (258–17; p<0.001) reduced  Pacing modality found to marginally increase bodily pain and vitality measures in the DDDR mode  Summary: No clear superiority of one pacing mode over another; QoL overall did not differ

## Data Supplement 32. Observational studies, for Type of Permanent Pacing in Carotid Sinus Hypersensitivity – (Section 5.2.2)

Study Acronym; Author; Year Published	Study Type/Design*; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Madigan, et al. 1984 <u>6702680</u> (290)	Study type: Prospective, observational DVI vs. VVI  Size: n=11 pts	Inclusion criteria: Cardioinhibitory with partial or complete reproduction of symptoms or dizziness, near or syncope compatible with cardiac	<u>1º endpoint</u> : Changes in BP after CSM in pts paced in DVI mode vs. VVI <u>Results:</u> Drop in BP in VVI vs. DVI (59 vs. 37 mm	VVI results in significant hemodynamic compromise resulting in increased symptoms
		origin  Exclusion criteria: N/A	Hg; p=0.001) and a higher rate of symptom persistence (91% vs. 27%; p=0.008)	
Sutton, et al. 1989 (291)	Study type: Case series AAI vs. DDD vs. VVI	Inclusion criteria: syncope or pre- syncope 98%; positive CSM; pacemaker inserted	1° endpoint: Syncope recurrence  Results: Failure to control syncope for various	The most effective pacing mode is DVI/DDD compared with other modes
Bae MH, et al.	Size: n=202 pts Study type:	Exclusion criteria: N/A Inclusion criteria: DS occurring	modes: AAI 50%, VVI 18% and DVI/DDD 9%  1º endpoint: Clinical characteristics ( using standard	DS occurred in older women, MS in
2011 22188510	Retrospective, observational study	during or immediately after defecation and during abdominal cramping or	statistics to compare btw groups)	middle-age men and drinking alcohol precipitator
(292)	comparing defecation, micturition and VVS	urge to defecate; MS - syncope occurring at the beginning of, during, at the termination of, or immediately	Results: DS occurred in older age of diagnosis (p=0.004) and first syncope (p=0.002); younger VVS; male more likely MS (p=0.036); frequency of drinking	
	Size: n= 680 consecutive DS n=38; MS n=38; VVS n=208	after urination  Exclusion criteria: Other cause of	alcohol higher in MS (<0.001) as was CV risk factor/underlying disease (p=0.031)	
		syncope or unknown not consistent with VVS (clinical & HUTT)		

#### Data Supplement 33. RCTs for Neurogenic Orthostatic Hypotension – (Section 6.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Anley C, et al.	Aim: To assess which	Inclusion: All collapsed	Intervention: OT, oral fluid and	1° endpoint: Time to discharge: no	With no difference in time to
2011	treatment protocol for	athletes at 2 Ironman	Trendelenburg position	significant difference between IV (52.5 +/- 18	discharge, but significantly less
<u>20584756</u>	exercise-associated	Triathlon competitions and		min) and OT group (58+/-23 min), p=0.47	fluid given in OT group
(293)	postural hypotension	one ultra-distance footrace	Comparator: IV, intravenous fluid	Secondary: heart rate and BP changes: NS	compared to IV group, the

	results in earlier discharge,  Study type: Analytical, Randomized controlled, prospective cohort  Size: n=28 pts	in 2006 and 2007  Exclusion: Abnormal serum sodium		changes were seen.  • Total volume of fluid in OT group was 204 +/- 149 ml, and was significantly less than IV group 1045+/-185 ml, p<0.001.	probable cause of exercise associated postural hypotension is peripheral vasodilatation resulting in venous pooling
Lu CC, et al. 2008 <u>18772858</u> (230)	Aim: Assess whether glucose water ingestion will reduce orthostatic tolerance in young healthy volunteers  Study type: Analytical, Randomized controlled crossover, prospective cohort  Size: n=15 pts	Inclusion: Healthy male  Exclusion: Hx of syncope, any medications	Intervention: 10% glucose water  Comparator: Pure water 5 min before 70 degree HUTT	<u>1º endpoint</u> : Orthostatic tolerance (time to presyncope during 70 degree HUTT): 13 of 15 (87%) ingesting pure water were able to complete the full tilt without presyncope, but 7 of 15 (47%) ingesting glucose water could complete the full tilt. Test was terminated sooner in glucose water group (40.0±6.9 min) vs. pure water group (43±5.6 min), p=0.008. There was no difference in symptom scores (p=0.26) between 2 groups.	Glucose water attenuates reflex role of PVR during orthostatic stress, perhaps by vasodilatation in splanchnic circulation or raising plasma osmolality which may enhance barorefex control of SNS.
Raj SR, et al. 2006 <u>16785332</u> (294)	Aim: To assess if ingestion of salt with water would increase magnitude of acute pressor response compared with water in OH  Study type: Analytical, randomized controlled, prospective crossover  Size: n=9 pts	Inclusion: OH pts with at least 6 mo Hx of orthostatic symptoms and were ≥18 y of age. All medications that could impair BP regulation were withdrawn for ≥5 half-lives before testing.  Exclusion: None	Intervention: Distilled water mixed with 2 g of NaCl added,  Comparator: 16 ox (473 mL) of distilled water then noninvasive heart rate and BP were measured for ≥ 60 mins after ingestion	1° endpoint: Hemodynamic response to water: SBP increased from 92±8 mmHg at baseline to 129±9 mmHg 30 min after ingestion (p<0.001), and 110±12 mmHg 60 min after ingestion (p=0.022). Plasma norepi significantly increased at 30 min (p=0.018) after water ingestion  1° endpoint: Hemodynamic response to salt water: SBP increased from 94 ±9 mmHg as baseline to 112 ±9 mmHg 30 min after ingestion (p=0.005), and 104 ±9 mmHg (p=0.139)	Water and salt water both increased SBP at 30 min post ingestion, with water having double the effect of salt water. By 60 m, only water ingestion continued to show significant increase in SBP. The osmolality of salt water may have reduced the gastropressor response which likely is not just due to blood volume.
Schroeder C, et al. 2002	Aim: To assess water drinking on orthostatic tolerance in healthy pts	Inclusion: Healthy volunteers	Intervention: 500 mL nonsparkling mineral water at room temperature	1° endpoint: Drinking 500 mL water prolonged time to presyncope in 11 pts from 31 ±3 min to 36 ±3 min (p<0.001). Supine	Water drinking 500 mL increases orthostatic tolerance, with the effect apparently

12451007		Exclusion: Regular		heart rate, BP, SV, and cardiac output were	mediated with factors beyond
(231)	Study type: Analytical,	medication except oral	Comparator: 50 mL	not significantly different with 500 mLwater	increasing plasma volume.
(201)	randomized controlled,	contraceptives	nonsparkling mineral water, then	drinking. With HUTT, 500 mL water drinking	Increase in peripheral resistance
	prospective crossover,		60 degree HUTT for 20 min	blunted decrease in SV from -45+/-2% to -38	and vasoconstrictor tone may
	p. 00p000 0.0000,		followed by LBNP for 10 m each	±3%, p<0.01	have role.
	Size: n=13 pts		at -20, then -40, -60 mmHg		
Jankovic JJ, et al.	Aim: Effect of	Inclusion: At 18 centers	Intervention: Midodrine 2.5 mg,	1° endpoint: Midodrine increased standing	• Scalp tingling (13.5%), supine
1993	midodrine in neurogenic	between 1989 to 1990, OH	5 mg, or 10 mg 3x daily, for 4 wk	SBP by 22 mmHg vs. 3 mmHg for placebo	HTN (8%)
<u>7687093</u>	OH	(≥15 mmHg fall from supine		(p<0.001). Midodrine increased standing DBP	Midodrine significantly improves
(295)		to standing position plus	Comparator: Placebo for 4 wk	by 15 mmHg vs. 3 mmHg for placebo	standing SBP and symptoms of
	Study type: Analytical,	symptoms) due to		(p<0.001). Supine SBP increased 13 mmHg	OH.
	Randomized double-	autonomic failure. (n=18,		vs2mmHg for placebo (p<0.001). Symptom	
	blind placebo	Shy Drager; n=22		improvement was significant with 10 mg for	
	controlled, prospective	Parkinson disease; n=27		blurred vision, syncope, and energy level	
	cohort,	DM)		(p<0.01). Improvement with energy level	
	0. 07 (			occurred with midodrine 2.5 and 5 mg doses.	
	Size: n=97 pts	Exclusion: Pre-existing			
		supine hypertension			
		(>180/110 mmHg), renal or			
		hepatic impairment,			
		pheochromocytoma, or severe cardiac			
		abnormalities			
Jordan J, et al.	Aim: To assess volume	Inclusion: Idiopathic OI	Intervention: Phenylephrine	1° endpoint: At 5 m HUTT compared to	Volume loading, alpha-agonist
1998	loading and alpha-	(>30 bpm increase in heart	(infusion rate increased until	placebo, volume loading significantly blunted	infusion, and alpha-blockade all
9774366	adrenergic agonism in	rate within 5 min of	either heart rate decreased by 5-	the increased upright heart rate (-20+/-3.2	blunted decrease in mean
(296)	idiopathic orthostatic	standing without a	10 bpm or SBP increased by 5-10	bpm, p<0.001) as did phenylephrine (-18+/-3.4	middle cerebral artery velocity
(200)	intolerance	concomitant decrease in	mmHg,	bpm, p<0.001), but effect diminished at end of	(despite worsening systemic
		SBP/DBP >20/10 mmHg);	3,	HUTT.	hemodynamics with alpha-
	Study type: Analytical,	plasma norepi level >600	Comparator 1: Phentolamine		blockade). Excessive
	Randomized placebo	pg/mL with standing; at	(infusion rate increased until heart	Phentolamine significantly increased upright	sympathetic activity contributes
	controlled, cross-	least 6 mo Hx of typical	rate increased by 5–10 bpm or	heart rate at 5 min (20+/-3.7 bpm, p<0.01) and	to decreased cerebral blood flow
	sectional cohort	symptoms of OI with	SBP decreased by 5–10 mmHg)	at end of HUTT (14+/-5 bpm (p<0.05)	during HUTT
		standing, which were	or	compared with placebo. With placebo, mean	
	Size: n=9 pts	significantly relieved by		cerebral blood flow velocity decreased by	
		lying down	Comparator 2: Normal saline	33+/-6% at HUTT, but phenylephrine infusion,	
			(placebo at rate similar to	volume loading, and phentolamine infusion all	
		Exclusion: Systemic illness	pheylephrine or phentolamine	attenuated the decrease in mean middle	

	T	Heat and affect the	[ :=£.=:==\	Language and the second	
		that could affect the	infusion).	cerebral artery velocity with upright posture	
		autonomic nervous system		(p<0.05 for each).	
		(DM, amyloidosis)	Comparator 3: All pts were		
			volume loaded with 2000 mL		
			normal saline over 3 H,then 75		
			degree HUT for 30 m		
Jordan J, et al.	Aim: To assess various	Inclusion: severe OH due	Seated BP effect of Intervention:	1° endpoint: Compared to placebo, the	<ul> <li>Not every pts received each</li> </ul>
1998	medication effect in	to multiple system atrophy	Phenylpropanolamine 12.5 mg	pressor response was significant for	drug so direct comparison was
<u>9727818</u>	severe OH from	or PAF	(25 mg in pts not responsive to	phenylpropanolamine (12.5 mg, standing SBP	not possible. Midodrine was
(297)	autonomic failure		12.5 mg),	+37+/-12 mmHg, p<0.05), yohimbine	described as having similar
, ,		Exclusion: Secondary	Comparator 1: yohimbine 5.4	(standing SBP 36+/-13 mmHg, p<0.05), and	effect to phenylopropanolamine
	Study type:	causes of autonomic failure	mg, Comparator 2: indomethacin	indomethacin (standing +28+/-2 mmHg,	with somewhat less effect seen
	Randomized placebo	(DM, amyloidosis),	50 mg,	p<0.05). Phenylpropanolamine and midodrine	in figure 4, but without specific
	controlled, prospective	contraindications to pressor	Comparator 3: Ibuprofen 600	elicited similar pressor responses. No	hemodynamic numbers.
	cohort,	agents (CAD, CHF)	mg,	association between drug response and	nomedynamie namedie.
	,	3 (,,	Comparator 4: Caffeine 250 mg,	autonomic function testing, or plasma	
	Size: n=35 pts		Comparator 5: Methylphenidate	catecholamine levels	
	<u> </u>		5 mg,	Satisficialism of totals	
			Comparator 6: Midodrine 5 mg		
Kaufmann H, et	Aim: To assess the	Inclusion: Several OH with	Intervention: Midodrine titrated	1° endpoint: Midodrine increased standing	Midodrine improves BP and
al.	effect of midodrine OH	multiple system atrophy, or	from 2.5 mg 4x daily to total daily	BP significantly in 3 of 7 pts (p<0.05) and	symptoms of OH in selected pts
1988	in autonomic failure	idiopathic OH.	dose of 0.5 mg/kg (25-40 mg/d)	these pts reported improved orthostatic	with autonomic failure. Pts with
2452997		idiopatino et il	for 7 days,	symptoms. In 4 pts, fludrocortisone,	increasing severity of autonomic
(298)	Study type: Analytical,	Exclusion: None	io. r dayo,	midodrine, and the combination did not	function may not respond to
(200)	Randomized double-	<u>Exolation</u> . None	Comparator: Placebo	increase standing BP or symptoms, and in	midodrine, and may worsen OH
	blind placebo controlled	Low dose fludrocortisone	<u> </u>	these pts the decrease paralleled decrease in	due to extracellular fluid loss
	crossover, prospective	0.1 mg daily continued		body weight.	due to extracellular fluid 1033
	cohort,	o. i mg dany continued		body weight.	
	COTION,				
	Size: n=7 pts				
Low PA, et al.	Aim: Assess midodrine	Inclusion: 18 y of age or	Intervention: Midodrine 10 mg	1° endpoint: Primary: improvement in	Piloerection 13%, pruritus
1997	in neurogenic OH	older, symptomatic	3x daily	standing SBP: mean increased SBP of 21.8	(scalp) 10%, paresthesia 9%,
9091692		neurogenic OH (due to a	on daily	mmHg, p<0.001. Midodrine effect was	supine HTN 4%
(299)	Study type: Multicenter	structural lesion of	Comparator: Placebo	independent of fludrocortisone (mean dose	Supino IIIIV = /0
(200)	analytical, randomized	adrenergic pathways,	<u> </u>	0.35+/-0.33 mg) and independent of wearing	Midodrine 10 mg 3 x daily
	double-blind placebo	central or peripheral), ≥15		compression garments. Symptoms of	increases standing BP and
	controlled, prospective	mmHg SBP postural		lightheadedess improved over entire study,	improves symptoms of OI.
	cohort			and reached significance at second wk of	improves symptoms or Or.
	COHOIL	change, postmenopausal		and reached significance at second WK of	

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		women or on contraception		medication, p=0.02. Global symptom relief	
	<u>Size</u> : n=171 pts	at 25 centers		score improved significantly.	
	(multiple system				
	atrophy, n=40 pts; PAF,	Exclusion: Pregnant or			
	n=37 pts, diabetic	lactating women,			
	neuropathy, n=37 pts,	preexisting sustained			
	Parkinsonism, n=19	supine HTN of ≥ 180/110			
	pts)	mmHg, concomitant			
		administration of			
		sympathomimetic agents,			
		adrenoreceptor alpha-			
		agonist or antagonists, or			
		vasoactive drugs, or			
		significant systemic illness			
Phillips AA, et al.	Aim: Assess effect of	Inclusion: SCI (n=10) and	Intervention: Midodrine 10 mg	1° endpoint: Tilt table (Progressively tilted	Only assessment of MCA and
2014	midodrine on OH and	age and sex matched able		from supine to 30, 45, and 60 degrees) and	PCA, without measurement of
24436297	cerebral blood flow in	bodied individuals (n=10)	Comparator: Baseline	symptoms. Stage and time at which	carotid, vertebral, or upstream
(300)	SCI compared to able-			participant withdrew or was withdrawn from tilt	arteries. No study of time of SCI
(000)	bodied	Exclusion: Smokers,	Then tilt table testing on 2	were recorded.	until assessment by tilt,
		history of CV disease	separate days	Word Todal dad.	and accomment by and,
	Study type: Analytical,			Steady state and dynamic cerebral blood	
	randomized controlled,			flow response to tilt is similar in SCI and AB;	
	prospective case-			midodrine improved orthostatic tolerance in	
	control			SCI by 59% (p=0.003) as calculated by	
	Control			orthostatic tolerance index calculated by the	
	Size: n=20 pts			formula orthostatic tolerance index calculated by the	
	<u> </u>			degree x time the last stage was tolerated.	
Ramirez CE, et	Aim: To assess	Inclusion: Pts with severe	Intervention: Atomoxetine 18 mg	<b>Primary:</b> Post-treatment upright SBP at 1 min.	Atomoxetine improved DBP
al.	whether atomoxetine	autonomic failure (PAF,	intervention. Atomoxetine to my	i innaiy. 1 Ost-deadinent upngnt ODF at 1 mm.	and symptoms greater than
2014	would be superior to	multiple systems atrophy,	Comparator 1: Midodrine 5–10	Secondary: Post-treatment seated SBP and	midodrine. Supine BP was not
25185131	midodrine in improving	Parkinson disease) with OH		DBP, upright DBp and heart rate, and OH	assessed. BP was measured
(301)	upright BP and OH	defined as SBP ≥20 mmHg	mg	Questionnaire and Q1 symptom scores.	
(301)	uprignt br and Ori	or DBP ≥10 mmHg within 3	Comparator 2: Placebo,	Atomoxetine improved upright SBP to a great	beyond 1 h after medication
	Study type: Analytical,	min of standing or 60	with SBP, DBP, and heart rate	extent than midodrine (means difference =7.5	administration
	randomized, single-	degree HUTT	assessed Q5 mins for 60 m	mmHg, p=0.03) and upright DBP (means	
	blind placebo	degree non	assessed Q3 1111115 101 00 111	difference =4.1mmHg, p=0.05). Atomoxetine	
	controlled, prospective	Exclusion: autonomic		improve OH related symptoms (p=0.02) but	
		failure secondary to DM,		not midodrine	
	crossover	Tallule Secondary to DM,		not midourine	

		amyloidosis, or			
	Size: n=65 pts	paraneoplastic syndrome			
Singer W, et al.	Aim: To assess	Inclusion: Adults >18 y of	Intervention: Pyridostigmine 60	Primary: Standing DBP at 1 h post drug:	Dividentiamina along and in
2006	pyridostigmine alone or	age with neurogenic OH	, <del></del> ,	pyridostigmine increased it from 49+/-14 to	Pyridostigmine alone and in combination with midodrine with
	in combination with		mg		
<u>16476804</u>		(multiple system atrophy,	Commence of a Divide of ancies	56+/-17 mmHg (p=0.02). Pyridostigmine with	resultant improvement in
(302)	midodrine in neurogenic	n=17; PAF, n=15;	Comparator 1: Pyridostigmine	midodrine 5 mg significantly increase standing	symptoms without significantly
	OH	autoimmune autonomic	60 mg + midodrine 2.5 mg	DBP compared to pyridostigmine + midodrine	affecting supine HTN.
	Charles have a Amelatical	neuropathy, n=9; diabetic	Commence of the Commission of	2.5 mg (p=0.03) and placebo (p=0.002) and	
	Study type: Analytical,	autonomic neuropathy,	Comparator 2: Pyridostigmine	almost significantly compared to	
	randomized, double-	n=11; or unspecified	60 mg + midodrine 5 mg	pyridostigmine alone (p=0.51)	
	blind, placebo	neurogenic OH, n=6). OH	Commente a 2 Discolor	Carandama Inflances on CDD and assista DD	
	controlled, prospective	defined as SBP drop ≥ 30	Comparator 3: Placebo	Secondary: Influence on SBP and supine BP:	
	crossover,	mmHg or mean BP drop ≥		no significant change, in SBP (p=0.36) or DBP	
	C:	20 mmHg within 3 min of		(p=0.85); relation of symptoms to change in	
	Size: n=58 pts	standing.		BP: significant association between change in	
		<b>-</b>		symptom score at 1 h to change in standing	
		Exclusion: Pregnant,		BP, p<0.001	
		lactating, evidence of failure			
		of other organ systems or			
		of systemic illness that			
		could affect autonomic			
		function, CHF, significant			
		CAD, significant arrhythmia,			
		renal disease, severe			
		anemia, hypothyroidism,			
		and cerebrovascular			
		accidents, concomitant			
		therapy with anticholinergic,			
		adrenergic antagonists,			
Mainlet DA et el	A: T d	vasoactive agents	Indonesia de Midadeira O.F.	40 1 14 15 05 5 5	- : HTN ::: 00
Wright RA, et al. 1998	Aim: To assess dose	Inclusion: >18 y of age,	Intervention 1: Midodrine 2.5 mg	1° endpoint: Midodrine 2.5 mg did not	Excessive HTN with 20 mg
	effect of midodrine in	neurogenic OH (≥15 mmHg	Intervention 2: Midadrina 10	significantly increase standing SBP at any	dose. Supine SBP >200 mmHg
9674789	neurogenic OH	SBP drop with standing;	Intervention 2: Midodrine 10 mg	time point.	occurred in 17% of pts on 10
(303)	Ctudy type, Applytical	PAF, n=14, and multiple	Intervention 2. Midadrina 20	Midodrine 10 mg increased standing SBP	mg, and in 41% of pts taking 20
	Study type: Analytical, randomized, double-	system atrophy n=7), and	Intervention 3: Midodrine 20 mg	significant 1 h post ingestion with a mean	mg.
	blind, placebo-	symptoms of OH, postmenopausal if a	Comparator: Placebo	increase of 34 mmHg, p<0.05.	. Midedwine at deepe of 10
	controlled, prospective	woman or taking	Comparator. Flacebo	Midodrine 20 mg increased standing SBP     increased standing SBP	Midodrine at doses of 10 mg     and 20 mg improves CRP with
	controlled, prospective	woman or taking		significantly at 1 to 4 h post ingestion with a	and 20 mg improves SBP with

	crossover,	contraception		mean increase of 43 mmHg, p<0.05.	standing in dose-dependent
	Size: n=25 pts	Exclusion: Pregnancy, lactating, supine hypertension ≥ 180/110 mmHg, concomitant administration of sympathomimetics or vasoactive drugs, significant systemic, cardiac, renal, or gastrointestinal illness, or clinically significant abnormalities on exam.		Significant improvement in symptoms occurred with 10 mg and 20 mg doses.	fashion with improvement in symptoms. With increasing dose, there is also increased frequency of supine HTN.
Biaggioni I, et al. 2015 25350981 (304)	Aim: To evaluate whether droxidopa is beneficial in treatment of neurogenic OH  Study type: Multinational, Analytical, Randomized placebo controlled, prospective cohort; parallel-groups phase 3 study  Size: n=101 pts	Inclusion: 18 y of age, symptomatic OH assoc with Parkinson disease, multiple system atrophy, PAF, dopamine beta-OHase deficiency, or non-diabetic autonomic neuropathy, with SBP decrease ≥ 20 mmHg or DBP decrease ≥ 10 mmHg within 3 mins standing  Exclusion: Severe HTN ≥ 180/110 mmHg; AF, or significant cardiac arrhythmia, current use of TCA, norepi reuptake inhibitors, current use of anti HTN meds, use of vasoconstrictive agents within 2 d.	Intervention: Droxidopa 100 mg TID and adjusted upward; mean dose at randomization was 389.6 +/- 180.9 mg 3x daily, then randomized to continue droxidopa  Comparator: After upward adjustment of droxidopa adjustment then withdraw to placebo for 14 days	Self Rated OH Questionnaire [6-item OHSA and 4-item OHDAS: Primary: pts change on OHSA item 1: dizziness/lightheadedness  Primary: OHSA item 1 increased by 1.3+/- 2.8 in droxidopa group vs. 1.9 +/-3.2 in placebo (p=0.509); Secondary: Favored droxidopa but not statistically  Secondary: Change in OHSA items 2–6: vision disturbance, weakness, fatigue, trouble concentrating, and head/neck discomfort).	During open label 58.6% reported ≥1AE, most commonly headache (11%); dizziness (8.3%); fatigue (5.5%); During double blind treatment, falls (2%), headache (4%), URI (4%), and dizziness (4%)      Unanticipated carryover effect of persistence of symptomatic improvement during withdrawal phase even in the placebo group. Secondary endpoints favor use of droxidopa in symptomatic neurogenic OH.
Freeman R, et al. 1999	Aim: To assess DL- DOPS in neurogenic	Inclusion: Autonomic failure pts with severe,	Intervention: 3-4-DL- threodihydroxyphenylserine (DL-	1° endpoint: DL-DOPS increased supine SBP (p<0.001), tilted SBP (p<0.05), supine	The norepi precursor DL- DOPS decreases BP fall with 60
10599797	OH,	symptomatic OH (n=6,	DOPS) 1000 mg	DBP (p<0.01) and tilted DBP (p<0.01) with	degree tilt orthostatic challenge.

(305)	Study type: Analytical, Randomized double- blind, placebo controlled crossover, prospective cohort, Size: n=10 pts	multiple system atrophy; n=4, PAF)  Exclusion: Alternative cause OH, systemic illness affecting autonomic function, significant CAD, cerebrovascular disease, or peripheral vascular disease, or malignant cardiac arrhythmias, pregnancy or child-bearing potential not on birth control, medication impairing vasomotor function except fludrocortisone	Comparator: Placebo then 60 degree tilt table	peak SBP occurring 300 m after medication ingestion. Plasma norepi increased in supine and tilt after DL-DOPS ingestion (p<0001). There was no significant effect on heart rate, forearm vascular resistance with DL-DOPS vs. placebo. Trend toward improvement in symptoms and quality of life of orthostatic intolerance seen with DL-DOPS (p<0.06)	
Hauser RA, et al. 2014 24326693 (306)	Aim: To assess droxidopa effect in neurogenic OH in Parksonson disease,  Study type: Multicenter analytical, randomized double-blind placebo controlled, prospective cohort phase 3 trial,  Size: n=51 pts	Inclusion: 51 pts with Parkinson disease enrolled in clinicaltrials.gov NCT01176240, droxidopa for neurogenic OH in Parkinson disease interim analysis;  Exclusion: N/A	Intervention: Droxidopa dosage optimization for ≤ 2 wk followed by 8 wk of maintenance therapy (100-600 mg 3x daily), mean study-drug dosage was 433 mg  Comparator: Placebo	Primary: Change in OH questionnaire composite score from baseline to wk 8  Secondary: OH questionnaire item 1 (dizziness, lightheadedness) and pts reported falls  Mean OH qiestionnaire composite score change at wk 8 was -2.2 vs21 (p=0.98).  Droxidopa group with 1.0 falls/wk vs. 1.9 falls/wk in placebo (p=0.16).	<ul> <li>17 droxidopa recipients (71%) with AE, nausea in 3 (13%), headache in 3 (13%), dizziness in 2 (8%)</li> <li>There was no benefit of droxidopa as measured by OHQ. There was a lower (insignificant) rate of falls with droxidopa, but this subgroup was too small to analyze benefit of droxidopa.</li> <li>98% of falls occurred in 22 pts (43%).</li> </ul>
Kaufmann H, et al. 2003 12885750 (307)	Aim: To assess L-DOPS effect on BP and orthostatic tolerance in severe neurogenic OH  Study type: Analytical, Randomized doubleblind placebo controlled	Inclusion: Severe symptomatic OH (n=11 with multiple system atrophy, n=8 with PAF  Exclusion: Sustained, severe HTN (>180/110 mmHg while sitting),	Intervention: L-threo-3,4-dihydroxyphenylserine (L-DOPS) with dose based on dose ranging study  Comparator: Placebo, then active standing	1° endpoint: L-DOPS significantly increased mean BP in supine (101+/-4 to 141 +/-5 mmHg) and standing (60 +/-4 to 100+/-6 mmHg, p<0.001)  • At 3 m of standing, 94% of pts were able to stand compared to 84% with placebo, p<0.001.  • L-DOPS showed increase in plasma NE	Supine HTN 45% vs. 23% in placebo, hyponatremia in 1 pts      L-DOPS improves BP and orthostatic tolerance in severe neurogenic OH, but the administration of carbidopa (which inhibits conversion of L-

	crossover, prospective cohort  Size: n=19 pts	clinically significant CAD, cerebrovascular disease, peripheral vascular disease, or cardiac arrhythmias		level that remained significantly elevated for at 46 H. Cardidopa abolished pressor response to L-DOPS.	DOPS to norepi peripherally) may limit L-DOPS effect in Parkinson disease pts
Kaufmann H, et al. 2014 24944260 (308)	Aim: To determine whether droxidopa improves neurogenic OH  Study type: Analytical, Randomized placebo controlled, prospective cohort; parallel-group trial of droxidopa responders,  Size: n=162 pts	Inclusion: Symptomatic neurogenic OH due to Parkinson disease, multiple system atrophy, PAF, or non-diabetic autonomic neuropathy  Exclusion: 95 titration failures (50 had treatment failure, 12 AEs, 6 withdrew consent, 4 protocol violations, 23 other failures, 6 randomized in error	Open-label droxidopa dose optimization (100 to 600 mg 3x daily) followed, in responders by 7 day washout and then  Intervention: 7 d double blind trial of droxidopa  Comparator: Placebo	1° endpoint: Responders to droxidopa defined as improvement on OHQ item 1 ≥ 1 unit, plus a ≥ 10 mmHg increase from baseline in standing SBP Primary: OHQ improvement from randomization to end of study • Secondary: changes in symptom and symptom-impact composite scores, and individual OHQ items • OHQ composite score improvement (1.83 vs. 0.93 units, p=0.003). Mean standing SBP increase of 11.2 vs. 3.9 mmHg, p <0.001)	<ul> <li>Headache (9.9%), dizziness (6.5%), nausea (4.6%), palpitations (1.9%)</li> <li>Only 1 w duration of therapy. No continuous BP monitoring</li> </ul>
Figueroa JJ, et al. 2015 25448247 (309)	Aim: Assess effect of abdominal compression on postural changes in SBP with OH,  Study type: Analytical, Randomized controlled, prospective crossover cohort  Size: n=13 pts	Inclusion: Moderately severe neurogenic OH, diagnosis of Parkinson disease, diabetic neuropathy, multiple system atrophy, autonomic failure, laboratory evidence of moderately severe adrenergic failure as measure by Valsalvainduced hypotension  OH defined as SBP ≥ 30 mmHg or DBP ≥ 15 mmHg  Exclusion: pregnancy, lactation, motor impairment affecting hand coordination, dementia, severe systemic illness, inability to tolerate	Moving from supine to standing  Comparator 1: Without abdominal compression;  Comparator 2: With abdominal binder in place;  Comparator 3: With maximal tolerable abdominal compression;  Comparator 4: With abdominal compression that pts believed would be tolerable for prolonged period	Primary: Postural changes in SBP. Mild abdominal compression (10 mmHg) prior to rising blunted drop in BP from -57 mmHg to -50 mmH (p=0.03) but other levels of compression did not have additional benefit.  Secondary: Pts assessment of preferences and ease of use. There was no difference in preference or ease of use.  Standing without binder: -57 mmHg (interquartile -40 to -76 mmHg). With 10 mmHg compression: -50 mmHg (interquartile range -33 to -70 mmHg, p=0.03)	Abdominal binders at minimal compression of 10 mmHg may blunt drop in BP. Additional compression did not have increasing effect unlike specialized shock garments which apply pressure over larger areas.

Di ii Oli di ii		withholding of anticholinergic-/alpha- and beta-adrenergic agonists for 5 half-lives prior to study, inability to withhold midodrine night before evaluation			
Platts SH, et al. 2009	Aim: To assess ability of 2 compression	Inclusion: n=19 healthy volunteers, 32–54 y of age,	(To mimic plasma volume loss due to spaceflight) pts given	<u>1º endpoint</u> : No significant difference in plasma volume loss between control (17.1%),	<ul> <li>Both the antigravity suit and Kentavr suits were able to</li> </ul>
<u>19456003</u> (310)	garments to prevent hypovolemia-related OI	and passing a modified Air Force Class III physical;	furosemide 0.5 mg/kg, consumed low-salt diet for 36 H	antigravity suit (16.9%), or Kentavr (18.4%).	resolve orthostatic intolerance during HUT, although the
(0.0)		and n=16 hypovolemic		Only 9 of 16 (56%) control pts were able to	Kentavr provided same benefit
	Study type: Analytical, randomized controlled,	control pts	Intervention: NASA antigravity suit inflatable in 25.9 mmHg	complete HUT. All antigravity suits (9 pts) and Kentavr (10 pts) were able to complete HUTT:	at approximately ½ of the compressive force.
	prospective cohort	Exclusion: none	increments, n=9	antigravity suit vs. control, p=0.03, Kentavr vs.	compressive force.
	Size: n=35 pts		Comparator: Russian Kentavr –	control, p=0.02. Change in SBP of control pts (-16 mmHg) was greater than antigravity suits	<ul> <li>Pts not exposed to all deconditioning effect of</li> </ul>
	<u>0126</u> . 11–35 pts		non-inflatable elastic shorts and	group (8 mmHg, p=0.005) and Kentavr group	microgravity, just acutely
			gaiters, n=10 then did 15 m 80 degree HUTT	(2 mmHg, p=0.035). No difference in diastolic BP.	reduced plasma volume
Podoleanu C, et	Aim: To assess lower	Inclusion: Pts with	Intervention: Leg compression	1° endpoint: Sham placebo leg bandage	Leg compression stocking is
al.	limb compression	symptoms signs of OI	bandages at 40 -60 mmHg for 10	and placebo abdominal bandage:	able to decrease the SBP drop
2006, 17010806	bandage effect on OH in elderly persons	(asymptomatic after standing in initial 3 m, but	m and then of the abdomen too (20 – 30 mmHg) for 10 m	SBP decreased from 125 +/- 18 mmHg to 112	with postural change, and
(311)	in elderly persons	cannot tolerate afterward	(20 – 30 Hilling) for 10 Hi	+/-25 mmHg with tilt for 10 m then to 106 +/- 25 mmHg after 20 m.	reduce symptoms over 1 mo follow-up
(5.1.)	Study type: Analytical,	due to increasing	Comparator: Sham compression,	With active bandage:	Tonon up
	randomized controlled	hypotensive symptoms,	then measured effect on 60	SBP was 129 +/-19 mmHg, then 127 +/-17	
	cross-over, prospective cohort	progressive decrease in BP pattern during diagnostic tilt	degree modified Italian HUTT	mmHg (p=0.03) at 10 m tilt, and then 127 +/- 21 mmHg, (p=0.02) at 20 min. Symptom	
	Conort	testing		burden vis SSS-OI questionnaire decreased	
	Size: n=21 pts			from 35.2 to 22.5 (p=0.01) after 1 mo of leg	
		Exclusion: Inability of pts to collaborate and to		compression stocking therapy.	
		perform tilt testing			
Protheroe CL, et	Aim: To assess effect	Inclusion: Healthy	HUTT and LBNP (-20 mmHg, -40	1° endpoint: Time to presyncope was not	There was no significant
al.	of graded calf	volunteers	mmHg, and -60 mmHg for 10 min	significantly different between compression	difference in time to presyncope
2011 22194814	compression stockings on orthostatic tolerance	Exclusion: CV or	each) on 3 occasions with different types of stocking:	stocking 26 +/- 2.0 m, calf placebo 29.9 +/- 1.8 m, and ankle placebo 27.6 +/- 2.4 m. Smaller	between compression stockings to placebo.
<u>ZZ 1340 14</u>	On orthostatic tolerance	EACIUSIOII. OV OI	unierent types of stocking.	in, and ankle placebo 27.0 +/- 2.4 m. Smaller	ιο μιασεύο.

(312)		neurological disease	Intervention: Calf-length graded	calf circumference may predict individuals who	
	Study type: Analytical,		compression stocking,	improve with compression stockings more	<ul> <li>Testing was performed in</li> </ul>
	randomized double-		Comparator 1: Standard calf-	than others.	healthy volunteers, and not pts
	blind placebo-controlled		length socks not designed to		with OI.
	crossover, prospective		provide compression (calf		
	cohort		placebo),		
			Comparator 1: Ankle-length		
	Size: n=15 pts		socks (ankle-placebo)		-
Clarke DA, et al.	Aim: Effect on isometric	Inclusion: Young pts	Intervention: Isometric	1° endpoint: With standing alone compared	<ul> <li>Maximum force isometric</li> </ul>
2010	handgrip on initial OH in	median age: 17 y of age	contraction of nondominant arm	to baseline, MAP decreased by 42+/-10%	handgrip before and during
<u>20350727</u>	young persons	range 15-22 y of age with	for 1 m then standing for 5 m	(p<0.01), heart rate increased by 62+/-18%	standing can blunt the decrease
(313)		initial OH (defined as	while maintaining isometric	(p<0.01), cardiac output decreased by 33+/-	in MAP and cardiac output in
	Study type: Analytical,	transient decrease in SBP	handgrip	17% (p<0.05), and TPR was unchanged at	younger pts with initial OH. No
	Randomized controlled,	>40 mmHg or a decrease in		17+/-21% (p=0.65).	formal evaluation of symptoms
	prospective cohort,	DBP >20 mmHg within 15 s	<u>Comparator</u> : Standing alone	On standing with isometric handgrip, MAP	performed. Less than maximal
	0. 44.1	of standing) with symptoms		decreased by 31+/-9% (p<0.01), heart rate	force handgrip not performed.
	Size: n=14 pts	F -1 -1 - 0 -1 -1		increased by 33+/-17% (p<0.01), cardiac	
		Exclusion: Systemic		output decreased by 2+/-14% (p<0.05), and	
		disease, vasovagal fainting,		TPR decreased by 30+/-15% (p<0.01)%.	
Vradiat CT at al	Aim. Acces les	chronic OI	Outh catatic talayanaa ahallayanad	40 and a late All along the late and a	1
Krediet CT, et al.	Aim: Assess leg	Inclusion: Healthy pts	Orthostatic tolerance challenged	1º endpoint: All pts sustained greater	Leg crossing increased
2006 16714361	crossing to increase orthostatic tolerance.	Exclusion: No medications	at same time	orthostatic challenge with leg crossing (34 +/-2	orthostatic tolerance in healthy
(314)	orthostatic tolerance,	except oral contraceptive.	Intervention, With los erossins	min), than during control (26 +/-2 min) or with	pts
(314)	Study type: Analytical,	No alcohol, tobacco, and	Intervention: With leg crossing	placebo (23+/-3 min, p<0.001). Heart rate	
	Randomized placebo	caffeine use.	Comparator 1: Without leg	increase was lower (+13 bpm) with leg	
	controlled crossover,	caneine use.	crossing	crossing during HUTT compared to control	
	cross-sectional cohort		Crossing	(+18 bpm, p<0.05)	
	CIOSS-SECTIONAL CONOR		Comparator 2: Placebo table		
	Size: n=9 pts		Comparator 2. Placebo table		
Thijs RD, et al.	Aim: To evaluate	Inclusion: Pts with	Intervention: Inspiratory	1° endpoint: IO increased MAP by 8 mmHg (-	Muscle tensing and inspiratory
2007	respiratory impedance	autonomic failure (PAF,	obstruction through narrowing of	1 to 13 mmHg), mean cerebral blood flow	impedance and muscle tensing
17679677	to reduce OH in	n=4; multiple system	inspiratory tube of 2 way	velocity (mCBFV) by 8% (2 to 23%).	had similar effects in increasing
(315)	autonomic failure	atrophy, n=3, amyloidosis,	nonrebreathing valve (IO)	1000kg (1110D1 1) by 070 (2 to 2070).	MAP and mean cerebral blood
(***)	/	n=1, anti-Hu neuropathy,		Muscle tensing increased MAP by 9 mmHg (1	flow velocity, but no difference in
	Study type: Analytical,	n=1, Parkinson disease,	Comparator 1: NS	to 10 mmHg), mCBFV by 9% (-7 to 18%).	symptom improvement was
	randomized controlled,	n=1) and symptomatic OH.			noted.
	prospective crossover	Healthy pts as control	Comparator 2: Muscle tensing of	Pursed lips during inspiration increased MAP	-

	<u></u>			<u></u>	
	<b>Size</b> : n=20 pts	(n=10)	legs without leg crossing	by 1 mmHg (-7 to 8 mmHg), mCBFV by 2% (-11 to 9%).	
		Exclusion: Cardiac disease or used antihypertensive medications	Comparator 3: Breathing through pursed lips during inspiration  Comparator 4: Inspiratory sniffing	No significant difference in symptom scores was noted between maneuvers	
Tutaj M, et al. 2006 <u>16096819</u> (316)	Aim: Assess effect of countermaneuveres in familial dysautonomia and active standing  Study type: Analytical, randomized controlled, prospective crossover  Size: n=17 pts	Inclusion: Familial dysatuonomia with IKBKAP gene mutation  Exclusion: Pts unable to comply with discontinuation of fludrocortisone or midodrine for 18 h.	Physical countermaneuvers Intervention: Leg crossing,  Comparator 1: Squatting,  Comparator 2: Bending forward with abdominal compression.  Medication affecting CV system (fludrocortisone, midodrine) held for 18 h prior to procedures	1° endpoint: 7 of 17 pts able to perform all 4 countermaneuvers. 16 of 17 pts able to perform at least 2 countermanuevers. SBP increase during bending forward (+23 mmHg, p=0.0005), squatting (+49 mmHg, p=0.002), leg crossing (+8.3 mmHg, p=0.01), abdominal compression (+27 mmHg), p=0.001). DBP increase during bending forward (+12 mmHg, p=0.0005), squatting (+38 mmHg, p=0.004), leg crossing (+11.6 mmHg, p=0.02) but no change during abdominal compression,	Squatting was most effective countermaneuver in increasing BP but only 7 of 17 pts with familial dysautonomia were able to perform it adequately. Other countermaneuvers increase BP to lesser degree, with leg crossing likely least effective
Singer W, et al. 2006 16476804 (302)	Aim: To assess pyridostigmine alone or in combination with midodrine in neurogenic OH  Study type: Analytical, randomized, double- blind, placebo controlled, prospective crossover,  Size: n=58 pts	Inclusion: Adults >18 y of age with neurogenic OH (multiple system atrophy, n=17; PAF, n=15; autoimmune autonomic neuropathy, n=9; diabetic autonomic neuropathy, n=11; or unspecified neurogenic OH, n=6). OH defined as SBP drop ≥ 30 mmHg or mean BP drop ≥ 20 mmHg within 3 m of standing.  Exclusion: Pregnant, lactating, evidence of failure	Intervention: Pyridostigmine 60 mg,  Comparator 1: pyridostigmine 60 mg + midodrine 2.5 mg,  Comparator 2: pyridostigmine 60 mg + midodrine 5 mg,  Comparator 3: Placebo	(+2.0 mmHg, p=0.30).  Primary: Standing DBP at 1 h post drug: pyridostigmine increased it from 49+/-14 to 56+/-17 mmHg (p=0.02). Pyridostigmine with midodrine 5 mg significantly increase standing DBP compared to pyridostigmine + midodrine 2.5 mg (p=0.03) and placebo (p=0.002) and almost significantly compared to pyridostigmine alone (p=0.51)  Secondary: Influence on SBP and supine BP: no significant change, in SBP (p=0.36) or DBP (p=0.85); relation of symptoms to change in BP: significant association between change in symptom score at 1 h to change in standing BP, p<0.001.	Pyridostigmine alone and in combination with midodrine with resultant improvement in symptoms without significantly affecting supine HTN.
		of other organ systems or of systemic illness that			

fi C	could affect autonomic unction, CHF, significant CAD, significant arrhythmia, enal disease, severe unemia, hypothyroidism, and cerebrovascular		
	ccidents, concomitant		
	herapy with anticholinergic,		
	drenergic antagonists, asoactive agents		

# Data Supplement 34. Nonrandomized Trials, Observational Studies, and/or Registries of Neurogenic Orthostatic Hypotension – (Section 6.1)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Jordan J, et al. 1999 <u>10073520</u> (317)	Study type: Analytical, observational, prospective case control,  Size: n=30 pts	Inclusion criteria: Severe OH due to autonomic failure (PAF, n=10; multiple system atrophy, n=9); healthy controls, n=11  Exclusion criteria: None	1º endpoint: 480 mL tap water  Results: In both autonomic failure and healthy controls, water ingestion raised SBP by 11 mmHg (p<0.001). No significant change in plasma volume was seen in healthy controls and 5 pts with autonomic failure. Norepi levels increased in controls with water ingestion.	Water ingestion increased BP in autonomic failure and healthy controls, possibly through sympathetic activation
Jordan J, et al. 2000 <u>10662747</u> (318)	Study type: Analytical, observational, prospective case control,  Size: n=66 pts	Inclusion criteria: primary autonomic failure with "disabling" OH. MSA, n=28; PAF, n=19. Healthy controls, n=19.  Exclusion criteria: Secondary causes of autonomic failure (DM, amyloidosis)	1º endpoint: 480 mL tap water  Vasoactive medications and fludrocortisone discontinued ≥5 half- lives before testing  Results: With water drinking, BP increased 33+/-5/16+/-3 mmHg (p<0.001) in MSA, and increased 37+/-7/14+/-3 mmHg in PAF (p<0.001). There was no difference between drinking cold vs. warm water. Drinking 480 mL had a greater pressor response than 240 mL water. Healthy controls also noted an increase in SBP of 11+/- 2.4 mmHg (p<0.001). Healthy controls undergoing ganglionic blockade did not have pressor effect with water. Enhanced pressor effect present with yohimbine plus water.	Water ingestion has a pressor response in autonomic failure, with BP increase also seen in healthy pts. The peak elevation in BP was 30 to 35 mins after ingestion. This effect is largely sympathetically driven.
Shannon JR, et al. 2002	Protocol 1: <u>Study type</u> : Analytical,	Inclusion criteria: 18 consecutive pts with primary autonomic failure	1° endpoint: Protocol 1:	<ul> <li>Rapid water ingestion of 480 mL at room temperature</li> </ul>

11904109 (319)	observational, prospective cohort Size: n=27 pts  Protocol 2: Study type: Analytical, observational, prospective cohort Size: n=27 pts	(multiple system atrophy n=9, and PAF n=9) with disabling OH, and n=9 pts with idiopathic orthostatic intolerance with 6 mo of symptoms,  Exclusion criteria: None	Intervention: 480 mL tapwater at room temperature in <5 min  Comparator: no tapwater  then active standing  Protocol 2: Intervention: eat a meal then 480 mL tapwater at room temperature  Comparator: no tapwater  then active standing  Results: Protocol 1: Seated BP increased from 117/67 mmHg before water drinking to 150/78 mmHg with water drinking (P<0.01).  After 1 min of standing, BP increased from 83/53 mmHg before water drinking to 114/66 mmHg with water drinking (p<0.01).  Maximal tolerated standing time increased from 5+/-3 min before water drinking to 11+/-10 min after drinking (p=0.06).  Protocol 2: Baseline BP was 138/77 mmHg, and with eating BP reached an average nadir of 95/57 mmHg. With water ingestion, BP increased to average peak 174/86 mmHg, and average nadir of 116/65 mmHg	improves orthostatic tolerance in pts with autonomic failure as well as post-prandial hypotension
Young TM, et al. 2004 15548493 (320)	Study type: Analytical, observational, prospective cohort  Size: n=14 pts	Inclusion criteria: chronic autonomic failure (7 pts with multiple system atrophy [MSA] which is preganglionic, and 7 pts with PAF which is postganglionic  Exclusion criteria: None	1º endpoint: 480 mL of distilled water at room temperature within 5 min then remained seated for 15 mins before standing for 5 min (Stand 1) then seated for 15 min then standing for 5 min again (Stand 2)  Results: Water ingestion raised SBP and DBP and lowered heart rate at 3 min and 5 min of Stand 1 compared to before water, all p<0.01.	Water ingestion increased standing BP and reduced symptoms due to OH. Increase in standing BP appeared related to increase in baseline BP after water ingestion. Pressor effect occurred sooner in PAF (within 5 mins) compared to MSA (13 mins)
Humm AM, et al. 2008 18469030	Study type: Analytical, randomized controlled crossover,	Inclusion: PAF with sympathetic and parasympathetic dysfunction with severe OH.	Water ingestion raised SBP and DBP and lowered heart rate at 3 min of Stand 2 compared to before water, all p<0.01, but at 5 min, only SBP and DBP had significance, p<0.01.  1° endpoint: 480 mL distilled room temperature water, then supine cycle ergometer followed by active standing	• N/A

(321)	prospective cohort		Results: Without water ingestion, with exercise there was SBP fall	
		Exclusion: None	(42.1+/-24.4 mmHg), DBP fall (25.9+/-10 mmHg).	
	Size: n=8 pts		With water ingestion, with exercise, SBP fall was still present	
			(49.8+/-18.9 mmHg), DBP fall (26.0+/-9.1 mmHg) but BP remained	
			higher after water intake although not quite significant (p=0.09).	
			Without water ingestion, 3 of 8 pts completed 5 min standing	
			protocol, whereas with water ingestion, 7 of 8 pts completed	
			protocol.	

### Data Supplement 35. Nonrandomized Trials, Observational Studies, and/or Registries of Neurogenic Orthostatic Hypotension – (Section 6.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Axelrod FB, et al. 1995 8690848 (322)	Aim: To assess midodrine effect in treating OH in familial dysautonomia,	Inclusion: Familial dysautonomia, OH  Exclusion: None	Intervention: Midodrine 2.5 3x daily titrated up  Comparator: No midodrine	Results: Average dose: 3.6 mg TID  All 9 pts had dizziness at baseline, and with midodrine 7 had improvement or resolution of dizziness. Mean increase in standing BP was not significant.	No placebo control, but most pts noted symptomatic improvement in this small open label study
	Study type: Analytical, observational, open label, prospective cohort  Size: n=9 pts	5 pts were on fludrocortisone which was continued			
Fouad-Tarazi FM, et al. 1995 7503082	Aim: To assess efficacy of midodrine with ephedrine,	Inclusion: autonomic insufficiency (idiopathic OH, n=7, multiple system atrophy, n=1), unable to tolerate other	Intervention: Midodrine (titrated from 2.5 to 10 mg 3x daily)	Results: Mean midodrine dose 8.4 mg 3x daily.  Mean ephedrine dose 22.3 mg 3x daily.  Midodrine and ephedrine both increased supine	Midodrine: supine HTN (n=1), scalp tingling (n=1) Midodrine was able to
(323)	Study type: Analytical, Randomized double- blind, placebo controlled crossover, prospective cohort	treatments because of physical disability, gastric irritation, fluid retention, or resistant hypokalemia	Comparator: ephedrine (titrated from 6 to 24 mg 3x daily) to where supine SBP between 140-180 mmHg, and supine DBP <100	BP vs. placebo (p<0.01 for both) not significantly different from each other. Ephedrine (vs. placebo) did not increase standing BP but did heart rate (p<0.05). Midodrine increased standing SBP and DBP vs.	significantly improve tolerance to standing with greater maintenance of SBP with standing compared to ephedrine and placebo
	Size: n=8 pts	Exclusion: recent history of persistent supine hypertension >180/100 mmHg unrelated to	mmHg and standing SBP≥ 80 mmHg	placebo (p<0.001) and vs. ephredine (p<0.001). Only midodrine produced a significant reduction in postural symptoms as	

	T	T		T	
		drug therapy, symptomatic CAD,		shown by increased ability to stand (5.3+/-4.4%	
		acute or chronic renal failure,		vs. 14.2+/-8.4%, p<0.01 vs. placebo) which	
		thyrotoxicosis, significant liver		correlated in increased percentage with	
		disease, pheochromocytoma,		standing SBP ≥ 80 mmHg	
		dementia, concomitant MAO			
		inhibitors			
Deng JC, et al.	Aim: Whether	Inclusion: Pts with neurogenic	Intervention/ Comparator:	Results: Order of efficacy in reducing	Compression of abdomen and
1997	compression of	OH (multiple system atrophy,	G suit with 5 separate	orthostatic symptoms from best to worst: All (13	legs, and even abdominal
9430805	different capacitance	PAF, or autonomic neuropathy)	compartments (lower	of 14, 93%) > abdomen (9 of 14, 64%) > calves	compression alone improves
(324)	beds can improve	, , , , , , , , , , , , , , , , , , , ,	abdominal, 2 thigh, and 2	+ thighs = calves alone > thighs.	orthostatic symptoms and
(0-1)	symptomatic	Exclusion: None	calf bladders).	angrie carres arene angrier	improves BP.
	neurogenic OH	<u>Excitation</u> . None	San Bladdoroj.	Maximal improvement in orthostatic BP	improved Br .
	Tiourogonio ori	OH defined as decrement in	Compartments were inflated	occurred with All (115.9+/-7.4 mmHg, p<0.005))	
	Study type: Analytical,	SBP $\geq$ 30 mmHg or mean BP $\geq$	to 40 mmHg as 1) bilateral	followed by Abdomen 102.0+/-6.7 mmHg,	
	observational,	20 mmHg	calves; 2) bilateral thighs, 3)	p<0.01) vs. noncompression (89.6+/-7.0	
	prospective cohort,	20 111111111111111111111111111111111111	combination of 1) and 2); 4)	mmHg). The other compartments compression	
	prospective conort,		low Abdomen; 5) All sites	results were not significantly different from	
	Cina, n=14 nto				
	Size: n=14 pts		combined; 6) baseline tilt (80	noncompression.	
			degrees for 5 min) without	lean and the second a	
Mathias Ol at	Aires Effect of L DODG	Includes 40 75 and and with	compression	Improvement correlated to increase in TPR.	La consensa la stata della della della consensa della
Mathias CJ, et	Aim: Effect of L-DOPS	Inclusion: 18–75 y of age with	Intervention: L-threo-DOPS	Results: L-DOPs blunted SBP decrease with	Increase lactate dehydrogenase
al. 2001	in management of	autonomic failure and symptoms	from 100 mg BID to 300 mg	standing (22+/-28 mmHg, p=0.0001) compared	(12.1%), urinary tract infection
<u>11710796</u>	neurogenic OH	(dizziness, syncope) and OH	BID	to baseline SBP. L-DOPs blunted DBP	(12.1%), akinesia (9.1%),
(325)		(drop in SBP ≥20 mmHg)		decrease with 2-min standing (8.1+/-17.2	headache (9.1%), and stomach
	Study type:			mmHg, p=0.0124) compared to baseline DBP.	upset (9.1%)
	Multicenter, analytical,	Exclusion: idiopathic		In 25 pts (78%), there was a decrease in OH.	
	observational, open-	Parkinson's disease, prior use or		In 14 ps (44%), OH was no longer observed by	L-DOPS reduces OH and
	label, prospective	current use of any		BP definition. Symptoms of light-headedness,	related symptoms in pts with
	cohort	antiparkinsonian drugs, mental		dizziness, and blurred vision improved	autonomic failure. No supine
		disorder, AF, serum creatinine		significantly from baseline with L-DOPS, but no	HTN was seen.
	<u>Size</u> : n=33 pts	>130 micromol/L, narcotic		correlation was found between change in	
		abuse, > moderate alcohol		postural SBP decrease and change in clinical	
		consumption (>1 L of beer or		symptom scores.	
		equivalent daily), child-bearing			
		potential, drug hypersensitivity			
Henry R, et al.	Aim: Effect of	Inclusion: elderly pts with	Intervention: Graduated	Results: Mean: 77.2 y of age (range 62-89 y	Graduated elastic compression
1999	compression hosiery in	reproducible, symptomatic OH	elastic compression hose	of age). Compression hosiery resolved	hose improves orthostatic
10406369	elderly persons with OH	(>20 mmHg)	,	symptoms of orthostatic dizziness in 7 of 10	tolerance and symptoms
		1 \ 2 \\	1	1 -7	

(326)	Study type: Analytical, observational, open label, prospective cohort  Size: n=10 pts	Exclusion: None	Comparator: baseline without compression hose then 90 degree HUTT	pts. Mean fall in SBP was 20.3+/-3.8 mmHg at baseline to 0.4 mmHg+/-8.2 mmHg with compression hose (p=0.005). Mean fall was significantly blunted with compression at HUTT mins 1, 2, and 3 (p<0.01, p<0.005, and p=0.01 respectively)	acutely. Long term studies are required.
Yamamoto N, et al. 2006 17003821 (327)	Aim: To assess abdominal compression with inflatable abdominal band in hemodialysis pts with OH  Study type: Analytical, observational, prospective cohort  Size: n=25 pts	Inclusion: Hemodialysis pts and OH for at least 6 mo before study enrolling between 7/2004 to 8/2004.  Exclusion: severe anemia (Hematocrit <25%), bleeding tendency, hypervolemic symptoms such as leg edema and pleural effusion, poor compliance, treatment for apparent infection, admission to hospital, chronic hypotension (defined as pre-dialysis SBP of <100 mmHg)	Intervention: Inflatable abdominal band then active standing test.  Intervention 2: Some pts received antihypotensive medications (L-threo-3,4-dihydroxyphenylserine [L-DOPS], n=5,  Intervention 3: midodrine, n=3	Results: Delta SBP was significantly less after hemodialysis with the abdominal band (-19.4 mm Hg) vs. without the abdominal band (-36.2 mm Hg, p<0.002). Supine SBP elevation was not seen with the abdominal band (149 vs. 153 mm Hg). Delta HR after hemodialysis was significantly greater with the band	Inflatable abdominal band was able to reduce post dialysis OH, in pts already receiving antihypotensive medications
Ten Harkel AD, et al. 1994 <u>7874844</u> (328)	Aim: Effect of leg muscle pumping and tensing on orthostatic pressure  Study type: Analytical, observational, cross-sectional cohort  Size: n=13 pts	Inclusion: normotensive pts (n=6); hypoadrenergic OH (OH, n=7) of which PAF comprised n=4.  Exclusion: None	Intervention: leg crossing  Comparator: no leg crossing	Results: Leg crossing resulted in increase in BP (13+/-2 mmHg vs. 9+/-7 mmHg), and cardiac output (49+/-13% vs. 38+/-15%) in normal pts vs. pts respectively. Pts with PAF and non-PAF noted increase in BP and cardiac output.	Leg crossing increases BP and cardiac output in both normal and hypoadrenergic OH.
Van Lieshout, et al. 1992 1348300 (329)	Aim: Whether physical maneuvers can improve orthostatic tolerance in autonomic failure  Study type: Analytical,	Inclusion: autonomic dysfunction (hypoadrenergic) with OH, n=7; healthy pts, n=6  Exclusion: None	Comparator: Standing upright until presyncopal, followed by  Intervention 1: leg-crossing and then standing upright	Results: In autonomic dysfunction group, 5 of 7 pts had orthostatic dizziness within 10 min of standing. (BP 139/75 mg supine decreasing to 75/50 mmHg upright, MAP 58 mmHg). Leg crossing improved SBP to 95/60 mmHg with MAP 72 mmHg. With recurrence of	Both leg crossing and squatting improved symptoms of orthostatic intolerance and improved BP, with squatting having larger effect.

	observational,		until prosynoonal	presyncope, BP was 74/47 mmHg with MAP 56	
			until presyncopal		
	prospective cohort,		Into months of Or followed by	mmHg. Squatting increased BP to 131/81	
	<b>6</b> :		Intervention 2: followed by	mmHg (MAP 100 mmHg). Symptoms improved	
	Size: n=13 pts		squatting and then standing	with both maneuvers.	
			upright until presyncopal		
				In healthy, there was much milder increase with	
				leg-crossing (+4/0 mmHg) and with squatting	
				(+12/4 mmHg).	
Singer W, et al.	Aim: To assess	Inclusion: at least 18 y of age	Intervention: Pyridostigmine	<b><u>Primary:</u></b> Heart rate: 1 h after pyridostigmine,	Acetylcholinesterase inhibition
2006	acetylcholinesterase	old with orthostatic intolerance	60 mg	heart rate was significantly lower in both supine	may enhance sympathetic
<u>17016160</u>	inhibition in orthostatic			(73.0 vs. 78.9 bpm) and upright position (110.6	ganglionic transmission and
(330)	intolerance during	Exclusion: Pregnancy or	Comparator: No	vs. 123.7 bpm, p<0.001)	improves orthostatic intolerance
, ,	HUTT	lactating, failure of other organ	pyridostigmine	. , ,	
		systems or of systemic illness	' '	Secondary: Other CV parameters: no	
	Study type: Analytical,	that could affect study results,	Then 70 degree HUTT for 5	significant difference in SBP, DBP, MAP, SV,	
	observational open-	autonomic function or pts ability	mins	cardiac index; Influence on baroreflex	
	label, prospective	to cooperate (CHF, significant		sensitivity (BRS): significantly higher after	
	cohort.	CAD, significant arrhythmia,		pyridostigmine (p<0.005);	
	Conort,	renal disease, severe anemia,		Influence on plasma catecholamines: plasma	
	<b>Size</b> : n=18 pts	hypothyroidism, and		norepi significantly higher 1 h after	
	<u>0120</u> : 11–10 pts	cerebrovascular accidents),		pyridostigmine for supine (p=0.03) and upright	
		therapy with anticholinergic,		(p=0.005) positions.	
				(p=0.003) positions.	
		adrenergic antagonists,		Lieut vete bijusting and increased places	
		vasoactive agents, or		Heart rate blunting and increased plasma	
		medications that could interfere		catecholamine levels were associated with	
		with autonomic function unless		significant amelioration of orthostatic symptoms	
		discontinued for 5 half-lives		(p=0.01)	
		before study			

# Data Supplement 36. RCTs Involving Dehydration and Drugs – (Section 6.2)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Anley C, et al.	Aim: To assess which	Inclusion: All	Intervention: OT, oral fluid and	1° endpoint: Time to discharge from the	With no difference in time to

2011 20584756	treatment protocol for exercise-associated	collapsed athletes at two Ironman Triathlon	Trendelenburg position	medical tent (in min)	discharge, but significantly less fluid given in OT group
(293)	postural hypotension results in earlier discharge	competitions and one ultra-distance footrace in 2006 and 2007	Comparator: IV	Results: No significant difference between IV (52.5 +/- 18 min) and OT group (58+/-23 min), p=0.47	compared to IV group, the probable cause of exercise associated postural hypotension is peripheral
	Study type: Analytical, randomized, prospective cohort	Exclusion: Abnormal serum sodium		Secondary endpoint: Heart rate and BP changes:	vasodilatation resulting in venous pooling
	<u>Size</u> : n=28 pts			Results: No significant changes were seen.  Total volume of fluid in OT group was 204 +/- 149 ml, and was significantly less than IV group 1045+/-185 ml, p<0.001.	
Atherly-John YC, et al. 2002 12444837 (331)	Aim: To compare oral rehydration therapy with IV therapy for moderate dehydration in children  Study type: Analytical, randomized, prospective cohort  Size: n=34 pts	Inclusion: Children with moderate dehydration (having at least 4 standard published criteria) at single center  Exclusion: Chronic illness, severe dehydration or shock, protracted vomiting, absent bowel sounds, no accompanying guardian, no contact telephone number, and those requiring IV access for reasons other than hydration	Intervention: Oral replacement therapy: 5 mL every 5 min if <4 y of age, 10 mL every 5 mins if ≥4 y of age, and intake was advanced to twice the initial volume if there was no vomiting during the first H; n=18  Comparator: IV therapy (initial bolus of 20 mL/kg of isotonic sodium chloride over 30 min period, and second bolus was given per treating physician discretion. This was followed by IV solution of 5% dextrose in 0.45% or 0.33% saline depending on age at a rate of 1.5 times daily maintenance; n=16	1º endpoint: Duration of pediatric emergency department stay: Results: Oral replacement therapy: 224.7 +/-77.8 min vs. IV 358 +/-160 min, p<0.01  Secondary endpoints: Staff time require for pts care: Results: ORT: 35.8+/-32 min vs. IV: 65+/-44 min, p=0.03  Parent satisfaction: Results: ORT: 77.7% vs. IV: 37.5%, p=0.01  Hospital admission rate: Results: ORT: 11.1% vs. IV: 25%, p=0.2  Relapse after being discharged:	Oral rehydration therapy shortens emergency department stay, reduces staff time required for pts care, and improves satisfaction with pts care compared to intravenous rehydration for pediatric pts presenting with moderate dehydration.
Keneflick RW, et al. 2006 17146319 (332)	Aim: To determine effects of rapid (<30 min) IV vs oral rehydration immediately after dehydration during subsequent exercise in	Inclusion: Healthy non heat acclimated men  Exclusion: N/A	Each subject performed 3 trials:  1) Dehydration phase, pts walked or ran for 75 min at 50% VO <sub>2</sub> max with airflow directed to enhance evaporative sweat loss	Results: 0% in both ORT and IV groups  1° endpoint: To determine effects of rapid (<30 min) IV vs. oral rehydration immediately after dehydration, on CV, thermoregulatory, and perceptual responses during subsequent exercise	Although IV hydration restored plasma volume more quickly than oral rehydration, there was no significant effect on exercise duration. Sensation of thirst was

	the heat  Study type: Analytical,		2) Rehydration phase Rehydration treatments were randomly assigned to receive amount of fluid lost during	Results:  IV rehydration resulted in more rapid plasma volume restoration (p<0.05)	improved with oral rehydration.
	randomized, prospective cohort  Size: n=8 pts		dehydration: Intervention 1: IV rehydration (0.45% saline) Intervention 2: Oral rehydration (0.45% saline) Intervention 3: No fluid	However, there was no significant improvement in exercise duration (IV: 72.6+/-28.9 min; oral: 70.6+/-8.2 min) during the heat tolerance testing with IV vs. oral rehydration.	
			Then: 3) heat-tolerance test: immediately after 30 min rehydration period, pts performed a 75 min heat tolerance test in 37°C chamber	Sensation of thirst was significantly lower in oral rehydration than IV fluid (p<0.05)	
Maughan RJ, et al. 1995 <u>8549573</u> (333)	Aim: To study the effect of sodium content of drinks on rehydration after exercise  Study type: Analytical, randomized, prospective cohort  Size: n=6 pts	Inclusion: Healthy males  Exclusion: N/A	Pts were dehydrated by intermittent cycle exercise in warm and humid environment then ingested 1.5 times body mass loss of:  Intervention 1:  Na content 2 mmol/L (108 mosmol/kg) Intervention 2:  Na content 26 mmol/L (158 mosmol/kg) Intervention 3:  Na content 52 mmol/L (206 mosmol/kg) Intervention 4:  Na content 100 mmol/L (300 mosmol/kg)	1º endpoint: Effect of sodium content of drinks on rehydration after exercise  Results: Net fluid balance at end of trial: Sodium content 2 mmol/L: -689 mL Sodium content 26 mmol/L: -359 mL Sodium content 52 mmol/L: 2 mL Sodium content 100 mmol/L: 98 mL  Plasma volume was higher with sodium contents of 52 and 100 mmol/L compared to 2 mmol/L  Cumulative urine output was higher on sodium content 2 mmol/L than with 52 mmol or 100 mmol/L.	Rehydration and retained volume is greater with ingestion of fluid with increasing sodium concentration
Merson SJ, et al. 2008 <u>18463891</u> (334)	Aim: To investigate differing sodium chloride concentrations affect rehydration  Study type: Analytical, randomized, prospective	Inclusion: Healthy men without Hx of CV or renal disease Exclusion: N/A	Exercise via cycle ergometer with measured VO <sub>2</sub> max then drinking 150% of fluid lost as sweat:  Intervention 1: NaCl 0 mmol Intervention 2: NaCL 30 mmol/L Intervention 3: 40 mmol/L Intervention 4: 50 mmol/L	1° endpoint: Sodium chloride concentration effect on rehydration after exercise and subsequent exercise capacity      Results:     • Pts retained more of test drink as the sodium concentration of the drink increased	Increased sodium content of the test drink improved hydration compared to lower sodium and no sodium test drinks. Higher sodium drinks did not affect repeat exercise performance.

	cohort  Size: n= 8 pts		Then exercised again to 95% of VO <sub>2</sub> peak or exhaustion	(as measured by corresponding decreasing urine output). ◆ Significantly more fluid was retained on 40 and 50 mmol/L NaCl compared to 0 mmol/L (p<0.01).  ◆ Greater net negative fluid balance was seen 4 h after finishing drinking with lower sodium concentration test drink.  ◆ There was no effect of the sodium content of the drink on time to exhaustion on repeat exercise (p>0.8)	
El- Sayed H, et al. 1996 8673750 (232)	Aim: To evaluated salt supplementation in syncope with OI  Study type: Analytical, Randomized placebo controlled, prospective cohort, Size: n=20 pts  Study type: Analytical, observational, open label, prospective cohort Size: n=11 pts	Inclusion: Recurrent syncope without etiology  Exclusion: N/A	RDBPCT: Intervention: sodium chloride 10 mmol  Comparator: Placebo 12x daily then 60 degree HUTT with LBNP up to -40 mmHg  Open label: Intervention: slow sodium 10 mmol 12x daily (pts told it was a "mineral dietary supplement") then 60 degree HUTT with LBNP up to -40 mmHg	1° endpoint: Effect of salt administration on plasma volume and orthostatic tolerance in pts with posturally related syncope  Results: RDBPCT: 8 of 10 pts taking salt, vs. 3 of 10 taking placebo showed significant increases in plasma and blood volumes (p<0.05); all pts with increased plasma and blood volumes showed improved tolerance to orthostatic stress (time to presyncope)  Open label: 7 of 11 taking salt had increased plasma and blood volumes, and these pts showed improved symptoms of orthostatic tolerance	Pts with salt supplementation (increasing plasma volume by >90 mL) had significant increase in orthostatic tolerance. Pts with signs of high salt intake at baseline (by 24 h urinary sodium excretion) did not benefit from additional salt loading

## Data Supplement 37. Nonrandomized Trials, Observational Studies, and/or Registries of Dehydration and Drugs – (Section 6.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Greenlead JE, et al.	Aim: To evaluate various	Inclusion: Healthy young	Pts dehydrated for 24 h with moderate dehydration confirmed by	<ul> <li>Sodium content appears to be</li> </ul>
1998	carbohydrate electrolyte fluid	men, nonsmokers, no drug	plasma osmolality (298-305 mOsm/kg) then drank 1 of 6 fluid	more important than total osmotic
<u>9737753</u>	formulations for consumption	use	formulations (12 mL/kg: 898-927 mL):	content for inducing hypervolemia.
(335)	by astronauts to restore plasma		Intervention 1: water	

	1			
	volume	Exclusion: N/A	Intervention 2: 19.6 mEq/L Na	
			Intervention 3: 157 mEq/L Na	
	Study type: Analytical,		Intervention 4: 19.6 mEq/L Na + glucose	
	observational, prospective		Intervention 5: Performance® ~20 mEq Na	
	cohort		Intervention 6: Power Surge® ~20 mEq Na	
	Size: n=7 pts		1° endpoint: Plasma volume and total body water	
			Results: At rest, drinking formulations with higher sodium had greater increases in plasma volume. 157 Na resulted in 7.6% increase in plasma volume. Lower sodium content beverages but with higher total osmolality did not hydrate as well.  At rest, drinking 157 Na (the largest Na content), induced the greatest hypervolemia: 7.6%, p<0.05. water ingestion did not increase plasma volume.	
			With exercise, high sodium intake beverages were no more	
			effective than low sodium beverages for plasma volume stabilization. However, water was the least effective with an initial loss (17%) of plasma volume within the first 9 min of exercise.	
Shirreffs SM, et al. 1996 <u>8897383</u> (336)	Aim: To study the interaction between volume and composition of fluids ingested for rehydration effectiveness  Study type: Analytical, observational, prospective cohort  Size: n=12 pts	Inclusion: Healthy men Exclusion: N/A	Each subject exercised to induce sweat loss of 2% of body mass then drank beverages with different sodium concentration and volumes:  Sodium concentration: Intervention 1: low sodium (23 mmol/L) Or Intervention 2: high sodium (61 mmol/L)  Both drinks also contained small amounts of potassium and glucose (90 mmol/L).	Drinking a large volume beverage may be inadequate to rehydrate if the sodium concentration is insufficient, and drinking a high-sodium concentration beverage may be inadequate if a large enough volume is not consumed.
			Volume:  Intervention A: 50% of body mass loss Intervention B: 150% of body mass loss Intervention C: 150% of body mass loss Intervention D: 200% of body mass loss	

2009 increasing carbohydrate and sodium content on fluid delivery (337)  Exclusion: N/A  Carbohydrate group (CHO, n=10 pts) Intervention 1: G0: water + 20 mmol/L sodium Intervention 2: G3: 3% glucose + 20 mmol/L sodium above 3% did not further incre fluid delivery. Sodium content not significantly affect fluid del although there was a trend for	2009 19232115	increasing carbohydrate and sodium content on fluid delivery  Study type: Analytical, observational, prospective case control,	,	<ul> <li>mL, C=602 mL, D=1001 mL</li> <li>Pts rehydrating with low sodium beverage were in a more negative state of fluid balance with Intervention A (-909 mL) than Intervention C (-128 mL) or D (-135 mL)</li> <li>Pts rehydrating with high sodium beverage were in a more negative state of fluid balance with Intervention A (-958 mL) than Intervention D (+427 mL).</li> <li>Each subject undertook 4 trials each &gt;7 days apart</li> <li>Carbohydrate group (CHO, n=10 pts)</li></ul>	Increasing the glucose content above 3% did not further increas fluid delivery. Sodium content di not significantly affect fluid delive although there was a trend for reaching plateau time more quick with higher sodium content.
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Beckett NS, et al. 1999 10618673 (338)	Aim: To assess OH prevalence and associated factors in elderly hypertensive pts,  Study type: Analytical, observational, cross-sectional study	Inclusion: Pts in HYVET trial (Hypertension in the Very Elderly Trial); at least 80 y of age with sustained systolic (average SBP 160-219 mmHg) and diastolic hypertension (average DBP	glucose.  • Sodium group: trend for decrease in time to plateau with increasing sodium content (Na0: 23 min, Na20:19 min, Na40:18 min, Na60:16 min)  • Plasma deuterium enrichment did not differ between groups (p=0.121)  1° endpoint: Orthostatic fall in BP in hypertensive pts  Results:  Mean sitting BP was 182/100 mmHg.  Average fall in SBP on standing was 8 mmHg (95% CI: 7.3–8.3) and in DBP was 1.3 mmHg (95% CI: 1.0–1.6). 96 (7.7%) had a drop of ≥20 mmHg systolic and 66 (5.4%) had a drop of ≥10	Prevalence of OH in elderly pts with hypertension was 12%
Blake AJ, et al. 1988,	Size: n=1,241 pts  Aim: To assess falls and their associated causes	90-109 mmHg)  Exclusion: Pts on BP lowering treatment for reasons other than HTN  Inclusion: Community survey (Activity and Ageing	mmHg diastolic  1° endpoint: Prevalence of and factors associated with falls in the elderly	Decreasing handgrip strength, arthritis, and foot difficulties were
<u>3266440</u> (339)	Study type: Descriptive cross sectional survey  Size: n=356 pts	survey conducted between 5/1985 and 9/1985 of individuals age ≥65 y of age who reported ≥ 1 fall in preceding y  Exclusion: Mental incompetence, dementia, acute organic brain syndrome	Results: Women were more likely to report falls than men (p<0.001). Older respondents were more likely to report falls (p<0.05). Increasing number of prescribed drugs correlated increased prevalence of falls (p<0.001). There was no significant difference in antihypertensives (p=NS) or diuretics (p=NS). Hypnotics (p<0.05) and antidepressants (p<0.01) were more associated falls	strongest predictors of falls. Hypnotics and antidepressants (tricyclic antidepressants) were the medication classes associated with falls.
Burke V, et al. 1992 <u>1484937</u> (340)	Aim: To assess relation of drug treatment to postural fall in BP in elderly,  Study type: Descriptive, cross-	Inclusion: Independent elderly volunteers (pts >60 y of age) in Perth, Australia;  Exclusion: N/A	1° endpoint: Factors associated with postural fall in SBP  Results: Postural fall in SBP was related to alcohol intake >20 mL/day, sleeping tablet use, higher anxiety level, and lower body mass	There was no relation of anti- hypertensive medication to postural fall, but sleeping aid use was associated.
	sectional survey  Size: n=843 pts	EXCIMINATION IN THE PROPERTY OF THE PROPERTY O	index.  Postural fall in SBP was not related to HTN, age, gender, diabetes, or cardiac medications [verapamil (p=0.092), BB (p=0.728),	• Postural fall in SBP defined as ≥ 20 mmHg decrease when changing from sitting to standing

			diuretics (p=0.356), or vasodilators (p=0.199)].	
Craig GM, et al. 1994 <u>7971628</u> (341)	Aim: Presentation of OH in elderly  Study type: Descriptive retrospective chart review  Size: n=50 pts	Inclusion: Elderly pts with OH (defined as ≥ 20 mmHg fall in SBP)  Exclusion: N/A	1° endpoint: Factors associated with orthostatic fall in SBP ≥ 20 mmHg  Results: Presenting features of OH: Falls 64%, poor mobility 44%, unsteadiness 38%, confusion 22%. Medication usage in OH pts: Diuretic 56%, benzodiazepine 26%, anti-depressant 24%, anti-parkinsonian therapy 22%, phenothiazine 18%, BB 12%, hydralazine 10%, calcium antagonist 8%, nitrates 6%.	Medication was primarily responsible for OH in 66%, and implicated in 80% of cases.
Fotherby MD, et al. 1994 7870633 (342)	Aim: Assess prevalence of OH in elderly HTN pts whether anti-HTN therapy was continued or not,  Study type: Analytical, observational, prospective cohort  Size: n=47 pts	Inclusion: Pts ≥ 65 y of age, BP <175/100 mmHg on pharmacological treatment >1.  Exclusion: MI or stroke within preceding 6 mo, having angina or known major illness, diabetes, Parkinson disease, or on medication other than antihypertensives known to affect BP  Following treatment withdrawal, pts whose SBP was ≥175 mmHg and/or whose DBP >100 mmHg on 2 occasions were withdrawn from the study and deemed unsuitable for anti-HTN withdrawal	1º endpoint: Prevalence of OH  Intervention: Anti-HTN medication withdrawal, BP measured at 1, 3, 6, 9, and 12 mo of anti-HTN therapy withdrawal  Comparator: Continuing on anti-HTN had BP measure at 6 and 12 mo.  Results: For pts stopping anti-HTN medication, the number of OH fell from 11 (23%) on anti-HTN treatment to 4 (11%, p<0.05) off treatment.  • The pts continuing anti-HTN medication showed no significant change in prevalence of OH, 5 (38% at baseline, and 4 (31%) at 12 mo.  • Pts with OH on treatment (vs. those with OH on treatment) were older (79 y of age vs. 74 y of age, p=0.05) and had higher prewithdrawal SBP (164+/-21 vs. 147 +/-17 mmHg, p=0.02)	Withdrawal of anti-HTN therapy can decrease OH occurrence. Those with OH on anti-HTN treatment tended to be older and had higher prewithdrawal SBP     13 of the 47 pts did not meet criteria for anti-hypertensive withdrawal     OH defined as mean SBP fall ≥ 20 mmHg on standing from supine
Jansen RW, et al. 1996 <u>8636581</u> (343)	Aim: To assess post-prandial hypotension and relation to chronic use of CV medications  Study type: Analytical, observational, prospective	Inclusion: Nursing home residents, sinus rhythm, be able to stand from supine position within 30 s and remain standing for 10 min	<ul> <li>Comparator: Standing test, then</li> <li>Intervention: repeat standing test after eating meal.</li> <li>Same protocol repeated in 3 to 14 days.</li> <li>1° endpoint: BP and heart rate before and after postural change;</li> </ul>	Post-prandial responses in BP and heart rate are similar, and CV medication administration did not affect post-meal findings. However, the CV medication did affect BP after standing suggesting this

	T	r <del></del>		
	cohort	Exclusion: Presence of	BP and heart rate before and after meals	response may be distinct from
		pacemaker, insulin-		postprandial hypotension
	Size: n=22 pts	dependent DM	Results:	<ul> <li>Post-prandial hypotension defined</li> </ul>
			Mean SBP, mean DBP, and MAP all declined 45 min after the meal	as SBP decline of ≥20 mmHg within
			(p<0.001 for each).	90 min study period
			<ul> <li>Mean SBP declined 16+/-4 mmHg (p&lt;0.001) at 45 min and by</li> </ul>	OH defined as SBP decline ≥
			12+/-4 mmHg (p<0.01) during second test with no difference	during first and/or third min after
			between the 2 tests. MAP similarly declined in each test after	standing
			means (p<0.001).	
			Postprandial hypotension occurred in 10 pts in first test and 1	
			additional pts in second test. Administration of CV medications did	
			not affect significantly subsequent BP response after meals but did	
			affect SBP after standing.	
Jodaitis L, et al.	Aim: Association of OH with	Inclusion: Older (pts ≥75 y	1° endpoint: Prevalence of OH	There was no association of any
2015	use of drugs with psychotropic,	of age) in pts screened for		medication with OH or falls, but
26135806	CV, or diuretic effect	OH (defined as reduction of	Results:	many pts in this study had frailty
(344)		≥20 mmHg in SBP or ≥10	Mean age was 85+/-5 y of age in pts with OH, and 84+/-4 y of age	which could affect response to
	Study type: Prospective	mmHg in DBP within 3 min	without OH. Prevalence of OH was 41% (30% for SBP, 23% for	medication
	observational, multicenter,	of standing	DBP). Pts with OH vs. without OH were more likely to have falls	
		, and the second	(62% vs. 40%, p<0.001) and syncope (29% vs. 4%, p<0.001).	
	Size: n=285 pts	Exclusion: N/A	There was no difference in proportions of pts receiving drugs or	
	· ·		drug potentially associated with falls and/or OH.	
Kamaruzzaman S, et	Aim: Association of OH and	Inclusion: British Women's	1° endpoint: Prevalence of OH	OH was associated with
al.	medication use in British	Heart and Healthy Study		increasing age, HTN, and death.
2010	Women's Heart and Health	cohort, OH (defined as SBP	Results:	Use of BB and use of 3 or more
<u>19897539</u>	Study	≥ 20 mmHg and/or diastolic	Higher prevalence of OH in women with HTN than without HTN	antihypertensive medications were
(345)	Cross-sectional analysis	$BP \ge 10 \text{ mmHg}$ ).	(79% vs. 64%, p<0.001). No association of OH to coronary heart	associated with OH. Polypharmacy
		<b>G</b> ,	disease, diabetes, COPD, or cancer.	in of itself was not associated with
	Study type: Retrospective,	Exclusion: N/A	<ul> <li>Prevalence of OH was 28% (95% CI: 26.6–29.4) among women</li> </ul>	OH.
	observational cross-sectional		60-80 y of age. Among BP lowering medication, only BB had	
	cohort		higher odds of OH (OR: 1.26, 95% CI: 1.09–1.47, p<0.01). Women	
			on multiple antihypertensive drugs (≥ 3 vs. 0) had increased odds	
	<u>Size</u> : n=3,775 pts		of OH (OR: 1.99, 95% CI: 1.30–3.05, p=0.003). OH was	
			associated with all-cause mortality (OR: 1.10, 95% CI:1.07–1.14,	
			p<0.001)	
McLachlan CY, et al.	Aim: To assess frequency,	Inclusion: All admissions	1° endpoint: Prevalence of ADE	ADE comprises a significant
2014	nature, and causality of ADE	at single center in New	· <del>-</del>	amount of admissions at this single
<u>24750276</u>	resulting in acute admissions	Zealand between 10/1/2011	Results:	center, with the syncope being the

(0.40)		1. 44/44/0044	00000 - 1.1.1.1	
(346)	Study type: Analytical, observational, prospective	to 11/11/2011 and 12/24/2011 to 4/4/2012.	Of 336 admissions, 96 (28.6%) were related to ADE. 65 (19.3%) were caused by ADE, and 31 (9.2%) were contributed to by an ADE.	most frequent effect. Vasodilators and diuretics comprise 39% of ADE-related admissions
	cohort	Exclusion: N/A	Most common adverse effects were postural hypotension and/or vasovagal syncope (29%)	
	<b>Size</b> : n=96 pts		Most common implicated medications were vasodilators (23%), psychotropic medications (18%), and diuretics (16%), chronotropic medications [amiodarone, BB, diltiazem, digoxin] (11%)	
Ooi WL, et al. 1997 <u>9109468</u> (347)	Aim: To assess for clinical correlates for orthostatic BP change,  Study type: Analytical, prospective observational cohort	Inclusion: Nursing home residents ≥ 60 y of age, life expectancy >3 mo, able to stand at least 1 min  Exclusion: N/A	1° endpoint: supine BP, 1-min standing BP, 3-min standing BP, and heart rate  Results: After multivariate analysis, significantly associated (p<0.05) with OH were: elevated supine BP before breakfast, lightheadedness with standing, male gender, Parkinson disease medications, lower	Antihypertensive medication use was not associated with OH, but lower body mass index and Parkinson disease medications were.
	<u>Size</u> : n=911 pts		body mass index. Diuretic, antianginal, antiarrhythmics, and ACE-inhibitors were not associated with OH.	
Panayiotou B, et al. 2002 11824858 (348)	Aim: To assess antihypertensive medications in acute stroke for OH  Study type: Analytical, prospective, observational cohort.  Size: n=80 pts	Inclusion: Pts ≥ 65 y of age, mild or moderate ischemic stroke, admitted to hospital ≤24 h of stroke onset, living at home, could be on antihypertensive medication ("treated group", n=40) or not ("untreated group", n=40)  Exclusion: Hemorrhagic stroke, comorbidity affecting BP regulation (DM or Parkinson disease), know postural hypotension, MI in previous 3 mo, severe HF (NYHA III or IV), AF, urea >10 mmol/L, hemoglobin <10 g/dL, antibiotic requirement, serious illness.	1º endpoint: BP and heart rate measurements while supine, sitting, and standing within 3 d of stroke onset ("day 1"), and again 4 to 7 days ("wk 1") after stroke onset  Results:  Between d 1 and wk 1, supine BP fell significantly in treated group (165 +/-24/87+/-14 mmHg to 155 +/-24/83+/-14 mmHg, p=0.003 for SBP and p=0.03 for diastolic BP, but no significant difference in untreated group. On day 1, OH was observed within 5 min in 11 treated and 5 untreated pts, p=0.09. At wk 1, OH occurred in 5 treated and 8 untreated pts, p=0.36. Only cardiac dysfunsion was associated with OH on multivariate analysis (OR: 3.5, 95% CI: 1.0–13.1, p=0.05) independent of age, HTN stroke score, and anti-HTN treatment. Anti-HTN medication was not associated with OH, p=0.48	In pts with mild to moderate ischemic stroke, antihypertensive therapy is not associated with OH. Presence of cardiac dysfunction was associated with OH

Poon IO, et al.	Aim: To describe prevalence of	Inclusion: Pts ≥75 y of	19 and nainty Drayalance of OL modication provalence	- With increasing number of
· ·	· —		1º endpoint: Prevalence of OH, medication prevalence	With increasing number of
2005	symptomatic and asymptomatic	age, with documented		causative medications, the
<u>15811171</u>	OH in elderly veterans and	sitting and standing BP	Results:	prevalence of OH increased. The
(349)	relation to medications	readings, who attended	189 (55%) pts had OH. Prevalence of OH in pts who had no	highest association among cardiac
		geriatric clinic in electronic	causative medication was 35%. Prevalence OH in pts on 1, 2, or ≥	medications included HCTZ and
	<b>Study type:</b> Retrospective	medical record database	3 causative medications was 58%, 60%, and 65% respectively,	lisinopril. The effect of work-up bias
	chart review,	(MEDVAMC) between	with a significant relationship x <sup>2</sup> =15.18, p=0.002)	is not accounted for, as there are
		6/2002 and 6/2003		many pts on these medications
	<b>Size</b> : n=342 pts		Associated with highest prevalence of OH was	without orthostatic symptoms or BP
		Exclusion: Pts unable to	hydrochlorothiazide (65%), lisinopril (60%), furosemide (56%), and	measurements.
		stand, no assessment of	terazosin (54%) for cardiac medications. Other medications	<ul> <li>OH defined as SBP reduction ≥20</li> </ul>
		sitting and standing BP,	associated with OH included paroxetine (86%), trazodone (58%),	mmHg or DBP ≥10 mmHg within 3
		autonomic dysfunction,	olanzapine (57%), and quetiapine (56%)	mins of standing +/- symptoms
		Parkinson disease.		Potentially causative medications
				of OH were those reported with >1%
				incidence of OH
Raiha I, et al.	Aim: To evaluate predisposing	Inclusion: Baseline and 10	1° endpoint: Prevalence of postural hypotension, 10 y mortality	Only supine HTN was associated
1995	factors to postural hypotension	y follow-up survey of elderly	1 enaponit. The valence of postural hypotension, To y mortality	with postural hypotension, but there
			Decultor	
<u>7726701</u>	in elderly	(pts >65 y of age) in Turku,	Results:	was not effect on mortality. No
(350)	Ct t. t	Finland in 347 pts.	Prevalence of postural hypotension was 28%.	medication (nitrates, diuretics, BB,
	Study type: Analytical,		Predisposing factors for postural hypotension: elevated supine BP	orther antihypertensives) was
	observational, prospective	Exclusion: Living in an	(p<0.001).	associated with postural
	cohort	institution	Chronic CV diseases, body mass index, medication, and	hypotension.
			abnormal ECG were not associated with postural hypotension	<ul> <li>Postural hypotension was defined</li> </ul>
	<b>Size</b> : n=347 pts			as ≥ 20 mmHg after 3 mins of
				standing.

## Data Supplement 38. Nonrandomized Trials, Observational Studies, and/or Registries of Pseudosyncope – (Section 8)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)

Moya, et al. 2009 <u>19713422</u> (351)	Study type: Practice guideline consensus (European Society of Cardiology)  Size: N/A	Inclusion criteria: N/A  Exclusion criteria: N/A	1° endpoint: N/A  Results: 1. Frequent attacks, often many times a d 2. Eyes closed 3. Prolonged episodes, often many mins in duration 4. No apparent trigger for attack 5. Prone to being 'suggestible' which favors triggering attacks in clinic/laboratory	Summarizes the key clinically useful markers to aid recognition of PPS/PNES
Tannamaat, et al. 2013 23873974 (126)	Study type: Tilt-test induction of PPS/PNES examined retrospectively to assess clinical features  Size: n=43 pts with PPS/PNES vs. 69 pts with vasovagal syncope	Inclusion criteria: Diagnosis of PPS/PNES by Tilt-test and video EEG  Exclusion criteria: N/A	1° endpoint: Pseudosyncope  Results PPS/PNES can be diagnosed and differentiated from vasovagal syncope by use of a tilt-test.	Provides a quantitative assessment of clinical features distinguishing PPS/PNES from vasovagal syncope
McKenzie PS, et al. 2010 21421771 (352)	Study type: Retrospective observational study of PNES pts diagnosed by inpatient or outpatient EEG or video-EEG  Size: n=187 pts	Inclusion criteria: Diagnosed PPS/ PNES  Exclusion criteria: N/A	1° endpoint: New onset of medically unexplained symptoms (MUS) in pts diagnosed with PPS/PNES.  Results: Approx. 25% of PNES pts develop new medically unexplained symptoms after initial diagnosis	Many PPS/PNES pts exhibit other medically unexplained symptoms, but in most cases the medically unexplained symptoms were present prior to diagnosis of PPS/PNES and only infrequently became manifest for the first time later during the approx. 1 y follow-up.
Iglesias, et al. 2009 <u>19250095</u> (353)	Study type: Single center prospective syncope evaluation  Size: n=131 PPS/PNES cases out of 939 pts undergoing TLOC evaluation	Inclusion criteria: Presentation of TLOC or apparent TLOC  Exclusion criteria: N/A	1° endpoint: Frequency of PPS/PNES in a TLOC population  Results: 14% of all pts were considered PPS/PNES. Approx. 60% are young woman with multiple presyncope and syncope	A stepwise evaluation of apparent TLOC cases in an ambulatory clinic may yield a diagnosis in 2/3. More than 50% of cases are either vasovagal syncope or PPS/PNES.

Elliot JO, et al. 2014 <u>25262500</u> (354)	Study type: Observational Quantitative assessment in PNES alone or PNES with epilepsy  Study size: PNES alone 84, PNES + epilepsy 281; No Controls	Inclusion criteria: Retrospective study of pts admitted to an epilepsy monitoring unit over a 6 y period  Exclusion criteria: N/A	1° endpoint: Predictors of video-EEG confirmed PPS/PNES in an epilepsy monitoring unit  Results:  • 5 Biologic predictors of PNES alone  • 1 Psychological predictor  • 2 Social predictors	Psychosocial issues (e.g., anxiety, physical/sexual abuse) as well as comorbidities (e.g., prior head injury, GERD) are important features of PPS/PNES pts.
Mayor, et al. 2012 23168089 (355)	Study type: Prospective observational  Size: n=44 previously diagnosed cases	Inclusion criteria: Prior diagnosis of PPS/PNES in which pts completed self-reporting symptom questionnaires or otherwise reported symptom frequency during follow-up  Exclusion criteria: N/A	1° endpoint: Symptom recurrence after being told the nature of the diagnosis  Results:  Median self-reported symptom frequency dropped from 10 to 7.5/mo over 6 mo. 7 of 44 became symptom free, and 10/44 had >50% reduction of event frequency. Nevertheless, baseline levels of life-style impairment did not improve.	Apart from identifying the diagnosis of PPS/PNES, further efforts are needed to diminish adverse life-style impact of this condition.
Mayor, et al. 2010 <u>20561022</u> (356)	Study type: Prospective observational of psychodynamic psychotherapy (no controls)  Size: n=66 pts of whom 47 were followed full study duration	Inclusion criteria: Diagnosed PPS/PNES  Exclusion criteria: N/A	1° endpoint: PPS/PNES event frequency  Results: With follow-up of 12–61 mo (mean 50 mo), 25% were symptom free and 40% achieved event reduction >50%. Health care utilization declined significantly (p=0.039)	Psychodynamic interpersonal therapy may be associated with reduction of symptom frequency and healthcare utilization.
Reuber M, et al. 2007 <u>18061753</u> (357)	Study type: Uncontrolled observational assessment of tailored psychotherapy in pts with functional neurologic impairment  Size: n=91 enrollees; 63 completed treatment and 34 completed final questionnaires	Inclusion criteria: Functional neurological symptoms but NOT just PPS/PNES  Exclusion criteria: N/A	1º endpoint: Therapeutic impact of individualized psychotherapy using validated questionnaires  Results:  Questionnaires throughout approx. 6 mo follow-up revealed that multiple patient-centered psychiatric instruments improved by at least 1 SD in 50% of pts	Individualized psychotherapy may be beneficial but one-size does not fit all.

LaFrance Jr WC, et al. 2010 20739647	Study type: Prospective double-blind RCT of sertraline in PPS/PNES	Inclusion criteria: Diagnosed PPS/PNES	1° endpoint: Symptom frequency sertraline vs. placebo	Sertraline initially appeared to be more effective than placebo with reduction of symptom frequency from baseline.
(358)	Size: 38 enrollees; n=26 completed study	Exclusion criteria: N/A	Results: Sertraline was associated with 48% symptom reduction vs. 8% with placebo. However, intention-to-treat not reported and baseline differences resulted in no significant difference	However, after adjustment for baseline differences the effect was deemed nonsignificant.
Santos, et al. 2014 <u>25650860</u> (359)	Study type: Observational_effects of psychoanalytic therapy; no controls  Size: n=37 pts	Inclusion criteria: PNES diagnosed by video- EEG  Exclusion criteria: N/A	1° endpoint: Symptom recurrence frequency during follow-up  Results: During 1 y follow-up, 30% had cessation of symptoms,	Individual psychoanalytic therapy proved beneficial in this uncontrolled study
	5.25 5. p.0		and 51% had reduced number of attacks.	

# Data Supplement 39. RCTs for Pseduosyncope – (Section 8)

Study Acronym; Author; Year Published	Aim of Study; Study Type; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Absolute Event Rates, P values; OR or RR; & 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
Goldstein LH, et al. 2010 20548043 (360)	Study type: RCT  Size: n=36 pts randomized to standard therapy vs. CBT psychotherapy	Inclusion criteria: Diagnosis of PPS/PNES  Exclusion criteria: N/A	Intervention: CBT in addition to standard therapy  Comparator: Standard therapy alone	1° endpoint: Symptom recurrence frequency  Result With short-term application of CBT, the CBT group tended to have a better 3-mo event freedom (OR: 3.125, p<0.086)  Safety endpoint (if relevant): N/A	CBT tended to improve short-term outcomes but larger controlled studies are needed.

## Data Supplement 40. Nonrandomized Trials, Observational Studies, and/or Registries of Pediatrics – (Section 10.1)

Study Acronym; Author; Year Published	Aim of Study; Study Type*; Study Size (N)	Patient Population	Study Intervention (# patients) / Study Comparator (# patients)	Endpoint Results (Include Absolute Event Rates, P value; OR or RR; and 95% CI)	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events; Summary
Zhang Q, et al. 2009 <u>19183119</u> (361)	Aim: Aimed to measure the diagnostic value of a protocol on the management of children and adolescents with syncope.  Study type: Multi center, prospective consecutive pts <18 y of age with syncope.  Size: n=474 consecutive pts presenting with syncope. (20 mo period)	Inclusion criteria: <18 y of age with syncope as defined as TLOC and postural tone caused by cerebral hypoperfusion  Exclusion criteria: Pts with symptoms compatible with seizures, vertigo, or shock were excluded.	Intervention:  1st Step: H&P, and ECG  2nd Step: Echo, Holter, CT, Psych evaluation. 2nd Step diagnostic maneuvers were only performed if 1st step did not yield a definitive diagnosis. HUTT was only used if unexplained syncope.  Comparator: None	1° endpoint: Initial diagnostic work-up (H&P & ECG) gave a definitive diagnosis in 59 (12.4%). 2nd Step diagnostic work-up required in 326 (87%).  1) n=382 HUTT identified VVS in 203, POTS in 87. No final diagnosis in 89 pts (TILT YIELD): 76%  2) n=10 had a neurological event (additional testing is unnecessary unless challenged by H&P).	HUTT can help with the diagnosis. An extensive neurological work-up is not indicated unless the H&P is suspicious for a neuro condition (i.e. vertigo seizure)  Summary: HUTT can help make the diagnosis of VVS. An extensive neurological work-up should be reserved for pts whose H&P is concerning for a neuro condition.
Miyake, et al. 2015 <u>26277987</u> (362)	Aim: Aimed to evaluate the incidence of cardiac disorders among children with midexertional syncope.  Study type: Single center, retrospective evaluation of children who presented for cardiac evaluation with exertional syncope (1999-2012)  Size: n=60 pts	Inclusion criteria: ≤18 y of age with mid-exertional syncope an EKG and ECHO and at least one of the following: TTT, EST, EPS  Exclusion criteria: Pts with known structural heart defects or known arrhythmia disorders	Intervention: None, Clinical Evaluation Only  Comparator: None	1° endpoint: 28 Non cardiac Diagnosis 32 Cardiac Diagnosis LQT (n=10) CPVT (n=6) SVT (n=5) VT (n=2) VF (n=2) HCM (n=2) LVNC (n=1)  No difference in symptoms between cardiac and noncardiac pts preceding syncope or following syncopal event.	Reported symptoms before and after a mid-exertional syncopal event may not distinguish between a benign noncardiac condition and a cardiac condition.      Summary:     Mid-exertional syncope in children carries a high-risk of being diagnosed with a cardiac condition.

Zhang, et al. 2013 22417947 (363)	Aim: Value of Hx taking in identifying children with cardiac syncope  Study type: Multicenter prospective consecutive series of pts in the Pediatric Syncope Unit  Size: n=275 pts <18 y of age	Inclusion criteria: ≤18 y of age with suspected syncope admitted to the Pediatric Syncope Unit of 5 hospitals in China  Exclusion criteria: Pts with known CHD or known arrhythmia disorders	Intervention: Clinical history, physical exam, BP measurements and ECG. All pts complete 118 item questionnaire  Comparator: None	1º endpoint: Clinical diagnosis made	Results Cardiac 31 (11%) Autonomic mediated 214 (78%) Unexplained 15 (5%) Summary: Multivariate analysis showed the history of exercise-triggered syncope or ECG abnormalities were independent predictors of cardiac syncope.
Qingyou, et al. 2004 14727100 (364)	Aim: To determine usefulness in children with unexplained syncope.  Study type: Single center prospective study of pts with unexplained syncope.  Size: n=47 pts divided into a positive response group (I) and a negative tilt response group (II)	Inclusion criteria: ≤18 y of age with unexplained syncope.  Exclusion criteria: Pts with known structural heart defects or known arrhythmia disorders	Intervention: All syncopal pts (all unexplained) had a normal exam, EKG, Echo, and head CT).  Comparator: Positive tilt vs. Negative tilt groups	1º endpoint: Clinical diagnosis made	Results HUTT positive results more common in 12– 16 y of age than younger children. Prodrome of syncope had an odds ratio of 17 in predicting positive TTT results.  Summary: Clinical history of a prodrome prior to syncope in conjunction with a positive HUTT supports diagnosis of vasovagal syncope.
Udani, et al. 2004 <u>15269465</u> (365)	Aim: Aimed to measure the diagnostic value of a HUTT  Study type: Single center, prospective consecutive pts <18 y of age with syncope.  Size: n=18 pts	Inclusion criteria: <18 y of age with strong clinical suspicion of neurocardiogenic syncope  Exclusion criteria: N/A	Intervention: HUTT following Hx and clinical examination  Comparator: None	1° endpoint: Recurrent syncope	16/18 (90%) with clinical suspicion of vasodepressor syncope had a positive tilt test      Summary: HUTT can help make the diagnosis of neurocardiogenic syncope.
Fouad, et al. 1993 7681189 (366)	Aim: Measure the diagnostic value of a HUTT in pts with syncope compared to healthy controls w/o syncope  Study type: Single center, retrospective study of syncopal pts and prospective	Inclusion criteria: <18 y of age with strong clinical suspicion of neurocardiogenic syncope  Exclusion criteria: N/A	Intervention: HUTT following Hx and clinical examination  Comparator: Healthy controls	1° endpoint: Syncope on tilt test	<ul> <li>25/44 (58%) of symptomatic pts ha a positive tilt</li> <li>3/18 (17%) normal volunteers had a positive tilt</li> <li>Sensitivity of a positive tilt 57% and specificity 83%</li> </ul> Summary: HUTT has a high specificity in

	study of healthy controls				diagnosing vasodepressor syncope.
	Size: n=44 syncope pts (16±3 y vs. 18 healthy controls (16±2 y)				
Lerman-Sagie, et al. 1991 2019920 (367)	Aim: Measure the diagnostic value of a HUTT in pts with syncope compared to healthy controls w/o syncope  Study type: Single center, prospective study  Size: n=15 syncope pts (10–18 y of age vs. n=10 healthy controls (11–18 y of age)	Inclusion criteria: <18 y of age with strong clinical suspicion of neurocardiogenic syncope.  Exclusion criteria: Healthy controls without syncope.	Intervention: HUTT following Hx and clinical examination  Comparator: Healthy controls	1° endpoint: Syncope on tilt test	6/15 (43%) of symptomatic pts ha a positive tilt     0/10 (0%) normal volunteers had a positive tilt     Summary: HUTT offers a simple, noninvasive, high-yielding diagnostic tool for the evaluation of syncope in children.
Al Dhahri, et al. 2009 <u>19694968</u> (368)	Aim: Measure the usefulness of ILR in children with unexplained syncope.  Study type: Retrospective study of pts with unexplained syncope after initial evaluation identified cause of syncope.  Size: 42 pts (25 males) with a median age of 11.5 y of age (1.4–19.0 y of age) underwent ILR implantation. There were 14 pts (33%) with normal ECGs and echocardiograms. In these pts, the ILR device was implanted at a median age of 12.4 y of age (2.7–17.5 y of age).	Inclusion criteria: Pts with unexplained syncope undergoing ILR after conventional diagnostic testing failed to provide a definitive diagnosis.  Exclusion criteria: None	Intervention: ILR implantation  Comparator: None	1° endpoint: Identification of a substrate on ILR interrogation to explain causal syncope.	Among the 21 pts who presented with syncope, 14 of 21 (67%) were diagnosed with reflex-mediated syncope, 2 of 21 (9%) with seizures, and 2 of 21 (9%) with arrhythmias, while in 3 of 21 (15%) other causes were found, but we were able to rule out arrhythmias as a possible etiology.  Summary: ILR may be beneficial in children with syncope of unknown etiology to rule-out arrhythmias as a cause of syncope. The risk of infection and need for device removal is rare.

Babikar, et al. 2007 <u>17764457</u> (369)	Aim: Measure the usefulness of ILR in children  Study type: Retrospective single center  Size: n=23 pts (11.4± 4.3 y of age) underwent ILR. 11 pts with syncope and 3 with presyncope underwent ILR.	Inclusion criteria: Pediatric pts undergoing ILR  Exclusion criteria: None	Intervention: ILR implantation  Comparator: None	1° endpoint: Identification of a substrate on ILR interrogation to explain causal syncope.	14 pts (61%) underwent ILR for recurrent syncope or presyncope. ILR uncovered:  • Polymorphic VT (n=1)  • SVT (n=1)  • Type II AV block (n=1)  1 pts would infection and 1 pts relocated for discomfort  Summary: ILR facilitated diagnosis in majority of pts with syncope or pre-syncope with a relatively low complication rate.
Rossano, et al. 2003 12949317 (370)	Aim: Measure the usefulness of ILR in children  Study type: Retrospective multi-center center  Size: n=21 pts (12.3± 5.3 y of age) underwent ILR. Of these, 16 underwent ILR for unexplained syncope.	Inclusion criteria: Pediatric pts undergoing ILR where conventional testing failed to produce a diagnosis.  Exclusion criteria: None	Intervention: ILR implantation  Comparator: None	1° endpoint: Identification of a substrate on ILR interrogation to explain causal syncope.	Of the 16 pts, 6 (40%) were identified as having an arrhythmia to explain syncope.  • Junctional bradycardia (1)  • SVT (2)  • TdP (1)  • Asystole (1)  • VT (1)  No complications of ILR  Summary: ILR facilitated diagnosis in majority of pts with syncope or presyncope with zero complication rates.
Ergul, et al. 2015 <u>25348219</u> (371)	Aim: Measure the usefulness of ILR in children  Study type: Retrospective single-center center  Size: n=12 pts (9.4± 4.3 y of age) underwent ILR. All had a structurally normal heart with exception 1 pts having TOF. Of the 12 pts 6 had exertional syncope.  Average monitoring period: 20 mo	Inclusion criteria: Pediatric pts with unexplained syncope undergoing ILR. All pts had a normal ECG and event recorder and 10/12 had a normal EST.  Exclusion criteria: None	Intervention: ILR implantation  Comparator: None	1° endpoint: Identification of a substrate on ILR interrogation to explain causal syncope.	6 pts, (50%) were identified as having presyncope:  PMVT (3) CPVT (1) Asystole (1) No complications of ILR Of the 6 pts with exertional syncope, 4 were identified as having a malignant arrhythmia.  Summary: ILR is useful in establishing symptom rhythm correlation in the majority of pts with unexplained syncope.  ILR should strongly be considered in pts with unexplained exertional syncope.

Vlahos, et al. 2008 <u>17899242</u> (372)	Aim: Understand the relationship of family Hx in diagnosing syncope  Study type: Retrospective single center, case-control  Size: n=76 pts (11.8±2.9 y of age) with syncope and n=29 control non syncopal pts (11.3±2.9 y of age)	Inclusion criteria: Syncope diagnosis  Exclusion criteria: None	Intervention: None  Comparator: None	1° endpoint: Comparison family Hx of syncope between 2 groups	Of the 76 pts with diagnosis of syncope, 68 had a positive family history of syncope (89%) compared to 1/29 (3.5%)  Summary: Family Hx of vasovagal symptoms should be meticulously sought and is of value in the diagnosis of neurocardiogenic syncope in pediatric pts.
Alehan, et al. 1996 <u>8833492</u> (373)	Aim: Assess sensitivity and specificity of TTT  Study type: Prospective single center, case-control  Size: n=20 pts (12.0±2.5) with unexplained syncope and 10 healthy controls	Inclusion criteria: Syncope diagnosis  Exclusion criteria: No identifiable cause of syncope following ECG, ECHO, EEG, Hx & physical exam	Intervention: HUTT 25 mins  Comparator: 10 healthy age-matched controls	1° endpoint: Tilt results	1) During TTT, symptoms were elicited in 15 (75%) of the pts with unexplained syncope but in only one (10%) of the control group (p<0.001). 2) Sensitivity 75% 3) Specificity 90% 4) 40% of positive tilt responders had a family Hx  Summary: The head-up tilt test is a noninvasive, sensitive, specific diagnostic tool for evaluating children with unexplained syncope.
Thilenius, et al. 1991 <u>2000273</u> (374)	Aim: Assess sensitivity and specificity of TTT  Study type: Prospective single center  Size: n=35 pts (8-19) with unexplained syncope	Inclusion criteria: Syncope diagnosis  Exclusion criteria: No identifiable cause of syncope following ECG, ECHO, EEG, H&P	Intervention: HUTT  Comparator: None	1° endpoint: Tilt results	During TTT, symptoms were elicited in 26 (75%) of the pts with unexplained syncope.      Summary: The head-up tilt test is a noninvasive, sensitive, specific diagnostic tool for evaluating children with unexplained syncope.

Salim, et al. 2005 <u>15708690</u> (238)	Aim: Effectiveness of salt and fludocrotisone in prevention of VVS in children  Study type: Randomized (pediatric)  Size: n=32; florinef 0.1mg/day and salt 1g/d n=18;	Inclusion criteria: >1 syncope or presyncope; +HUTT; <18 y of age; no prior therapy for syncope  Exclusion criteria: No structural heart disease	Intervention: Florinef 0.1mg/day and salt 1g/d Comparator: Placebo	1° endpoint: Syncope or pre-syncope recurrence  1° Safety endpoint (if relevant):	Follow up 176+117d; recurrence 36% in controls and 55% active arm (p<0.04).      Summary: Symptoms were more frequent in the placebo group.
Massin MM, et al. 2004 15289772 (375)	control n=14  Aim: Analyzed the etiology of consecutive cases of syncope presenting to a pediatric emergency room.  Study type: Prospective cohort study  Size: n=252 presentations of syncope in 226 pts (mean age 10.8 ± 3.6 y of age)	Inclusion criteria: Primary complaint of syncope (witnessed and unwitnessed) upon presentation to the emergency department.  Exclusion criteria: None	Intervention: None  Comparator: None	1º endpoint: Clinical diagnosis  Safety endpoint: None	Of the 226 pts presenting with syncope, neurocardiogenic accounted for 80% of the diagnosis. Neurologic disorders were identified in 9%. A prodrome was a significant (p<. 05) factor in diagnosing neurocardiogenic syncope (present in 88% of cases); however a prodrome was also observed in 52% of those with a neurologic disorder.  Clinical Hx with particular attention to the events is the most critical piece of information required.  Limitation: ECG were not obtained in 58% of the pts and as such the utility of an ECG cannot be measured in this study.
Chen L, et al. 2011 21629199 (376)	Aim: Analyze the spectrum of underlying diseases in children presenting with syncope.  Study type: Multicenter retrospective chart review  Size: n=888 children (median age 12.0 ± 3.0 y of age)	Inclusion criteria: Presentation with syncope  Exclusion criteria: None	Intervention: All pts underwent H&P, orthostatic vital sign measurements and an ECG.  Comparator: None	1º endpoint: Clinical diagnosis  Safety endpoint: None	Vasovagal syncope was diagnosed in 32% of pts. POTS was diagnosed in 32% of pts. Cardiogenic syncope accounted for 1.5% of the cases. Approximately 31.5% of the cases of syncope were undiagnosed.

Colman N, et al. 2009 <u>19482852</u> (377)	Aim: To determine whether Hx taking can be used as a tool in identifying pts presenting with syncope who are more likely to have LQT	Inclusion criteria: All LQT pts confirmed genotype positive.  Exclusion criteria: >40 y of	Intervention: Clinical assessment with detailed Hx and detailed family Hx.  Comparator: LQT pts	1° endpoint: Clinical comparison  Safety endpoint: None	Results: 72% of pts with LQTS had a family Hx of syncope and 66% had a family Hx of sudden death. This is in contradistinction to pts presenting to the ED with syncope without LQT where the family Hx of syncope
	syndrome.  Study type: Retrospective study comparing 2 populations. The control cohort was evaluated as part of a Dutch Fainting	age.	compared to a consecutive heterogeneous group of patients with syncope presenting to the emergency department		was 9% and sudden death 10% (p<0.001).  Syncope while supine and syncope with exercise were significantly more common in the LQTS cohort compared to the ED cohort.
	Assessment Trial  Size: n=32 LQTS pts, n=113 pts in ED with syncope, and n=69 known vasovagal syncope pts.				Summary: A family Hx or syncope and sudden cardiac death are important questions that should be asked when evaluating a young group of pts with syncope.
Tretter JT, et al. 2013. 23992679 (378)	Aim: To identify characteristics that distinguishes VVS from cardiac syncope.  Study type: Retrospective review of pts presenting a vasovagal syncope vs. cardiac syncope.	Inclusion criteria: All pts (newborn to 18 y of age) presenting to the outpatient faculty with diagnosis of syncope)  Exclusion criteria: None	Intervention: None  Comparator: Vasovagal Symptoms vs. Cardiac Syncope Symptoms (identified from the ICD database and the cardiac stress lab database)	1º endpoint: Syncope at follow-up and comparison between 2 groups of etiology  Safety endpoint: None	Results: 1. There was no difference between the 2 groups with respect to chest pain or palpitations. 2. Preceding symptoms of lightheadedness, dizziness, visual and hearing changes were significantly less common in the cardiac group (41% vs. 84%). 3. ECG established the diagnosis 47% of
	Size: n=89 pts 4–18 y of age presenting to cardiology outpatient. Compared to 17 pediatric pts over the same era that were diagnosed with cardiac syncope.				time compared to 0% in vasovagal cohort. 4. 11/17 (65%) with cardiac syncope had episodes of syncope surrounding exertion.  Summary: Any one of the following 4 parts of a cardiac screen: (1) abnormal cardiac physical exam ± (2) abnormal findings on ECG ± (3) concerning family Hx ± (4) exertional syncope has 100% specificity and 60% specificity.

Ritter S, et al. 2000. 10799622. (379)	Aim: Understand the clinical symptoms in pts with syncope.  Size: n=480 pts (1.5 to 18 y of age)	Inclusion criteria: Syncope diagnosis  Exclusion criteria: Pts with previously known cardiac disease (cardiomyopathies, arrhythmias, or CHD)	Intervention: None  Comparator: None	1° endpoint: Use of H&P, and ECG in identifying pts with cardiac syncope.  Safety endpoint: None	Results: Of the 21 pts with cardiac related syncope, a (1) personal Hx of exercise induced syncope; (2) positive family Hx, (2) abnormal ECG, and 4) normal echo.
MacCormick JM, et al. 2011 21616715 (380)	Aim: Understand the signs and symptoms before the cardiac syncope and before the patient was diagnosed with a channelopathy.  Study type: Retrospective review of consecutive gene positive probands and symptoms before syncope.  Size: n=35 pts (8-19) with unexplained syncope	Inclusion criteria: Syncope diagnosis amongst consecutive gene positive probands.  Exclusion criteria: Pts with syncope and LQT that was not genetically confirmed.	Intervention: None  Comparator: Comparison was done on a historical and literature based control not in the same time period or by same authors.	1° endpoint: Clinical presentation of syncope.  Safety endpoint: None	Results: 20 pts with syncope (median age 13.9 y of age) with 17 describing symptoms prior to syncope (lightheadedness and dizziness in 47%). Similarly drowsiness and weakness post –syncope were noted in 64% of cases.  Summary: Young pts with cardiac syncope frequently have symptoms similar to neurocardiogenic syncope. The presence of symptoms before and after fainting may not completely distinguish between benign neurocardiogenic and cardiac syncope.
Grubb BP, et al. 1992 <u>1382276</u> (381)	Aim: Understand the utility of HUTT testing in the evaluation of recurrent syncope of unknown etiology in children and adolescents.  Study type: Prospective study  Size: 30 pts (15 males and 15 females; mean age: 14 ± 6 y of age)	Inclusion criteria: A minimum of 3 episodes of syncope in the preceding 6 mo with the cause of syncope unknown by H&P, ECG, echocardiogram, and exercise stress test.  Exclusion criteria: None	Intervention: Baseline HUTT (30 mins) with or without isoproterenol.  Comparator: None	1° endpoint: Clinical outcomes following HUTT results.  Safety endpoint: None	Results: During the baseline HUTT 6 pts (20%) had a positive HUTT and 15 additional pts (50%) during an isoproterenol infusion (total 70%) had a positive HUTT. A variety of treatments were used including BB, Florinef, and transdermal scopolamine. No further syncope occurred. This study was not designed to look at one particular treatment arm over another but asses the utility of the HUTT itself.
Numan M, et al. 2015. 25087055 (382)	Aim: To report experience with pts with cardiac asystole during HUTT	Inclusion criteria: Cardiac asystole (defined as absence of ventricular activity of >3 s)	Intervention: No uniform treatment strategy follow-up of cardiac asystole. All pts received education of	1° endpoint: Clinical recurrent syncope  Safety endpoint: None	25 pts with cardiac asystole (mean pause 9.2± 5.8 s) were managed with education, symptom awareness, and one of the following Florinef, BB, alpha agonists and all

Yilmaz S, et al. 2012. 22459868 (383)	Study type: Retrospective study, no placebo group.  Size: Retrospective analysis of 537 pts (age 6-22 y of age) and follow-up of 25 pts with cardiac asystole. Follow-up 19 ± 10 mo  Aim: Define predictors of recurrence of vasovagal syncope.  Study type: Retrospective observational study  Size: 150 pts (8–18 y of age) between 2007–2011. Group I HUTT positive (N=97)	Exclusion criteria: None  Inclusion criteria: 8–18 y of age with clinical VVS.  Exclusion criteria: Excluded CHD, LQT, Brugada, or medications that affect the heart rate.	symptom awareness, fluids and salt and additional treatment.  Comparator: None  This study did not compare medical management vs. pacemaker therapy.  Intervention: VVS pts follow after HUTT.  Comparator: Compare Recurrent syncope group (n=40) and Non-recurrent syncope group (n=110).  Average Follow up: 3.8±4.7 y	1° endpoint: Syncope recurrence Safety endpoint: None	but one responded to medical management. Only 1 patient required a pacemaker for failing numerous pharmacologic strategies.  Summary: Children and young adults (<25 y of age) with cardiac asystole at time of HUTT can be managed with pharmacologic agents and do not necessarily need a pacemaker immediately.  Recurrent syncope predictors: age at initial syncope, positive family Hx of syncope, and number of previous syncopal episodes were predictive of recurrent syncope. Positive HUTT did not predict recurrence of VVS.  Summary: Number of prior syncopal episodes and family Hx of syncope predict clinical recurrence of VVS. Result HUTT does not
Liu JF, et al. 2011 21329841 (197)	and Group II HUTT negative (n=53 pts) and follow to see if clinical VVS reoccurs. Average age of 1st syncope (12.3±3.1 y)  Aim: Identify risk factors for recurrent syncope in children and adolescents with LQT syndrome.  Study type: Retrospective review of data from the International Long QT Registry.  Size: n=1,648 pts <20 y of age with LQT (genotype or genotype and phenotype)	Inclusion criteria: QTc ≥450 msec, or a known pathogenic QT mutation, and syncope.  Exclusion criteria: QTc ≤450 ms without pathogenic mutation.	Intervention: Registry follow-up  Comparator: Different LQT genotypes and BB utilization with recurrent syncope.	1° endpoint: Occurrence of recurrent syncopal episodes.  Safety endpoint: Aborted cardiac arrest and LQT related sudden cardiac death as a defined endpoint.	Results: A QTc ≥ 500 ms was a significant predictor of a first syncopal event (HR: 2.16). LQT1 male pts had the highest rate of first syncope and LQT2 females had the highest rate of first and subsequent syncopal events. BB treatment for LQT1 & LQT 2 pts significantly (>70%) reduced subsequent syncopal events.

Younoszai AK, et al. 1998 <u>9491043</u> (384)	Aim: Assessment of oral fluid therapy in children with vasodepressor syncope on clinical recurrence.  Study type: Retrospective, non comparison study  Size: 58 pts (8.7–27.6 y)	Inclusion criteria: Clinical diagnosis of VDS and positive TTT  Exclusion criteria: Tilt positive with isoproterenol.	Intervention: Following a positive TTT pts were prescribed oral fluid therapy (64 oz/daily) and encouragement to drink more fluid and avoid caffeine.  Comparator: None	1° endpoint: 90% had resolution of syncope  Safety endpoint: Tolerance of fluid bolus.	Results:  Treatment of neurally-mediated syncope with oral rehydration reduced the number of syncopal events.  No control and not randomized, cannot account for placebo effect.
Chu W, et al. 1998 <u>25577227</u> (385)	Aim: Whether oral rehydration salts is effective in treatment of children with VVS  Study type: Single center, randomized; placebocontrolled. 6 mo-f/u  Size: Group I (n=87) conventional therapy (health education, tilt training, and oral rehydration salts) vs. Group II (n=79) conventional therapy.	Inclusion criteria: At least 2 episodes of syncope in prior 6 mo. Positive HUTT with clinical diagnosis of VVS. (Children 7-17)  Exclusion criteria: Other disease ruled-out by ECG, EEG, and head imaging.	Intervention: Conventional therapy +/- oral rehydration salts (oral rehydration salts: glucose, NaCl, KCl, dissolved in 500 ml H2O) for 6 mo  Comparator: Placebo plus conventional therapy (education – symptom awareness)	1° endpoint: Clinical symptoms  Safety endpoint:	Results:  Group I (oral rehydration salts): No recurrence (56%), Decrease in syncope (39%) and No change in syncope (5%).  Group II (Placebo): No recurrence (39%), Decrease in syncope (47%) and No change in syncope (14%). p<0.05  Summary:  Oral Rehydration Salts significantly reduced the recurrence rate of syncope in children 7–17 y of age.
Strieper MJ, et al. 1993. <u>8101533</u> (386)	Aim: Whether alpha- adrenergic agonist prevents syncope  Study type: Single center, prospective study  Size: n=16 pts (mean 13 y of age)	Inclusion criteria: Recurrent syncope and a positive HUTT  Exclusion criteria: Free of any other cardiac medication.	Intervention: Following HUTT discharged on pseudoephedrine 60 mg PO BID  Comparator: None	1° endpoint: Clinical symptoms  Safety endpoint: Tolerance of alpha-adrenergic medication.	Results: Follow up: 11.6 mo; 15/16 (94%) pts reported control of clinical symptoms.  Summary: Pseudoephedrine alleviates syncope in children without significant side effects.
Qingyou Z, et al. 2006. 17137891 (243)	Aim: Efficacy of midodrine in preventing VVS in children.  Study type: Single center, randomized control trial between 2003-2004. Not	Inclusion criteria: At least 3 episodes of syncope in prior 12 mo and "positive" tilt with clinical diagnosis of VVS.  Exclusion criteria: At least 3	Intervention: Conventional therapy + midodrine (Group I) or sole conventional therapy without midodrine (Group II).	1° endpoint: Syncope recurrence (AIM 1) and repeat HUTT (AIM 2) Safety endpoint:	Results:  • Group I (Midodrine): Effective rate of repeat HUTT evaluation 75%. Recurrence rate of clinical syncope: 22%.  • Group II (Conventional): Effective rate of repeat HUTT evaluation 20%. Recurrence

	blinded, no placebo.  Size: Group I (n=13) midodrine & conventional therapy (health education, tilt training, and salt) vs. Group II (n=13) conventional therapy only. 6 mo follow-up plus repeat HUTT.	episodes of syncope in prior 12 mo AND "positive" tilt with clinical diagnosis of VVS.	Comparator: Midodrine vs. no additional pharmacotherapy 6 mo follow-up. Not blinded, no placebo, no control.		rate of clinical syncope 80% (p<0.05)  Summary:  • Midodrine is effective in reducing clinical recurrence of syncope.  • No significant adverse side effects of midodrine.
Zhang Q, et al. 2008. 18376348 (387)	Aim: Efficacy of BB in conjunction with conventional treatment in reducing VVS in children.  Study type: Single center, prospective randomized. (2001-2003)  Size: n=28 pts; Age 12.3±3 y of age with 22±10 mo. Group I (n=14 pts) Metoprolol and Group II (n=14 pts) control	Inclusion criteria: At least 3 episodes of syncope in prior 12 mo along with a positive tilt.  Exclusion criteria: Other causes of CV or systemic causes of syncope.	Intervention: Conventional therapy + metoprolol (Group I) or sole conventional therapy without metoprolol (Group II).  Comparator: Metoprolol vs. conventional therapy.	1° endpoint: Recurrence of syncope in 2 wk after beginning therapy. Presyncope symptoms were not considered a failure of therapy.  Safety endpoint: None	Results: Group I (Metoprolol): Syncope recurrence 6/14 (43%)  • Group II (Conventional): Syncope recurrence 4/14 (29%)  Summary: In a prospective randomized study Metoprolol was not effective in reducing VVS in children.
Scott WA, et al. 1995 7639169 (388)	Aim: Comparison of Atenolol vs. Florinef in treatment of neurally mediated syncope  Study type: Prospective randomized  Size: n=58 pts	Inclusion criteria: ≥2 episodes of syncope in preceding 6 mo and a positive TTT (BL or Isuprel). All pts had a normal H&P, ECP, and echocardiogram.  Exclusion criteria: None	Intervention: Following a positive TTT randomized to Atenolol (25 or 50 mg) or Florinef (0.1 mg) followed 6 mo  Comparator: Atenolol (N=29 pts) vs. Florinef (N=29 pts) No placebo group	1° endpoint: 48/58 (82%) cured or improved. No difference was observed between the 2 groups.  Safety endpoint: No	Secondary Comment: 11/29 (38%) o Atenolol had an adverse event. (depression, suicide ideation, headaches)  Summary: Oral treatment of neurally mediated syncope with Florinef or Atenolol is safe and efficacious.  However, a major limitation of this paper is the absence of a placebo group.
Balaji S, et al. 1994. 7906701	Aim: Outcomes of children with neurocardiogenic syncope.  Study type: Single center	Inclusion criteria: Age <20 y of age with ≥3 episode of syncope in preceding 12 mo. Structurally normal heart, normal ECG (normal QT)	Intervention: Of 100 pts positive orthostatic response, 84 were treated with fludrocortisone and NaCl.	1° endpoint: Response to medical management. Syncope present, absent, improved over a 12 mo period	Results: Of the 100 orthostatic positive responders, 84 treated with fludrocortisone and NaCl. Of these 65% complete resolution and 17% some improvement Of the 11 nonresponders 10 were treated BB

(389)	study comparing pts with positive autonomic maneuver vs. negative autonomic response.  Size: n=162 pts with syncope (12.8 y of age) compared 100 positive orthostatic response to 62 negative orthostatic response	Exclusion criteria: Other disease ruled-out by ECG, EEG, and head imaging.	Comparator: Orthostatic (autonomic abnormal) response compared to orthostatic negative response	Safety endpoint: No	and 4 responded.  Summary: Benefit to combination salt and Fludrocortisone in pts with orthostatic intolerance.  Cannot exclude placebo effect
McLeod KA, et al. 1999 <u>10573501</u> (390)	Aim: To determine whether reflex bradycardic seizures can be prevented by cardiac pacing  Study type: Randomized double blind study  Size: n=12 pts (median 2.8 y of age, mean 4 y). Duration of documented asystole (10-40 s)	Inclusion criteria: Children >2y of age, clinical Hx reflex anoxic seizures, documented asystole >4 s, reflex anoxic seizures at least 1/wk  Exclusion criteria: None	Intervention: Pacing strategy DDD, VVI, or ODO. Parent and patient blinded to PM strategy. 4 mo randomization to a different pacing protocol.  Comparator: None This study did not compare medical management vs. pacemaker therapy.	1° endpoint: Clinical recurrent syncope Safety endpoint: None	Results: Children paced either VVI or DDD significant reduction in number of syncopal events compared to a "sensing only" mode. 6 pts no further syncope when paced DDD/VVI compared to sensing only. 3 pts no further syncope regardless paced or not paced. 2 pts continued to have episodes of syncope when paced.  Summary: First blinded study demonstrating efficacy of pacing in severe neurally mediated syncope secondary to pallid breath holding spells. No control group of pts without a pacemaker. Cannot exclude placebo effect from pacemaker alone (though pts <3 y of age)  **Recommend hysteresis and rate drop features be applied

Kelly AM, et al. 2001 11533339 (391)	Aim: Determine resolution of significant bradycardia related pallid-breatholding spells with permanent pacemaker (PM) implantation	Inclusion criteria: Pallid breath-holding spells requiring PM implantation.  Exclusion criteria: None	Intervention: Pacemaker Implantation  Comparator: None	1º endpoint: Clinical Outcome Safety endpoint: None	10 pts (mean asystolic pauses 11.9 s). 5 pts had complete resolution of syncope (spells), 2 only had minor color changes without loss of consciousness, and 3 continued to have minor brief spells.
	Study type: Retrospective review				Tillion blich spelle.
	Size: n=10 pts (median PM implant at 14.5 mo)				

## Data Supplement 41. Nonrandomized Trials, Observational Studies, and/or Registries of Adult Congenital Heart Disease – (Section 10.2)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Khairy P, et al. 2004 <u>15051640</u> (392)	Study type: Retrospective Cohort Multicenter (6) Size: n=252 pts	Inclusion criteria: Programmed ventricular stimulation between 1985 and 2002  Exclusion criteria: Unrepaired TOF, pulmonary atresia, AV canal	<u>1° endpoint</u> : Composite of sustained VT or SCD <u>Results:</u> Age at EPS ≥18 y, palpitations, prior palliative surgery, Modified Lown ≥2, cardiothoracic ratio ≥0.6	Programmed ventricular stimulation is of diagnostic and prognostic value in risk stratifying pts with repaired TOF.
Khairy P, et al. 2004 <u>19808416</u> (393)	Study type: Multicenter cohort study  Size: n=37 pts	Inclusion criteria: TGA atrial baffle with ICD  Exclusion criteria: N/A	1° endpoint: Risk factors for shocks  Results: Annual rates of appropriate shocks were 0.5% and 6.0% in primary and secondary prevention, respectively (p=0.0366)	High rates of appropriate shocks are noted in secondary but not primary prevention.  Supraventricular arrhythmias may be implicated in the etiology of ventricular tachyarrhythmias; BB seem protective, and inducible VT does not seem to predict future events.

## Data Supplement 42. Nonrandomized Trials, Observational Studies, and/or Registries of Geriatrics – (Section 10.3)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Paling D, et al. 2011 22067373 (394)	Aim: To assess for CCS mediated falls in older adults (comparing those ≥80 y of age vs. 61–79 y pf age)  Study type: Prospective Observational  Size: n=101 pts with unexplained falls	Inclusion criteria: Unexplained Falls  Exclusion criteria: Pts with clear cardiac or neurological etiology of their syncope were treated as appropriate and excluded from this analysis.	1° endpoint: Combination of TT/CSM provided diagnosis in 62% of pts, and was significantly more likely to be positive in pts ≥80 y of age (68% vs. 50%, p=.001)  Safety endpoint (if relevant): N/A	Summary Diagnosis using TT/CSM in 62% pts; diagnostic sensitivity was relatively higher in those ≥80 yrs.
Cooke J, et al. 2011 21382922 (395)	Aim: To assess type of syncope wth age  Study type: Retrospective, observational  Size: n=3,002 pts	Inclusion criteria: All consecutive pts referred to a tertiary referral syncope unit over a decade were included.  Exclusion criteria: N/A	1° endpoint: Type of Syncope in relation to age.  1° Safety endpoint (if relevant): N/A	Summary:  OH was the most commonly observed abnormality (test positivity of 60.3%). Neurocardiogenic syncope demonstrated a bimodal age distribution. Of 194 pts with carotid sinus hypersensitivity, the median age (IQR) was 77 (68–82) y of age. Those with vasovagal syncope (n=80) had a median (IQR) y of age of 30 (19–44). There were 57 pts with isolated postural orthostatic tachycardia syndrome. Of the total pts, 75% were female. They had a median (IQR) y age of 23 (17–29).
Duncan GW, et al. 2010 <u>20444805</u> (396)	Aim: To clarify prevalence and character of VVS in OA  Study type: Prospective, observational  Size: n=1,060 pts	Inclusion: Pts presenting to syncope clinic Comparisons of those <60 to those ≥60  Exclusion criteria: <18 y of age	1° endpoint: Diagnosis  1° Safety endpoint (if relevant): N/A	<u>Summary:</u> Older pts even more likely than young to have VVS. The clinical presentation differed significantly between older vs. younger pts. Older pts were less likely to give a typical Hx.
Anpalaham M, et al. 2012 <u>22284256</u> (397)	Aim: To explore the relationship between falls and NMS Age 76.8±5.7 y  Study type: Proxpective Observational	Inclusion criteria: Study of consecutive admissions for falls aged ≥65 y  Exclusion criteria: those with an identifiable medical cause for the fall or a Hx of loss of	1° endpoint: 5/21 of those with nonaccidental falls had NMS  1° Safety endpoint (if relevant): N/A	Summary: Syncope underestimated in older adults as many have NMS with associated amnesia often confounding assessment

	Size n=200 nto	consciousness		
Richardson DA, et al. 1997 9080518 (398)	Size: n=200 pts  Aim: to assess for CSS- mediated syncope in pts with falls	Inclusion: Unexplained fallers age ≥50 y	1° endpoint: diagnosis of CSS with cardiac inhibition	Summary: 65/279 had cardioinhibitory carotid hypersensitivity, raising question of pacing.
, ,	Study type:  Prospective, observational  Size: n=279 pts	Exclusion criteria: (1) presented with a single simple accidental fall (simple slip or		
	Size: n=279 pts	trip); (2) presented with a readily or subsequently diagnosed medical cause; (3) were cognitively impaired (4) unable to speak English or illiterate; (5) lived outside a 15 mile radius of the RVI; (6) were immobile; or (7) were registered blind. Exclusions to CSM were: (1) MI within 3 mo; (2) stroke within 3 mo; (3) history of ventricular dysrhythmia; or (4) presence of carotid bruit		
GIS Ungar A, et al. 2006	Aim: Older adults (≥65 referred to ER) (mean age 79±7), 160 ≥75	Inclusion criteria: 65 and older with transient LOC	<u>1º endpoint</u> : Diagnosis	Summary: Definite diagnosis in 40.1%, suspected in 57.9%
<u>17038070</u> (399)	Study type: Observational	Exclusion criteria: Presyncope or cognitive impairment	1° Safety endpoint (if relevant): N/A	
	<u>Size</u> : n=231 pts			

	T			-
GIS	Aim: To study 2 y f/u of	Inclusion criteria: Pts	1° endpoint:	Summary:
Ungar A, et al.	guideline algorithm on	assessed using GIS diagnostic	Recurrent syncope and mortality	Total mortality 17.5% and syncope 32.5%;
2011	outcomes in older adults (age	algorithm		
21908471	≥60, mean 78.7±6.8)		1° Safety endpoint (if	Higher death in pts with cardiac syncope
(400)	,	Pts referred to clinic for	relevant): N/A	
,	Study type: Controlled, 2 y f/u	syncope/falls or dizziness	<u>1010 vanity</u> : 1477 t	Increased recurrence and mortality with age
				, ,
	Size: n=242 pts	Exclusion criteria:		Recurrence corresponded to age and disability
	<u> </u>	Exclusion criteria were		Tresumented seriespended to age and disability
		symptoms limited to pre-		
		syncope, severe cognitive		
		impairment, active (<5 y)		
		malignancies and disability in		
		more than 4 activities of daily		
		living		
O'Mahony, et al.	Aim: Diagnostic sensitivity of	Inclusion criteria:	<u>1° endpoint</u> :	Summary:
1998	algorithm in pts 61–91 y of age	Pts with unexplained syncope,	Diagnostic sensitivity and	High aggregate sensitivity of clinical thought process. Utility of TT
<u>9823747</u>		falls, or dizziness were	specificity	esp in context of syncopal amnesia.
(401)	Study type: Observational	referred for assessment		
			1° Safety endpoint (if	
	Size: n=54 pts	Exclusion criteria: N/A	relevant):	
Aging Clin Exp Res	Aim: To assess w/u of protocol	Inclusion criteria: Pts ≥65 with	1° endpoint: Diagnosis	Summary:
Ungar, et al.	in pts with dementia	dementia		Pts with dementia and high comorbidity, still with successful w/
2015	,	(83±6 yo) with falls or syncope.	1° Safety endpoint (if	workup
25820493	Study type:	(52% falls, 45% syncope and	relevant): N/A	r
(53)	Observational	3% overlap); 60% did not	iolovanij. WA	
(00)	222.74401141	remember episode		
	<b>Size</b> : n=296 pts	Tomombol opisode		
	<u>σίες</u> . 11–230 μισ	Exclusion criteria: Absence of		
		informed consent		

## Data Supplement 43. Nonrandomized Trials, Observational Studies, and/or Registries of Syncope in Athletes – (Section 10.5)

Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (P values; OR or RR; & 95% CI)	Summary/Conclusion Comment(s)
Maron BJ, et al. 2015 <u>19221222</u> (402)	Study type: National registry  Size: n=1,866 athletes	Inclusion criteria: Athletes who died suddenly or survived cardiac arrest; 19 y of age (+/- 6 y of age)  Exclusion criteria: N/A	1º endpoint: SCD or cardiac arrest  Results: Most common CV cause were HCM (36%) and congenital coronary artery anomalies (17%)	SCD in young US athletes was higher than previously estimated, but low nonetheless (<100 per y)
Maron BJ, et al. 2007 <u>17652294</u> (403)	Study type: Multicenter registry  Size: n= 506 pts	Inclusion criteria: ICDs implanted between 1986 and 2003  Exclusion criteria: N/A	1° endpoint: ICD intervention terminating VT or VF  Results: ICD intervention terminated VT or VF in 103 pts (20%)	ICD interventions effective in pts with HCM
Corrado, et al. 2006 <u>17018804</u> (404)	Study type: Longitudinal cohort  Size: Population based, per 100,000 person years	Inclusion criteria: Athletic and non athletic population 12–35 y of age in Veneto, Italy between 1974–2004  Exclusion criteria: N/A	1º endpoint: Incidence of CV death and cause specific CV death in screened athletes and unscreened non athletes  Results: 55 SCD in screened athletes (1.9 deaths/100,000 person-years) and 265 sudden deaths in unscreened non athletes (0.79 deaths/100,000 person-years). Incidence of SCD in athletes decreased by 89%. The incidence of SCD in unscreened nonathletic pts did not change significantly.	Incidence of SCD declined after implementation of pre participation screening program for young athletes
James CA, et al. 2013 <u>23871885</u> (405)	Study type: Longitudinal cohort  Size: n=87 pts	Inclusion criteria: Pts with desmosomal mutations  Exclusion criteria: N/A	<u>Results:</u> Compared to those who did not exercise, pts in the second (OR: 6.64 p= 0.013) third (OR: 16.7, p= 0.001) and top (OR: 25.3, p<0.001) quartiles were increasingly likely to meet Task Force Criteria for ARVC/D. Survival from first VT/VF event was lowest among those in top quatile before (p=0.036) and after (p=0.005) exercise. For pts in top quartile, a reduction in exercise decreased VT/VF risk (p=0.04)	Endurance and frequent exercise increased the risk of VT/VF, HF and ARVC/D in pts with desmosomal mutations.

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