2016 AHA/ACC Guideline on the Management of Patients With Lower Extremity Peripheral Artery Disease: Executive Summary

A Report of the American College of Cardiology/American Heart Association **Task Force on Clinical Practice Guidelines**

Developed in Collaboration With the American Association of Cardiovascular and Pulmonary Rehabilitation, Inter-Society Consensus for the Management of Peripheral Arterial Disease, Society for Cardiovascular Angiography and Interventions, Society for Clinical Vascular Surgery, Society of Interventional Radiology, Society for Vascular Medicine, Society for Vascular Nursing, Society for Vascular Surgery, and Endovascular Surgery Society

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Preamble

Since 1980, the American College of Cardiology (ACC) and American Heart Association (AHA) have translated scientific evidence into clinical practice guidelines with recommendations to improve cardiovascular health. These guidelines, based on systematic methods to evaluate and classify evidence, provide a cornerstone of quality cardiovascular care.

In response to reports from the Institute of Medicine (1, 2) and a mandate to evaluate new knowledge and maintain relevance at the point of care, the ACC/AHA Task Force on Clinical Practice Guidelines (Task Force) modified its methodology (3-5). The relationships among guidelines, data standards, appropriate use criteria, and performance measures are addressed elsewhere (5).

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 broader target. Although guidelines may be used to inform regulatory or payer decisions, the intent is to improve 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. Guidelines are reviewed annually by the Task Force and are official policy of the ACC and AHA. Each guideline is considered current until it is updated, revised, or superseded by published addenda, statements of clarification, focused updates, or revised full-text guidelines. To ensure that guidelines remain current, new data are reviewed biannually to determine whether recommendations should be modified. In general, full revisions are posted in 5-year cycles (3-6).

Modernization

Processes have evolved to support the evolution of guidelines as "living documents" that can be dynamically updated. This process delineates a recommendation to address a specific clinical question, followed by concise text (ideally <250 words) and hyperlinked to supportive evidence. This approach accommodates time constraints on busy clinicians and facilitates easier access to recommendations via electronic search engines and other evolving technology.

Evidence Review

Writing committee members review the literature; weigh the quality of evidence for or against particular tests, treatments, or procedures; and estimate expected health outcomes. In developing recommendations, the writing committee uses evidence-based methodologies that are based on all available data (3-7). 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 selected references are cited.

The Task Force recognizes the need for objective, independent Evidence Review Committees (ERCs) that include methodologists, epidemiologists, clinicians, and biostatisticians who systematically survey, abstract, and assess the evidence to address systematic review questions posed in the PICOTS format (P=population, I=intervention, C=comparator, O=outcome, T=timing, S=setting) (2, 4-6). Practical considerations, including time and resource constraints, limit the ERCs to evidence that is relevant to key clinical questions and lends itself to systematic review and analysis that could affect the strength of corresponding recommendations. Recommendations developed by the writing committee on the basis of the systematic review are marked "SR".

Guideline-Directed Management and Treatment

The term "guideline-directed management and therapy" (GDMT) refers to care defined mainly by ACC/AHA Class I recommendations. For these and all recommended drug treatment regimens, the reader should confirm dosage with product insert material and carefully evaluate for contraindications and interactions. Recommendations are limited to treatments, drugs, and devices approved for clinical use in the United States.

Class of Recommendation and Level of Evidence

The Class of Recommendation (COR; ie, the strength of the recommendation) encompasses the anticipated magnitude and certainty of benefit in proportion to risk. The Level of Evidence (LOE) rates evidence supporting the effect of the intervention on the basis of the type, quality, quantity, and consistency of data from clinical trials and other reports (Table 1) (3-5). Unless otherwise stated, recommendations are sequenced by COR and then by LOE. Where comparative data exist, preferred strategies take precedence. When >1 drug, strategy, or therapy exists within the same COR and LOE and no comparative data are available, options are listed alphabetically.

Relationships With Industry and Other Entities

The ACC and AHA sponsor the guidelines without commercial support, and members volunteer their time. The Task Force zealously avoids actual, potential, or perceived conflicts of interest that might arise through relationships with industry or other entities (RWI). All writing committee members and reviewers are required to disclose current industry relationships or personal interests, from 12 months before initiation of the writing effort. Management of RWI involves selecting a balanced writing committee and assuring that the chair and a majority of committee members have no relevant RWI (Appendix 1). Members are restricted with regard to writing or voting on sections to which their RWI apply. For transparency, members' comprehensive disclosure information is available online (http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIR.0000000000000471/-/DC1). Comprehensive disclosure information for the Task Force is also available at http://www.acc.org/guidelines/about-guidelines-and-clinical-documents/guidelines-and-documents-task-forces.

The Task Force strives to avoid bias by selecting experts from a broad array of backgrounds representing different geographic regions, sexes, ethnicities, intellectual perspectives/biases, and scopes of clinical practice, and by inviting organizations and professional societies with related interests and expertise to participate as partners or collaborators.

Individualizing Care in Patients With Associated Conditions and Comorbidities

Managing patients with multiple conditions can be complex, especially when recommendations applicable to coexisting illnesses are discordant or interacting (8). The guidelines are intended to define practices meeting the needs of patients in most, but not all, circumstances. The recommendations should not replace clinical judgment.

Clinical Implementation

Management in accordance with guideline recommendations is effective only when followed. Adherence to recommendations can be enhanced by shared decision making between clinicians and patients, with patient engagement in selecting interventions on the basis of individual values, preferences, and associated conditions and comorbidities. Consequently, circumstances may arise in which deviations from these guidelines are appropriate.

The reader is encouraged to consult the full-text guideline (9) for additional guidance and details with regard to lower extremity peripheral artery disease (PAD) because the executive summary contains limited information.

Jonathan L. Halperin, MD, FACC, FAHA Chair, ACC/AHA Task Force on Clinical Practice Guideline

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 RECOMMENDA	TION
CLASS I (STRONG) B	enefit >>> Ris
 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/india preference to treatment B Treatment A should be chosen over treatment 	
CLASS IIa (MODERATE)	Benefit >> Ris
 Suggested phrases for writing recommendations: Is reasonable Can be useful/effective/beneficial Comparative-Effectiveness Phrases†: Treatment/strategy A is probably recommend preference to treatment B It is reasonable to choose treatment A over treatment B 	led/indicated i
CLASS IIb (WEAK)	Benefit \geq Ris
Suggested phrases for writing recommendations: May/might be reasonable May/might be considered Usefulness/effectiveness is unknown/unclear/mor not well established	uncertain
CLASS III: No Benefit (MODERATE) (Generally, LOE A or B use only)	Benefit = Ris
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 > Benef
Suggested phrases for writing recommendations: Potentially harmful Causes harm Associated with excess morbidity/mortality	

- 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

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

LEVEL B-NR

- 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)

(Nonrandomized)

- 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

I FVFL C-FO

(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 January through September 2015. Key search words included but were not limited to the following: *acute limb ischemia, angioplasty, ankle-brachial index, anticoagulation, antiplatelet therapy, atypical leg symptoms, blood pressure lowering/hypertension, bypass graft/bypass grafting/surgical bypass, cilostazol,*

claudication/intermittent claudication, critical limb ischemia/severe limb ischemia, diabetes, diagnostic testing, endovascular therapy, exercise rehabilitation/exercise therapy/exercise training/supervised exercise, lower extremity/foot wound/ulcer, peripheral artery disease/peripheral arterial disease/peripheral vascular disease/lower extremity arterial disease, smoking/smoking cessation, statin, stenting, and vascular surgery.

Additional relevant studies published through September 2016, during the guideline writing process, were also considered by the writing committee, and added to the evidence tables when appropriate. The final evidence tables included in the Online Data Supplement

(<u>http://circ.ahajournals.org/lookup/suppl/doi:10.1161/CIR.000000000000470/-/DC2</u>) summarize the evidence utilized by the writing committee to formulate recommendations. Additionally, the writing committee reviewed documents related to lower extremity PAD previously published by the ACC and AHA (10, 11). References selected and published in this document are representative and not all-inclusive.

As stated in the Preamble, the ACC/AHA guideline methodology provides for commissioning an independent ERC to address systematic review questions (PICOTS format) to inform recommendations developed by the writing committee. All other guideline recommendations (not based on the systematic review questions) were also subjected to an extensive evidence review process. For this guideline, the writing committee in conjunction with the Task Force and ERC Chair identified the following systematic review questions: 1) Is antiplatelet therapy beneficial for prevention of cardiovascular events in the patient with symptomatic or asymptomatic lower extremity PAD? 2) What is the effect of revascularization, compared with optimal medical therapy and exercise training, on functional outcome and quality of life (QoL) among patients with claudication? Each question has been the subject of recently published, systematic evidence reviews (12-14). The quality of these evidence reviews was appraised by the ACC/AHA methodologist and a vendor contracted to support this process (Doctor Evidence [Santa Monica, CA]). Few substantive randomized or nonrandomized studies had been published after the end date of the literature searches used for the existing evidence reviews, so the ERC concluded that no additional systematic review was necessary to address either of these critical questions.

A third systematic review question was then identified: 3) Is one revascularization strategy (endovascular or surgical) associated with improved cardiovascular and limb-related outcomes in patients with critical limb ischemia (CLI)? This question had also been the subject of a high-quality systematic review that synthesized evidence from observational data and an RCT (15); additional RCTs addressing this question are ongoing (16-18). The writing committee and the Task Force decided to expand the survey to include more relevant randomized and observational studies. Based on evaluation of this additional evidence the ERC decided that further systematic review was not needed to inform the writing committee on this question. Hence, the ERC and writing committee concluded that available systematic reviews could be used to inform the development of recommendations addressing each of the 3 systematic review questions specified above. The members of the Task Force and writing committee thank the members of the ERC that began this process and their willingness to participate in this volunteer effort. They include Aruna Pradhan, MD, MPH (ERC Chair); Natalie Evans, MD; Peter Henke, MD; Dharam J. Kumbhani, MD, SM, FACC; and Tamar Polonsky, MD.

1.2. Organization of the Writing Committee

The writing committee consisted of clinicians, including noninvasive and interventional cardiologists, exercise physiologists, internists, interventional radiologists, vascular nurses, vascular medicine specialists, and vascular surgeons, as well as clinical researchers in the field of vascular disease, a nurse (in the role of patient representative), and members with experience in epidemiology and/or health services research. The writing committee included representatives from the ACC and AHA, American Association of Cardiovascular and Pulmonary Rehabilitation, Inter-Society Consensus for the Management of Peripheral Arterial Disease, Society for Cardiovascular Angiography and Interventions, Society for Clinical Vascular Surgery, Society of Interventional Radiology, Society for Vascular Medicine, Society for Vascular Nursing, Society for Vascular Surgery, and Vascular and Endovascular Surgery Society.

1.3. Document Review and Approval

This document was reviewed by 2 official reviewers nominated by the ACC and AHA; 1 to 2 reviewers each from the American Association of Cardiovascular and Pulmonary Rehabilitation, Inter-Society Consensus for the Management of Peripheral Arterial Disease, Society for Cardiovascular Angiography and Interventions, Society for Clinical Vascular Surgery, Society of Interventional Radiology, Society for Vascular Medicine, Society for Vascular Nursing, Society for Vascular Surgery, and Vascular and Endovascular Surgery Society; and 16 additional 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 and the AHA and endorsed by the American Association of Cardiovascular and Pulmonary Rehabilitation, Inter-Society Consensus for the Management of Peripheral Arterial Disease, Society for Cardiovascular Angiography and Interventions, Society for Clinical Vascular Surgery, Society of Interventional Radiology, Society for Vascular

Medicine, Society for Vascular Nursing, Society for Vascular Surgery, and Vascular and Endovascular Surgery Society.

1.4. Scope of Guideline

Lower extremity PAD is a common cardiovascular disease that is estimated to affect approximately 8.5 million Americans above the age of 40 years and is associated with significant morbidity, mortality, and QoL impairment (19). It has been estimated that 202 million people worldwide have PAD (20). The purpose of this document is to provide a contemporary guideline for diagnosis and management of patients with lower extremity PAD. This document supersedes recommendations related to lower extremity PAD in the "ACC/AHA 2005 Guidelines for the Management of Patients With Peripheral Arterial Disease" (10) and the "2011 ACCF/AHA Focused Update of the Guideline for the Management of Patients With Peripheral Artery Disease" (11). The scope of this guideline is limited to atherosclerotic disease of the lower extremity arteries (PAD) and includes disease of the aortoiliac, femoropopliteal, and infrapopliteal arterial segments. It does not address nonatherosclerotic causes of lower extremity arterial disease, such as vasculitis, fibromuscular dysplasia, physiological entrapment syndromes, cystic adventitial disease, and other entities. Future guidelines will address aneurysmal disease of the abdominal aorta and lower extremity arteries and diseases of the renal and mesenteric arteries.

For the purposes of this guideline, key terms associated with PAD are defined in Table 2.

Table 2. Definition of PAD K	ey Terms
Term	Definition
Claudication	Fatigue, discomfort, cramping, or pain of vascular origin in the muscles of the lower extremities that is consistently induced by exercise and consistently relieved by rest (within 10 min).
Acute limb ischemia (ALI)	 Acute (<2 wk), severe hypoperfusion of the limb characterized by these features: pain, pallor, pulselessness, poikilothermia (cold), paresthesias, and paralysis. One of these categories of ALI is assigned (Section 10): Viable—Limb is not immediately threatened; no sensory loss; no muscle weakness; audible arterial and venous Doppler. Threatened—Mild-to-moderate sensory or motor loss; inaudible arterial Doppler; audible venous Doppler; may be further divided into IIa (marginally threatened) or IIb (immediately threatened). III. Irreversible—Major tissue loss or permanent nerve damage inevitable; profound sensory loss, anesthetic; profound muscle weakness or paralysis (rigor); inaudible arterial and venous Doppler (21, 22).
Tissue loss	 Type of tissue loss: Minor—nonhealing ulcer, focal gangrene with diffuse pedal ischemia. Major—extending above transmetatarsal level; functional foot no longer salvageable (21).
Critical limb ischemia (CLI)	 A condition characterized by chronic (≥2 wk) ischemic rest pain, nonhealing wound/ulcers, or gangrene in 1 or both legs attributable to objectively proven arterial occlusive disease. The diagnosis of CLI is a constellation of both symptoms and signs. Arterial disease can be proved objectively with ABI, TBI, TcPO₂, or skin perfusion pressure. Supplementary parameters, such as absolute ankle and toe pressures and pulse volume recordings, may also be used to assess for significant arterial

Table 2. Definition of PAD Key Term

Term	Definition
	occlusive disease. However, a very low ABI or TBI does not necessarily mean
	the patient has CLI. The term CLI implies chronicity and is to be distinguished
	from ALI (23).
In-line blood flow	Direct arterial flow to the foot, excluding collaterals.
Functional status	Patient's ability to perform normal daily activities required to meet basic needs,
	fulfill usual roles, and maintain health and well-being. Walking ability is a
	component of functional status.
Nonviable limb	Condition of extremity (or portion of extremity) in which loss of motor function,
~	neurological function, and tissue integrity cannot be restored with treatment.
Salvageable limb	Condition of extremity with potential to secure viability and preserve motor function
Standard I and a standard standard	to the weight-bearing portion of the foot if treated.
Structured exercise program	Planned program that provides individualized recommendations for type, frequency, intensity, and duration of exercise.
	 Program provides recommendations for exercise progression to assure that the
	body is consistently challenged to increase exercise intensity and levels as
	functional status improves over time.
	 There are 2 types of structured exercise program for patients with PAD:
	1. Supervised exercise program
	 Structured community- or home-based exercise program
Supervised exercise program	Structured exercise program that takes place in a hospital or outpatient facility in
	which intermittent walking exercise is used as the treatment modality.
	 Program can be standalone or can be made available within a cardiac ociation.
	rehabilitation program.
	 Program is directly supervised by qualified healthcare provider(s).
	• Training is performed for a minimum of 30 to 45 min per session, in sessions
	performed at least 3 times/wk for a minimum of 12 wk (24-34). Patients may
	not initially achieve these targets, and a treatment goal is to progress to these
	levels over time.
	• Training involves intermittent bouts of walking to moderate-to-maximum
	claudication, alternating with periods of rest.
Structured community- or	• Warm-up and cool-down periods precede and follow each session of walking. Structured exercise program that takes place in the personal setting of the patient
home-based exercise program	rather than in a clinical setting (29, 35-39).
nome-based excicise program	 Program is self-directed with the guidance of healthcare providers who
	prescribe an exercise regimen similar to that of a supervised program.
	 Patient counseling ensures that patients understand how to begin the program,
	how to maintain the program, and how to progress the difficulty of the walking
	(by increasing distance or speed).
	• Program may incorporate behavioral change techniques, such as health
	coaching and/or use of activity monitors.
Emergency versus urgent	• An <i>emergency</i> procedure is one in which life or limb is threatened if the patient
	is not in the operating room or interventional suite and/or where there is time for
	no or very limited clinical evaluation, typically within <6 h.
	• An <i>urgent</i> procedure is one in which there may be time for a limited clinical
	evaluation, usually when life or limb is threatened if the patient is not in the
Testendiaria linear	operating room or interventional suite, typically between 6 and 24 h.
Interdisciplinary care team	A team of professionals representing different disciplines to assist in the evaluation and management of the national with PAD
	and management of the patient with PAD.For the care of patients with CLI, the interdisciplinary care team should include
	• For the care of patients with CLI, the interdisciplinary care team should include individuals who are skilled in endovascular revascularization, surgical
	revascularization, wound healing therapies and foot surgery, and medical
	evaluation and care.
	 Interdisciplinary care team members may include:
	• Vascular medical and surgical specialists (ie, vascular medicine,
	vascular surgery, interventional radiology, interventional
	cardiology)

Term	Definition					
	0 Nurses					
	 Orthopedic surgeons and podiatrists 					
	 Endocrinologists 					
	 Internal medicine specialists 					
	 Infectious disease specialists 					
	 Radiology and vascular imaging specialists 					
	 Physical medicine and rehabilitation clinicians 					
	 Orthotics and prosthetics specialists 					
	 Social workers 					
	 Exercise physiologists 					
	 Physical and occupational therapists 					
	 Nutritionists/dieticians 					
Cardiovascular ischemic events	Acute coronary syndrome (acute MI, unstable angina), stroke, or cardiovascular					
	death.					
Limb-related events	Worsening claudication, new CLI, new lower extremity revascularization, or new					
	ischemic amputation.					

ABI indicates ankle-brachial index; ALI, acute limb ischemia; CLI, critical limb ischemia; MI, myocardial infarction; PAD, peripheral artery disease; TBI, toe-brachial index; and TcPO₂, transcutaneous oxygen pressure.

2. Clinical Assessment for PAD

Evaluating the patient at increased risk of PAD (Table 3) begins with the clinical history, review of systems, and physical examination. The symptoms and signs of PAD are variable. Patients with PAD may experience the classic symptom of claudication or may present with advanced disease, including CLI. Studies have demonstrated that the majority of patients with confirmed PAD do not have typical claudication but have other non–joint-related limb symptoms (atypical leg symptoms) or are asymptomatic (40, 41). Patients with PAD who have atypical leg symptoms or no symptoms may have functional impairment comparable to patients with claudication (42). The vascular examination for PAD includes pulse palpation, auscultation for femoral bruits, and inspection of the legs and feet. Lower extremity pulses are assessed and rated as follows: 0, absent; 1, diminished; 2, normal; or 3, bounding. See Table 4 for history and physical examination findings suggestive of PAD. To confirm the diagnosis of PAD, abnormal physical examination findings must be confirmed with diagnostic testing (Section 3), generally with the ankle-brachial index (ABI) as the initial test.

Patients with confirmed diagnosis of PAD are at increased risk for subclavian artery stenosis (43-45). An inter-arm blood pressure difference of >15 to 20 mm Hg is abnormal and suggestive of subclavian (or innominate) artery stenosis. Measuring blood pressure in both arms identifies the arm with the highest systolic pressure, a requirement for accurate measurement of the ABI (46). Identification of unequal blood pressures in the arms also allows for more accurate measurement of blood pressure in the treatment of hypertension (ie, blood pressure is taken at the arm with higher measurements).

See Online Data Supplements 1 and 2 for data supporting Section 2.

2.1. History and Physical Examination: Recommendations

Recommendations for History and Physical Examination					
COR	LOE	Recommendations			
I	B-NR	Patients at increased risk of PAD (Table 3) should undergo a comprehensive medical history and a review of symptoms to assess for exertional leg symptoms, including claudication or other walking impairment, ischemic rest pain, and nonhealing wounds (40-42, 47-49).			
I	B-NR	Patients at increased risk of PAD (Table 4) should undergo vascular examination, including palpation of lower extremity pulses (ie, femoral, popliteal, dorsalis pedis, and posterior tibial), auscultation for femoral bruits, and inspection of the legs and feet (48, 50, 51).			
Ι	B-NR	Patients with PAD should undergo noninvasive blood pressure measurement in both arms at least once during the initial assessment (43- 45).			

Table 3. Patients at Increased Risk of PAD

- Age ≥65 y
- Age 50–64 y, with risk factors for atherosclerosis (eg, diabetes mellitus, history of smoking, hyperlipidemia, hypertension) or family history of PAD (52)
- Age <50 y, with diabetes mellitus and 1 additional risk factor for atherosclerosis
- Individuals with known atherosclerotic disease in another vascular bed (eg, coronary, carotid, subclavian, renal, mesenteric artery stenosis, or AAA)

AAA indicates abdominal aortic aneurysm; PAD, peripheral artery disease.

Table 4. History and/or Physical Examination Findings Suggestive of PAD

- Claudication
 - Other non-joint-related exertional lower extremity symptoms (not typical of claudication)
 - Impaired walking function
 - Ischemic rest pain

Physical Examination

- Abnormal lower extremity pulse examination
- Vascular bruit
- Nonhealing lower extremity wound
- Lower extremity gangrene
- Other suggestive lower extremity physical findings (eg, elevation pallor/dependent rubor)

PAD indicates peripheral artery disease.

3. Diagnostic Testing for the Patient With Suspected Lower Extremity PAD (Claudication or CLI): Recommendations

History or physical examination findings suggestive of PAD need to be confirmed with diagnostic testing. The resting ABI is the initial diagnostic test for PAD and may be the only test required to establish the diagnosis and institute GDMT. The resting ABI is a simple, noninvasive test that is obtained by measuring systolic blood

pressures at the arms (brachial arteries) and ankles (dorsalis pedis and posterior tibial arteries) in the supine position by using a Doppler device. The ABI of each leg is calculated by dividing the higher of the dorsalis pedis pressure or posterior tibial pressure by the higher of the right or left arm blood pressure (46). Segmental lower extremity blood pressures and Doppler or plethysmographic waveforms (pulse volume recordings) are often performed along with the ABI and can be used to localize anatomic segments of disease (eg, aortoiliac, femoropopliteal, infrapopliteal) (22, 53, 54).

Depending on the clinical presentation (eg, claudication or CLI) and the resting ABI values, additional physiological testing studies may be indicated, including exercise treadmill ABI testing, measurement of the toebrachial index (TBI), and additional perfusion assessment measures (eg, transcutaneous oxygen pressure [TcPO₂], or skin perfusion pressure [SPP]). Exercise treadmill ABI testing is important to objectively measure functional limitations attributable to leg symptoms and is useful in establishing the diagnosis of lower extremity PAD in the symptomatic patient when resting ABIs are normal or borderline (54-59). The TBI is used to establish the diagnosis of PAD in the setting of noncompressible arteries (ABI >1.40) and may also be used to assess perfusion in patients with suspected CLI. Studies for anatomic imaging assessment (duplex ultrasound, computed tomography angiography [CTA], or magnetic resonance angiography [MRA], invasive angiography) are generally reserved for highly symptomatic patients in whom revascularization is being considered. Depending on the modality, these studies may confer procedural risk.

See Table 5 for alternative causes of leg pain in the patient with normal ABI and physiological testing; Figure 1 for the algorithm on diagnostic testing for suspected PAD and claudication; Table 6 for alternative causes of nonhealing wounds in patients without PAD; Figure 2 for the algorithm on diagnostic testing for suspected CLI; and Online Data Supplements 3 to 7 for data supporting Section 3.

Recommendations for Resting ABI for Diagnosing PAD					
COR	LOE	Recommendations			
I	B-NR	In patients with history or physical examination findings suggestive of PAD (Table 4), the resting ABI, with or without segmental pressures and waveforms, is recommended to establish the diagnosis (60-65).			
I	C-LD	Resting ABI results should be reported as abnormal (ABI ≤0.90), borderline (ABI 0.91–0.99), normal (1.00–1.40), or noncompressible (ABI >1.40) (46, 63-66).			
IIa	B-NR	In patients at increased risk of PAD (Table 3) but without history or physical examination findings suggestive of PAD (Table 4), measurement of the resting ABI is reasonable (41, 42, 67-89).			
III: No Benefit	B-NR	In patients not at increased risk of PAD (Table 3) and without history or physical examination findings suggestive of PAD (Table 4), the ABI is not recommended (87, 90).			

3.1. Resting ABI for Diagnosing PAD

3.2. Physiological Testing

Recommendations for Physiological Testing				
COR	LOE	Recommendations		
Т	B-NR	Toe-brachial index (TBI) should be measured to diagnose patients with		
1	D-INK	suspected PAD when the ABI is greater than 1.40 (66, 91-94).		
		Patients with exertional non-joint-related leg symptoms and normal or		
I	B-NR	borderline resting ABI (>0.90 and ≤1.40) should undergo exercise treadmill		
		ABI testing to evaluate for PAD (54-59).		
		In patients with PAD and an abnormal resting ABI (≤0.90), exercise		
Ha	B-NR	treadmill ABI testing can be useful to objectively assess functional status		
		(54-59).		
	B-NR	In patients with normal (1.00–1.40) or borderline (0.91–0.99) ABI in the		
Ha		setting of nonhealing wounds or gangrene, it is reasonable to diagnose CLI		
		by using TBI with waveforms, TcPO ₂ , or SPP (95-99).		
		In patients with PAD with an abnormal ABI (≤0.90) or with		
IIa	B-NR	noncompressible arteries (ABI >1.40 and TBI ≤0.70) in the setting of		
	D-IVK	nonhealing wounds or gangrene, TBI with waveforms, TcPO2, or SPP can		
		be useful to evaluate local perfusion (95-99).		

Circulation

PAD-Related)						
Condition	Location	Characteristic	Effect of Exercise	Effect of Rest	Effect of Position	Other Characteristics
Symptomatic Baker's cyst	Behind knee, down calf	Swelling, tenderness	With exercise	Also present at rest	None	Not intermittent
Venous claudication	Entire leg, worse in calf	Tight, bursting pain	After walking.	Subsides slowly	Relief speeded by elevation	History of iliofemoral deep vein thrombosis; edema; signs of venous stasis
Chronic compartment syndrome	Calf muscles	Tight, bursting pain	After much exercise (jogging)	Subsides very slowly	Relief with rest	Typically heavy muscled athletes
Spinal stenosis	Often bilateral buttocks, posterior leg	Pain and weakness	May mimic claudication	Variable relief but can take a long time to recover	Relief by lumbar spine flexion	Worse with standing and extending spine
Nerve root compression	Radiates down leg	Sharp lancinating pain	Induced by sitting, standing, or walking	Often present at rest	Improved by change in position	History of back problems; worse with sitting; relief when supine or

After variable

After variable

degree of

exercise

degree of

exercise

Not quickly

Not quickly

relieved

relieved

Improved

when not

May be

weight bearing

relieved by not

bearing weight

 Table 5. Alternative Diagnoses for Leg Pain or Claudication With Normal Physiological Testing (Not PAD-Related)

Modified from Norgren L, et al. (23).

Hip arthritis

Foot/ankle

arthritis

PAD indicates peripheral artery disease.

Lateral hip,

thigh

Ankle,

foot, arch

Aching

discomfort

Aching pain

sitting

Symptoms

variable;

history of degenerative arthritis

Symptoms

rest

variable; may be

related to activity level or present at

Condition	Location	Characteristics and Causes
Venous ulcer	Distal leg, especially above medial mellolus	 Develops in regions of skin changes due to chronic venous disease and local venous hypertension Typically wet (ie, wound drainage) rather than dry lesion
Distal small arterial occlusion (microangiopathy)	Toes, foot, leg	 Diabetic microangiopathy End-stage renal disease Thromboangiitis obliterans (Buerger's) Sickle-cell anemia Vasculitis (eg, Churg-Strauss, Henoch-Schonlein purpura, leukocytoclastic vasculitis, microscopic polyangiitis, polyarteritis nodosa) Scleroderma Cryoagglutination Embolic (eg, cholesterol emboli, thromboemboli, endocarditis) Thrombotic (eg, antiphospholipid antibody syndrome, Sneddon's syndrome, warfarin skin necrosis, disseminated intravascular coagulation, livedoid vasculitis, protein C or S deficiency, prolonged vasospasm)
Local injury	Toes, foot, leg	Trauma Insect or animal bite Burn
Medication related	Toes, foot, leg	 Drug reactions (eg, erythema multiforme) Medication direct toxicity (eg, doxorubicin, hydroxyurea, some tyrosine kinase inhibitors)
Neuropathic	Pressure zones of foot	 Hyperkeratosis surrounds the ulcer Diabetes mellitus with peripheral neuropathy Peripheral neuropathy without diabetes mellitus Leprosy
Autoimmune injury	Toes, foot, leg	 With blisters (eg, pemphigoid, pemphigus, epidermolysis bullosa) Without blisters (eg, dermatomyositis, lupus, scleroderma)
Infection	Toes, foot, leg	 Bacterial (eg, pseudomonas, necrotizing streptococcus) Fungal (eg, blastomycosis, Madura foot, chromomycosis) Mycobacterial Parasitic (eg, Chagas, leishmaniasis) Viral (eg, herpes)
Malignancy	Toes, foot, leg	 Primary skin malignancy Metastatic malignancy Malignant transformation of ulcer
Inflammatory	Toes, foot, leg	 Necrobiosis lipoidica Pyoderma gangrenosum Granuloma annulare

 Table 6. Alternative Diagnoses for Nonhealing Wounds With Normal Physiological Testing (Not PAD-Related)

PAD indicates peripheral artery disease.

Figure 1. Diagnostic Testing for Suspected PAD



Colors correspond to Class of Recommendation in Table 1.

ABI indicates ankle-brachial index; CLI, critical limb ischemia; CTA, computed tomography angiography; GDMT, guideline-directed management and therapy; MRA, magnetic resonance angiography; PAD, peripheral artery disease; and TBI, toe-brachial index.

Figure 2. Diagnostic Testing for Suspected CLI



Colors correspond to Class of Recommendation in Table 1.

*Order based on expert consensus.

†TBI with waveforms, if not already performed.

ABI indicates ankle-brachial index; CLI, critical limb ischemia; CTA, computed tomography angiography; MRA, magnetic resonance angiography; TcPO₂, transcutaneous oxygen pressure; and TBI, toe-brachial index.

3.3. Imaging for Anatomic Assessment

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Recommendations for Imaging for Anatomic Assessment				
COR	LOE	Recommendations		
I	B-NR	Duplex ultrasound, CTA, or MRA of the lower extremities is useful to diagnose anatomic location and severity of stenosis for patients with symptomatic PAD in whom revascularization is considered (100-103).		
I	C-EO	Invasive angiography is useful for patients with CLI in whom revascularization is considered.		
IIa	C-EO	Invasive angiography is reasonable for patients with lifestyle-limiting claudication with an inadequate response to GDMT for whom revascularization is considered.		
III: Harm	B-R	Invasive and noninvasive angiography (ie, CTA, MRA) should not be performed for the anatomic assessment of patients with asymptomatic PAD (104-106).		

4. Screening for Atherosclerotic Disease in Other Vascular Beds for the Patient With PAD: Recommendations

See Online Data Supplement 8 for data supporting Section 4.

4.1. Abdominal Aortic Aneurysm

PAD has been recognized as a risk factor for abdominal aortic aneurysm (AAA). In observational studies, the prevalence of AAA (aortic diameter \geq 3 cm) was higher in patients with symptomatic PAD than in the general population (107, 108) and in a population of patients with atherosclerotic risk factors (109). The prevalence of AAA among patients with PAD increased with age, beginning in patients \geq 55 years of age, and was highest in patients \geq 75 years of age (107). There are no data on AAA screening in patients with asymptomatic PAD. This section refers to screening patients with symptomatic PAD for AAA. Recommendations for screening the general population with risk factors for AAA (based on age, sex, smoking history, and family history) have been previously published (10).

Recommendation for Abdominal Aortic Aneurysm		
COR	LOE	Recommendation
Ha	B-NR	A screening duplex ultrasound for AAA is reasonable in patients with symptomatic PAD (107-109).

4.2. Screening for Asymptomatic Atherosclerosis in Other Arterial Beds (Coronary, Carotid, and Renal Arteries)

The prevalence of atherosclerosis in the coronary, carotid, and renal arteries is higher in patients with PAD than in those without PAD (109-115). However, intensive atherosclerosis risk factor modification in patients with PAD is justified regardless of the presence of disease in other arterial beds. Thus, the only justification for

screening for disease in other arterial beds is if revascularization results in a reduced risk of myocardial infarction (MI), stroke, or death, and this has never been shown. Currently, there is no evidence to demonstrate that screening all patients with PAD for asymptomatic atherosclerosis in other arterial beds improves clinical outcome. Intensive treatment of risk factors through GDMT is the principle method for preventing adverse cardiovascular ischemic events from asymptomatic disease in other arterial beds.

5. Medical Therapy for the Patient With PAD: Recommendations

Patients with PAD should receive a comprehensive program of GDMT, including structured exercise and lifestyle modification, to reduce cardiovascular ischemic events and improve functional status. Smoking cessation is a vital component of care for patients with PAD who continue to smoke. A guideline-based program of pharmacotherapy to reduce cardiovascular ischemic events and limb-related events should be prescribed for each patient with PAD and is customized to individual risk factors, such as whether the patient also has diabetes mellitus. Pharmacotherapy for the patient with PAD includes antiplatelet and statin agents and is customized to additional risk factors, such as whether the patient also has diabetes mellitus or hypertension. Previous studies have demonstrated that patients with PAD are less likely to receive GDMT than patients with other forms of cardiovascular disease, including coronary artery disease (116-118). Cilostazol is an effective medical therapy for treatment of leg symptoms and walking impairment due to claudication (119). However, side effects include headache, diarrhea, dizziness, and palpitations and in 1 trial, 20% of patients discontinued cilostazol within 3 months (120).

See Online Data Supplements 13 to 19 for data supporting Section 5.

Recomme	Recommendations for Antiplatelet, Statin, and Antihypertensive Agents		
COR	LOE	Recommendations	
Antiplatel	et Agents		
I	Α	Antiplatelet therapy with aspirin alone (range 75–325 mg per day) or clopidogrel alone (75 mg per day) is recommended to reduce MI, stroke, and vascular death in patients with symptomatic PAD (121-124).	
Ha	C-EO	In asymptomatic patients with PAD (ABI ≤0.90), antiplatelet therapy is reasonable to reduce the risk of MI, stroke, or vascular death.	
IIb	B-R	In asymptomatic patients with borderline ABI (0.91–0.99), the usefulness of antiplatelet therapy to reduce the risk of MI, stroke, or vascular death is uncertain (67, 68, 121, 124).	
IIb	B-R	The effectiveness of dual-antiplatelet therapy (aspirin and clopidogrel) to reduce the risk of cardiovascular ischemic events in patients with symptomatic PAD is not well established (125, 126).	
IIb	C-LD	Dual-antiplatelet therapy (aspirin and clopidogrel) may be reasonable to reduce the risk of limb-related events in patients with symptomatic PAD after lower extremity revascularization (127-130).	
IIb	B-R	The overall clinical benefit of vorapaxar added to existing antiplatelet therapy in patients with symptomatic PAD is uncertain (131-134).	

5.1. Antiplatelet, Statin, Antihypertensive Agents, and Oral Anticoagulation

Statin Age	Statin Agents			
Ι	Α	Treatment with a statin medication is indicated for all patients with PAD (88, 135-139).		
Antihyper	tensive Age	nts		
I	А	Antihypertensive therapy should be administered to patients with hypertension and PAD to reduce the risk of MI, stroke, heart failure, and cardiovascular death (140-144).		
IIa	A	The use of angiotensin-converting enzyme inhibitors or angiotensin-receptor blockers can be effective to reduce the risk of cardiovascular ischemic events in patients with PAD (143, 145, 146).		
Oral Anticoagulation				
IIb	B-R	The usefulness of anticoagulation to improve patency after lower extremity autogenous vein or prosthetic bypass is uncertain (147-149).		
III: Harm	А	Anticoagulation should not be used to reduce the risk of cardiovascular ischemic events in patients with PAD (148, 150-152).		

5.2. Smoking Cessation

Recommen	Recommendations for Smoking Cessation		
COR	LOE	Recommendations	
Ι	Α	Patients with PAD who smoke cigarettes or use other forms of tobacco should be advised at every visit to quit (153-155).	
I	А	Patients with PAD who smoke cigarettes should be assisted in developing a plan for quitting that includes pharmacotherapy (ie, varenicline, bupropion, and/or nicotine replacement therapy) and/or referral to a smoking cessation program (153, 156-158).	
Ι	B-NR	Patients with PAD should avoid exposure to environmental tobacco smoke at work, at home, and in public places (159, 160).	

5.3. Glycemic Control

Recommendations for Glycemic Control		
COR	LOE	Recommendations
I	C-EO	Management of diabetes mellitus in the patient with PAD should be coordinated between members of the healthcare team.
IIa	B-NR	Glycemic control can be beneficial for patients with CLI to reduce limb-related outcomes (161, 162).

5.4. Cilostazol, Pentoxifylline, and Chelation Therapy

Recommendations for Cilostazol, Pentoxifylline, and Chelation Therapy			
COR	LOE Recommendations		
Cilostazol	Cilostazol		
Ι	Α	Cilostazol is an effective therapy to improve symptoms and increase walking distance in patients with claudication (119, 163).	
Pentoxifylline			
III: No Benefit	B-R	Pentoxifylline is not effective for treatment of claudication (119, 164).	

Chelation Therapy		
III: No	B-R	Chelation therapy (eg, ethylenediaminetetraacetic acid) is not beneficial for
Benefit		treatment of claudication (165).

5.5. Homocysteine Lowering

Recommendation for Homocysteine Lowering		
COR	LOE	Recommendation
III: No Benefit	B-R	B-complex vitamin supplementation to lower homocysteine levels for prevention of cardiovascular events in patients with PAD is not recommended (166-168).

5.6. Influenza Vaccination

Recommendation for Influenza Vaccination		
COR	LOE	Recommendation
Ι	C-EO	Patients with PAD should have an annual influenza vaccination.

6. Structured Exercise Therapy: Recommendations



Studies supporting structured community- or home-based programs for patients with PAD are more recent than studies supporting supervised exercise programs and have provided strong evidence in support of the community- or home-based approach (35, 37, 39, 80, 86, 171). Unstructured community- or home-based walking programs that consist of providing general recommendations to patients with claudication to simply walk more are not efficacious (38).

See Online Data Supplements 32 and 33 for data supporting Section 6.

Recomm	Recommendations for Structured Exercise Therapy		
COR	LOE	Recommendations	
I	Α	In patients with claudication, a supervised exercise program is recommended to improve functional status and QoL and to reduce leg symptoms (24-26, 28-34, 36, 169, 170).	
Ι	B-R	A supervised exercise program should be discussed as a treatment option for claudication before possible revascularization (24-26).	

IIa	А	In patients with PAD, a structured community- or home-based exercise program with behavioral change techniques can be beneficial to improve walking ability and functional status (37, 80, 86, 171).
IIa	A	In patients with claudication, alternative strategies of exercise therapy, including upper-body ergometry, cycling, and pain-free or low-intensity walking that avoids moderate-to-maximum claudication while walking, can be beneficial to improve walking ability and functional status (27, 173, 175, 176).

Table 7. Structured Exercise Programs for PAD: Definitions

Supervised exercise program (COR I, LOE A)

- Program takes place in a hospital or outpatient facility.
- Program uses intermittent walking exercise as the treatment modality.
- Program can be standalone or within a cardiac rehabilitation program.
- Program is directly supervised by qualified healthcare provider(s).
- Training is performed for a minimum of 30–45 min/session; sessions are performed at least 3 times/wk for a minimum of 12 wk (24-34).
- Training involves intermittent bouts of walking to moderate-to-maximum claudication, alternating with periods of rest.
- Warm-up and cool-down periods precede and follow each session of walking.

Structured community- or home-based exercise program (COR IIa, LOE A)

- Program takes place in the personal setting of the patient rather than in a clinical setting (29, 35-39).
- Program is self-directed with guidance of healthcare providers.
- Healthcare providers prescribe an exercise regimen similar to that of a supervised program.
- Patient counseling ensures understanding of how to begin and maintain the program and how to progress the difficulty of the walking (by increasing distance or speed).
- Program may incorporate behavioral change techniques, such as health coaching or use of activity monitors.
- COR indicates Class of Recommendation; LOE, Level of Evidence; and PAD, peripheral artery disease.

7. Minimizing Tissue Loss in Patients With PAD: Recommendations

Prevention of wounds through patient education, foot examination, and prompt recognition of foot infection is important to minimize tissue loss among patients with PAD. Education includes teaching patients about healthy foot behaviors (eg, daily inspection of feet, wearing of shoes and socks; avoidance of barefoot walking), the selection of proper footwear, and the importance of seeking medical attention for new foot problems (177). Educational efforts are especially important for patients with PAD who have diabetes mellitus with peripheral neuropathy.

Foot infections (infection of any of the structures distal to the malleoli) may include cellulitis, abscess, fasciitis, tenosynovitis, septic joint space infection, and osteomyelitis. Because of the consequences associated with untreated foot infection—especially in the presence of PAD—clinicians should maintain a high index of suspicion (178). Foot infection is suspected if the patient presents with local pain or tenderness; periwound erythema; periwound edema, induration, or fluctuance; pretibial edema; any discharge (especially purulent); foul odor; visible bone or a wound that probes to bone; or signs of a systemic inflammatory response (including temperature >38°C or <36°C, heart rate >90/min, respiratory rate >20/min or PaCO₂ <32 mm Hg, white blood cell count >12,000 or <4,000/mcL or >10% immature forms) (179). It is recognized that the presence of diabetes

mellitus with peripheral neuropathy and PAD may make the presentation of foot infection more subtle than in patients without these problems.

See Online Data Supplement 34 for data supporting Section 7.

Recommendations for Minimizing Tissue Loss in Patients With PAD		
COR	LOE	Recommendations
Ι	C-LD	Patients with PAD and diabetes mellitus should be counseled about self-foot examination and healthy foot behaviors (177, 180).
Ι	C-LD	In patients with PAD, prompt diagnosis and treatment of foot infection are recommended to avoid amputation (178, 179, 181-183).
Па	C-LD	In patients with PAD and signs of foot infection, prompt referral to an interdisciplinary care team (Table 8) can be beneficial (178, 184, 185).
Па	C-EO	It is reasonable to counsel patients with PAD without diabetes mellitus about self-foot examination and healthy foot behaviors.
IIa	C-EO	Biannual foot examination by a clinician is reasonable for patients with PAD and diabetes mellitus.

Table 8. Interdisciplinary Care Team for PAD

A team of professionals representing different disciplines to assist in the evaluation and management of the patient with PAD. For the care of patients with CLI, the interdisciplinary care team should include individuals who are skilled in endovascular revascularization, surgical revascularization, wound healing therapies and foot surgery, and medical evaluation and care.

Interdisciplinary care team members may include:

- Vascular medical and surgical specialists (ie, vascular medicine, vascular surgery, interventional radiology, interventional cardiology)
- Nurses
- Orthopedic surgeons and podiatrists
- Endocrinologists
- Internal medicine specialists
- Infectious disease specialists
- Radiology and vascular imaging specialists
- Physical medicine and rehabilitation clinicians
- Orthotics and prosthetics specialists
- Social workers
- Exercise physiologists
- Physical and occupational therapists
- Nutritionists/dieticians

CLI indicates critical limb ischemia; and PAD, peripheral artery disease.

8. Revascularization for Claudication: Recommendations

A minority of patients with claudication (estimated at <10% to 15% over 5 years or more) will progress to CLI (186-189). Therefore, the role of revascularization in claudication is improvement in claudication symptoms and functional status, and consequently in QoL, rather than limb salvage. Revascularization is reasonable when the patient who is being treated with GDMT (including structured exercise therapy) presents with persistent lifestyle-limiting claudication (13, 25, 26, 190, 191). Lifestyle-limiting claudication is defined by the patient rather than by any test. It includes impairment of activities of daily living and/or vocational and/or recreational

activities due to claudication. An individualized approach to revascularization for claudication is recommended for each patient to optimize outcome. Revascularization is but one component of care for the patient with claudication, inasmuch as each patient should have a customized care plan that also includes medical therapy (Section 5), structured exercise therapy (Section 6), and care to minimize tissue loss (Section 7). If a strategy of revascularization for claudication is undertaken, the revascularization strategy should be evidence based and can include endovascular revascularization, surgery, or both.

Due to the variability of ischemic limb symptoms and impact of these symptoms on functional status and QoL, patients should be selected for revascularization on the basis of severity of their symptoms. Factors to consider include a significant disability as assessed by the patient, adequacy of response to medical and structured exercise therapy, status of comorbid conditions, and a favorable risk–benefit ratio. Patient preferences and goals of care are important considerations in the evaluation for revascularization. The revascularization strategy should have a reasonable likelihood of providing durable relief of symptoms. There should be clear discussion with the patient about expected risks and benefits of revascularization, as well as discussion of the durability of proposed procedures. A general recommendation for revascularization as a treatment option for claudication is provided below followed by specific recommendations for endovascular (Section 8.1.1) and surgical (Section 8.1.2) procedures if a revascularization strategy is undertaken.

See Online Data Supplements 35 to 38 for data supporting Section 8.

8.1. Revascularization for Claudication

Recommendation for Revascularization for Claudication				
COR	LOE	Recommendation		
Па	А	Revascularization is a reasonable treatment option for the patient with lifestyle- limiting claudication with an inadequate response to GDMT (13, 25, 26, 190, 191).		

8.1.1. Endovascular Revascularization for Claudication

Endovascular techniques to treat claudication include balloon dilation (angioplasty), stents, and atherectomy. These techniques continue to involve and now include covered stents, drug-eluting stents, cutting balloons, and drug-coated balloons. The technique chosen for endovascular treatment is related to lesion characteristics (eg, anatomic location, lesion length, degree of calcification) and operator experience. Assessment of the appropriateness of specific endovascular techniques for specific lesions for the treatment of claudication is beyond the scope of this document.

Revascularization is performed on lesions that are deemed to be hemodynamically significant, and stenoses selected for endovascular treatment should have a reasonable likelihood of limiting perfusion to the distal limb. Stenoses of 50% to 75% diameter by angiography may not be hemodynamically significant, and resting or provoked intravascular pressure measurements may be used to determine whether lesions are significant (192, 193). Multiple RCTs have compared endovascular procedures to various combinations of medical treatment with or without supervised or unsupervised exercise programs (13, 25, 26, 190, 191, 194-

206). These trials have used different endpoints and enrolled patients with anatomic disease distribution at different levels. Long-term patency is greater in the aortoiliac than in the femoropopliteal segment. Furthermore, for femoropopliteal disease, durability is diminished with greater lesion length, occlusion rather than stenosis, the presence of multiple and diffuse lesions, poor-quality runoff, diabetes mellitus, chronic kidney disease, renal failure, and smoking (207-210).

Recommendations for Endovascular Revascularization for Claudication		
COR	LOE	Recommendations
I	Α	Endovascular procedures are effective as a revascularization option for patients with lifestyle-limiting claudication and hemodynamically significant aortoiliac occlusive disease (13, 25, 26, 190, 194, 196, 201).
IIa	B-R	Endovascular procedures are reasonable as a revascularization option for patients with lifestyle-limiting claudication and hemodynamically significant femoropopliteal disease (190, 197-200, 205, 206).
IIb	C-LD	The usefulness of endovascular procedures as a revascularization option for patients with claudication due to isolated infrapopliteal artery disease is unknown (211-213).
III: Harm	B-NR	Endovascular procedures should not be performed in patients with PAD solely to prevent progression to CLI (186-189, 214-216).

8.1.2. Surgical Revascularization for Claudication

Systematic reviews have concluded that surgical procedures are an effective treatment for claudication and have a positive impact on QoL and walking parameters but have identified sparse evidence supporting the effectiveness of surgery compared with other treatments (12, 191, 217, 218). Although symptom and patency outcomes for surgical interventions may be superior to those for less invasive endovascular treatments, surgical interventions are also associated with greater risk of adverse perioperative events (219-225). Treatment selection should therefore be individualized on the basis of the patient's goals, perioperative risk, and anticipated benefit. Surgical procedures for claudication are usually reserved for individuals who a) do not derive adequate benefit from nonsurgical therapy, b) have arterial anatomy favorable to obtaining a durable result with surgery, and c) have acceptable risk of perioperative adverse events. Acceptable risk is defined by the individual patient and provider on the basis of symptom severity, comorbid conditions, and appropriate GDMT risk evaluation.

The superficial femoral and proximal popliteal arteries are the most common anatomic sites of stenosis or occlusion among individuals with claudication. Femoral-popliteal bypass is therefore one of the most common surgical procedures for claudication. The type of conduit and site of popliteal artery anastomosis (above versus below knee) are major determinants of outcomes associated with femoral-popliteal bypass. Systematic reviews and meta-analyses have identified a clear and consistent primary patency benefit for autogenous vein versus prosthetic grafts for popliteal artery bypass (226, 227).

Accommendations for Surgical Acvascularization for Claudication		
COR	LOE	Recommendations
I	Α	When surgical revascularization is performed, bypass to the popliteal artery with autogenous vein is recommended in preference to prosthetic graft material (226-234).
IIa	B-NR	Surgical procedures are reasonable as a revascularization option for patients with lifestyle-limiting claudication with inadequate response to GDMT, acceptable perioperative risk, and technical factors suggesting advantages over endovascular procedures (190, 230, 235-237).
III: Harm	B-R	Femoral-tibial artery bypasses with prosthetic graft material should not be used for the treatment of claudication (238-240).
III: Harm	B-NR	Surgical procedures should not be performed in patients with PAD solely to prevent progression to CLI (186-189, 241).

Recommendations for Surgical Revascularization for Claudication

9. Management of CLI: Recommendations

Patients with CLI are at increased risk of amputation and major cardiovascular ischemic events. Care of the patient with CLI includes evaluation for revascularization and wound healing therapies, with the objective to minimize tissue loss, completely heal wounds, and preserve a functional foot. Medical therapy to prevent cardiovascular ischemic events is also an important component of care for the patient with CLI (Section 5).

See Online Data Supplements 39 and 40 for data supporting Section 9.

9.1. Revascularization for CLI

The goal of surgical or endovascular revascularization in CLI is to provide in-line blood flow to the foot through at least 1 patent artery, which will help decrease ischemic pain and allow healing of any wounds, while preserving a functional limb. The BASIL (Bypass versus Angioplasty in Severe Ischemia of the Leg) RCT (242, 243) demonstrated that endovascular revascularization is an effective option for patients with CLI as compared with open surgery. The primary endpoint of amputation-free survival was the same in the endovascular and surgical arms. Of note, the endovascular arm used only percutaneous transluminal angioplasty (242, 243). Multiple RCTs comparing contemporary surgical and endovascular treatment for patients with CLI are ongoing (17, 18, 244). Table 9 addresses factors that may prompt an endovascular versus surgical approach to the patient with CLI.

The angiosome concept has been described in the literature and entails establishing direct blood flow to the infrapopliteal artery directly responsible for perfusing the region of the leg or foot with the nonhealing wound. Multiple retrospective studies and 1 small nonrandomized prospective study assessing the efficacy of this concept have been published (245-257). Meta-analyses of these studies found improved wound healing and limb salvage with angiosome-guided therapy but cautioned that the quality of the evidence was low (258, 259). Although the angiosome concept is theoretically satisfying, randomized data comparing the establishment of inline flow versus angiosome-guided therapy have yet to be published. Furthermore, there is no evidence yet to

demonstrate the potential benefit of treating additional infrapopliteal arteries once in-line flow has been

established in one artery, regardless of angiosome.

Recomme	Recommendation for Revascularizations for CLI		
COR	LOE	Recommendation	
Ι	B-NR	In patients with CLI, revascularization should be performed when possible to minimize tissue loss (260).	
Ι	С-ЕО	An evaluation for revascularization options should be performed by an interdisciplinary care team (Table 8) before amputation in the patient with CLI.	

9.1.1. Endovascular Revascularization for CLI

Recomme	Recommendations for Endovascular Revascularization for CLI		
COR	LOE	Recommendations	
Ι	B-R	Endovascular procedures are recommended to establish in-line blood flow to the foot in patients with nonhealing wounds or gangrene (242, 243).	
Ha	C-LD	A staged approach to endovascular procedures is reasonable in patients with ischemic rest pain (261, 262).	
Ha	B-R	Evaluation of lesion characteristics can be useful in selecting the endoyascular approach for CLI (263, 264).	
IIb	B-NR	Use of angiosome-directed endovascular therapy may be reasonable for patients with CLI and nonhealing wounds or gangrene (245, 247-249, 251-253, 255-257).	

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Revascularization Findings That Favor Consideration of Surgical Revascularization	Examples
Factors associated with technical failure or poor durability with endovascular treatment	Lesion involving common femoral artery, including origin of deep femoral artery
	Long segment lesion involving the below-knee popliteal and/or infrapopliteal arteries in a patient with suitable single-segment autogenous vein conduit
	Diffuse multilevel disease that would require endovascular revascularization at multiple anatomic levels
	Small-diameter target artery proximal to site of stenosis or densely calcified lesion at location of endovascular treatment
Endovascular treatment likely to preclude or complicate subsequent achievement of in-line blood flow through surgical revascularization	Single-vessel runoff distal to ankle
Findings That Favor Consideration of Endovascular Revascularization	Examples
The presence of patient comorbidities may place patients at increased risk of perioperative complications from surgical revascularization. In these patients, an endovascular-first approach should be used regardless of anatomy	Patient comorbidities, including coronary ischemia, cardiomyopathy, congestive heart failure, severe lung disease, and chronic kidney disease
Patients with rest pain and disease at multiple levels may undergo a staged approach as part of endovascular-first approach	In-flow disease can be addressed first, and out-flow disease can be addressed in a staged manner, when required, if clinical factors or patient safety prevent addressing all diseased segments at one setting
	Some patients have had veins harvested for previous coronary

 Table 9. Therapy for CLI: Findings That Prompt Consideration of Surgical or Endovascular

 Revascularization

CLI indicates critical limb ischemia.

Recommendations for Surgical Revascularization for CLI		
COR	LOE	Recommendations
Ι	А	When surgery is performed for CLI, bypass to the popliteal or infrapopliteal arteries (ie, tibial, pedal) should be constructed with suitable autogenous vein (228, 231, 234, 265).
Ι	C-LD	Surgical procedures are recommended to establish in-line blood flow to the foot in patients with nonhealing wounds or gangrene (266-268).
IIa	B-NR	In patients with CLI for whom endovascular revascularization has failed and a suitable autogenous vein is not available, prosthetic material can be effective for bypass to the below-knee popliteal and tibial arteries (269-271).
Ha	C-LD	A staged approach to surgical procedures is reasonable in patients with ischemic rest pain (272-274).

9.1.2. Surgical Revascularization for CLI

9.2. Wound Healing Therapies for CLI

A comprehensive plan for treatment of CLI includes a plan to achieve an intact skin surface on a functional foot. The management of patients with CLI and nonhealing wounds includes coordinated efforts for both revascularization and wound healing among members of an interdisciplinary care team (Table 8). The structure and activities of interdisciplinary care teams for CLI may vary according to several factors, including the local availability of resources. Revascularization is coordinated with the efforts of clinicians who manage foot infections, provide offloading, and achieve complete wound healing, either through medical therapy, surgical options, or a combination of these options.

See Online Data Supplement 34a for a complete list of functions of the interdisciplinary care team.

Recommen	Recommendations for Wound Healing Therapies for CLI			
COR	LOE	Recommendations		
I	B-NR	An interdisciplinary care team should evaluate and provide comprehensive care for patients with CLI and tissue loss to achieve complete wound healing and a functional foot (184, 275-277).		
I	C-LD	In patients with CLI, wound care after revascularization should be performed with the goal of complete wound healing (275).		
IIb	B-NR	In patients with CLI, intermittent pneumatic compression (arterial pump) devices may be considered to augment wound healing and/or ameliorate severe ischemic rest pain (278).		
IIb	C-LD	In patients with CLI, the effectiveness of hyperbaric oxygen therapy for wound healing is unknown (279).		
III: No Benefit	B-R	Prostanoids are not indicated in patients with CLI (280).		

10. Management of Acute Limb Ischemia: Recommendations

Acute limb ischemia (ALI) is one of the most treatable and potentially devastating presentations of PAD. Timely recognition of arterial occlusion as the cause of an ischemic, cold, painful leg is crucial to successful

treatment. The writing committee has used a standard definition of ALI in which symptom duration is <2 weeks (Table 2) (21, 22). Category I refers to viable limbs that are not immediately threatened. Category II refers to threatened limbs. Category IIa limbs are marginally threatened and salvageable, if promptly treated. Category IIb are immediately threatened limbs that require immediate revascularization if salvage is to be accomplished. Category III are irreversibly damaged limbs, in which case resultant major tissue loss or permanent nerve damage is inevitable (22).

Patients with ALI should be rapidly evaluated by a vascular specialist if one is available. Depending on local clinical expertise, the vascular specialist may be a vascular surgeon, interventional radiologist, cardiologist, or a general surgeon with specialized training and experience in treating PAD. If such expertise is not locally or rapidly available, there should be strong consideration of transfer of the patient to a facility with such resources. The more advanced the degree of ischemia, the more rapidly the communication (eg, with regard to potential patient transfer) needs to occur.

ALI is a medical emergency and must be recognized rapidly. The time constraint is due to the period that skeletal muscle will tolerate ischemia—roughly 4 to 6 hours (281). A rapid assessment of limb viability and ability to restore arterial blood flow should be performed by a clinician able to either complete the Association revascularization or triage the patient (282). Lower extremity symptoms in ALI can include both pain and loss of function. The longer these symptoms are present, the less likely the possibility of limb salvage (283, 284). Clinical assessment must include symptom duration, pain intensity, and motor and sensory deficit severity to distinguish a threatened from a nonviable extremity (Figure 3). The bedside assessment includes arterial and venous examination with a handheld continuous-wave Doppler because of the inaccuracy of pulse palpation (22). The loss of Dopplerable arterial signal indicates that the limb is threatened. The absence of both arterial and venous Doppler signal indicates that the limb may be irreversibly damaged (nonsalvageable). Comorbidities should be investigated and managed aggressively, but this must not delay therapy. Even in the setting of rapid and effective revascularization, the 1-year morbidity and mortality rates ALI are high (283, 285).

See Figure 3 for the algorithm on diagnosis and management of ALI and Online Data Supplements 45 to 50 for data supporting Section 10.

Recommendations for Clinical Presentation of ALI			
COR	LOE	Recommendations	
I	С-ЕО	Patients with ALI should be emergently evaluated by a clinician with sufficient experience to assess limb viability and implement appropriate therapy.	
I	C-LD	In patients with suspected ALI, initial clinical evaluation should rapidly assess limb viability and potential for salvage and does not require imaging (282-284, 286, 287).	

10.1. Clinical Presentation of ALI

Figure 3. Diagnosis and Management of ALI (21, 22)



Colors correspond to Class of Recommendation in Table 1. ALI indicates acute limb ischemia.

10.2. Medical Therapy for ALI

Recommendation for ALI Medical Therapy			
COR	LOE	Recommendation	
Ι	C-EO	In patients with ALI, systemic anticoagulation with heparin should be administered unless contraindicated.	

10.3. Revascularization for ALI

For marginally or immediately threatened limbs (Category IIa and IIb ALI), revascularization should be performed emergently (within 6 hours). For viable limbs (Category I ALI), revascularization should be performed an on urgent basis (within 6–24 hours). The revascularization strategy can range from catheter-directed thrombolysis to surgical thromboembolectomy. Available facilities and clinical expertise are factors

that should be considered when determining the revascularization strategy. The technique that will provide the most rapid restoration of arterial flow with the least risk to the patient should be selected. For example, catheterdirected thrombolysis can provide rapid restoration of arterial flow to a viable or marginally threatened limb, particularly in the setting of recent occlusion, thrombosis of synthetic grafts, and stent thrombosis (288). If this is not available locally, surgical options for timely revascularization should be considered, along with the feasibility of timely transfer to a facility with the necessary expertise.

Prolonged duration of ischemia is the most common factor in patients requiring amputation for treatment of ALI. The risks associated with reconstruction outweigh the potential benefit in a limb that is already insensate or immobile because of prolonged ischemia. Patients who have an insensate and immobile limb in the setting of prolonged ischemia (>6 to 8 hours) are unlikely to have potential for limb salvage with revascularization.

Recomme	Recommendations for Revascularization for ALI		
COR	LOE	Recommendations	
Ι	C-LD	In patients with ALI, the revascularization strategy should be determined by local resources and patient factors (eg, etiology and degree of ischemia) (288-290).	
Ι	Α	Catheter-based thrombolysis is effective for patients with ALI and a salvageable limb (288-292).	
Ι	C-LD	Amputation should be performed as the first procedure in patients with a nonsalvageable limb (293, 294).	
Ι	C-LD	Patients with ALI should be monitored and treated (eg, fasciotomy) for compartment syndrome after revascularization (293, 294).	
Ha	B-NR	In patients with ALI with a salvageable limb, percutaneous mechanical thrombectomy can be useful as adjunctive therapy to thrombolysis (295-299).	
Ha	C-LD	In patients with ALI due to embolism and with a salvageable limb, surgical thromboembolectomy can be effective (300-302).	
IIb	C-LD	The usefulness of ultrasound-accelerated catheter-based thrombolysis for patients with ALI with a salvageable limb is unknown (303-305).	

10.4. Diagnostic Evaluation of the Cause of ALI

ALI may be related to underlying PAD (including prior lower extremity bypass graft) or may be related to other conditions that can result in ALI through either thrombotic (eg, hypercoagulable state) or embolic mechanisms. Treatment of ALI should not be delayed for testing for the underlying cause of the limb ischemia because delay from symptom onset to revascularization is a major determinant of outcome (283, 284). The evaluation of a cardiovascular (ie, embolic) cause for ALI is most useful in the patient without underlying PAD and can be completed after revascularization. Evaluation for cardiovascular cause includes electrocardiogram or additional heart rhythm monitoring to detect atrial fibrillation, electrocardiogram to detect evidence of MI, and echocardiography to further determine whether there is a cardiac etiology for thromboembolism, such as valvular vegetation, left atrial or left ventricular thrombus, or intracardiac shunt.

Recommendations for Diagnostic Evaluation of the Cause of ALI			
COR	LOE	Recommendations	
I	С-ЕО	In the patient with ALI, a comprehensive history should be obtained to determine the cause of thrombosis and/or embolization.	
Ha	C-EO	In the patient with a history of ALI, testing for a cardiovascular cause of thromboembolism can be useful.	

11. Longitudinal Follow-Up: Recommendations

PAD is a lifelong chronic medical condition. A comprehensive care plan for patients with PAD includes periodic clinical evaluation by a healthcare provider with experience in the care of vascular patients. Ongoing care focuses on cardiovascular risk reduction with medical therapy, optimizing functional status with structured exercise, and, when indicated, revascularization. The care plan is further customized depending on whether the patient has undergone a revascularization procedure.

See Online Data Supplements 51 and 52 for data supporting Section 11.

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Recommendations for Longitudinal Follow-Up		
COR	LOE	Recommendations
I	C-EO	Patients with PAD should be followed up with periodic clinical evaluation, including assessment of cardiovascular risk factors, limb symptoms, and functional status.
I	C-EO	Patients with PAD who have undergone lower extremity revascularization (surgical and/or endovascular) should be followed up with periodic clinical evaluation and ABI measurement.
Ha	B-R	Duplex ultrasound can be beneficial for routine surveillance of infrainguinal, autogenous vein bypass grafts in patients with PAD (306-312).
Па	C-LD	Duplex ultrasound is reasonable for routine surveillance after endovascular procedures in patients with PAD (313-315).
IIb	B-R	The effectiveness of duplex ultrasound for routine surveillance of infrainguinal prosthetic bypass grafts in patients with PAD is uncertain (310, 316-318).

12. Evidence Gaps and Future Research Directions

In performing the evidence review and in developing the present guidelines, the writing committee identified the following critical evidence gaps and future directions for PAD-related research:

- Basic science and translational studies to better understand the vascular biology of endovascular therapies and bypass grafting and to develop new methods for preventing restenosis after revascularization.
- Determination of risk factors for progression from asymptomatic PAD to symptomatic disease, including CLI.

- RCTs needed to determine the value of using the ABI to identify asymptomatic patients with PAD for therapies to reduce cardiovascular risk (eg, antiplatelet agents, statins, and other therapies).
- Advancement in PAD diagnostics, such as technologies for simplified yet highly accurate measurement of the ABI and tools for more reliable noninvasive perfusion assessment in CLI.
- Comparative-effectiveness studies to determine the optimal antiplatelet therapy (drug or drugs and dosage) for prevention of cardiovascular and limb-related events in patients with PAD.
- Development of additional medical therapies for claudication—an area of unmet medical need with a currently limited research pipeline (319).
- Studies to investigate the role of dietary intervention, in addition to statin therapy, to improve outcome and modify the natural history of PAD.
- Additional research to identify the best community- or home-based exercise programs for patients with PAD to maximize functional status and improve QoL, as well as the role of such exercise programs before or in addition to revascularization.
- Development and validation of improved clinical classification systems for PAD that incorporate symptoms, anatomic factors, and patient-specific risk factors and can be used to predict clinical outcome and optimize treatment approach. An example of a recently developed classification system is the Society for Vascular Surgery limb classification system, based on wound, ischemia, and foot infection (WIfI), which has been validated in different populations and may permit more meaningful prognosis in patients with CLI (320-324).
- Comparative- and cost-effectiveness studies of the different endovascular technologies for treatment of claudication and CLI, including drug-coated balloons and drug-eluting stents. Studies should include patient-centered endpoints, such as functional parameters, time to wound healing, and QoL, in addition to standard patency-focused outcomes. These studies could then be incorporated into value-based clinical algorithms for approach to revascularization for claudication and CLI.
- Additional studies to demonstrate the impact of multisocietal registries on clinical outcomes and appropriate use. At present, these include: the Vascular Quality Initiative (VQI), the National Cardiovascular Data Registry Peripheral Vascular Intervention Registry[™] (PVI Registry[™]), and the National Radiology Data Registry for Interventional Radiology (NRDR). These registries provide an opportunity to obtain "real-world" data on surgical and endovascular procedures for PAD and improve quality by providing feedback to participating centers. Future efforts should incorporate these registries into interventional RCTs and post-marketing studies of PAD-related devices.
13. Advocacy Priorities

The writing committee identified 3 priorities for multisocietal advocacy initiatives to improve health care for patients with PAD. First, the writing committee supports the availability of the ABI as the initial diagnostic test to establish the diagnosis of PAD in patients with history or physical examination findings suggestive of PAD (Table 4). Although the ABI test is generally reimbursed by third-party payers for patients with classical claudication or lower extremity wounds, payers may not provide reimbursement for the ABI with other findings suggestive of PAD, such as lower extremity pulse abnormalities or femoral bruits. The writing committee affirms the importance of confirming the diagnosis of PAD in such patients to allow for GDMT as delineated in this document. Second, the writing committee supports the vital importance of insuring access to supervised exercise programs for patients with PAD. Although extensive high-quality evidence supports supervised exercise programs to improve functional status and QoL, only a minority of patients with PAD participate in such programs because of lack of reimbursement by third-party payers. Third, the writing committee recognizes the need for incorporation of patient-centered outcomes into the process of regulatory approval of new medical therapies and revascularization technologies. For revascularization technologies, regulatory approval is driven primarily by data on angiographic efficacy (ie, target-lesion patency) and safety endpoints. The nature of the functional limitation associated with PAD warrants the incorporation of patient-centered outcomes, such as functional parameters and QoL, into the efficacy outcomes for the approval process.

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Key Words: AHA Scientific Statements, peripheral artery disease, claudication, critical limb ischemia, acute limb ischemia, antiplatelet agents, supervised exercise, endovascular procedures, bypass surgery, limb salvage, smoking cessation.

2016 AHA/ACC Lower Extremity PAD Guideline: Executive Summar	Gerhard-Herman MD, et al.
AD Guideline: Executive Summary	

Appendix 1. Author Relationships With Industry and Other Entities (Relevant)—2016 AHA/ACC Guideline on the Management of Patients With Lower Extremity Peripheral Artery Disease (March 2016)

NoneNoneNoneNoneNone4.8.1.1-None9.1.2, and10.2.2.NoneNoneNoneNoneNone5.1-5.3,NoneNone5.6, 5.10, 7,NoneNone5.6, 5.10, 7,NoneNone5.6, 5.10, 7,NoneNone4, 5.6,NoneNone4, 5.6,NoneNone4, 5.6,NoneNone81.1, 9.1.1,10.2.1 and10.2.1 and	Medtronic† No The Medicines Commany*				Associate Chief, Cardiology Director. Cardiac Catheterization	
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None None	None No	None	None	None	Harvard Medical School—	Marie D. Gerhard-
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 Boston Scientific; Medtronic; 	 Cook Medical 	None	None	 AstraZeneca Merck Novartis Plurestem 	None	None	 Boston Scientific Medrad Interventional Possis The Medicines Company 	Consultant
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None	None	None	None	Northwind†	None	None	None	Ownership/ Partnership/ Principal
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 Atrium Medical AstraZeneca† 	None	None	None	None	None As	None	None	Institutional, Organizational, or Other Financial Benefit
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4, 8.1.1– 9.1.2, and 10.2.2.	4, 8.1.1, 9.1.1 and 10.2.2.	None	None	5.1–5.3, 5.6, 5.10, and 12.	None	4, 7, 8, and10.2.2.	4, 5.6, 8.1.1, 9.1.1, 10.2.1 and 10.2.2.	Voting Recusals by Section*

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							of Nursing—Professor	Jacobson
None	None	None	None	None	None	None	University of Minnesota, School	Diane Treat-
							Exercise Physiology	
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							Bayview Medical	
							of Medicine; Johns Hopkins	
None	None	None	None	None	None	None	Johns Hopkins University, School	Kerry J. Stewart
		Benefit						
by Section*		Other Financial		Principal				
Recusals	Witness	Organizational, or	Research	Partnership/	Bureau			Member
Voting	Expert	Institutional,	Personal	Ownership/	Speakers	Consultant	Employment	Committee

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pertain to those in the full-text guideline. *Writing committee members are required to recuse themselves from voting on sections to which their specific relationships with industry and other entities may apply. Section numbers

*Significant relationship. *No financial benefit.

Medicine; SVN, Society for Vascular Nursing; SVS, Society for Vascular Surgery; TASC, Trans-Atlantic Inter-Society Consensus for the Management of Peripheral Arterial Disease; VA, cell; SCAI, Society for Cardiovascular Angiography and Interventions; SCVS, Society for Clinical Vascular Surgery, SIR, Society of Interventional Radiology, SVM, Society for Vascular Health; NHLBI, National Heart, Lung, and Blood Institute; PCORI, Patient-Centered Outcomes Research Institute; PI, primary investigator; PLX-PAD, placental-derived adherent stromal Artery Disease; FDA, U.S. Food and Drug Administration; HRS, Heart Rhythm Society; MI, myocardial infarction; NCDR, National Cardiovascular Data Registry; NIH, National Institutes of American Heart Association; AMA, American Medical Association; DSMB, data and safety monitoring board; EUCLID, Effects of Ticagrelor and Clopidogrel in Patients with Peripheral AACVPR indicates American Association of Cardiovascular and Pulmonary Rehabilitation; ACC, American College of Cardiology; ACE, Accreditation for Cardiovascular Excellence; AHA, Veterans Affairs; VESS, Vascular and Endovascular Surgery Society; and VIVA, Vascular Intervention Advances

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Patients Wit	h Lower Extremit	Patients With Lower Extremity Peripheral Artery Disease (March 2016)	Disease (March 2	016)				
Reviewer	Representation	Employment	Consultant	Speakers	Ownership/	Personal Research	Institutional,	Expert
				Bureau	Partnership/ Principal		Organizational, or Other Financial Benefit	Witness
Deepak L.	Official Reviewer-	Brigham and	 Elsevier 	None	None	 Amarin* 	 Belvoir 	None
Bhatt	ACC Board of	Women's Hospital-				 Amgen* 	Publications	
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						 PLx Pharma⁺ 	 Journal of 	
						Regado	Cardiology	
						Biosciences	(Editor)*	
						• Roche*	 Medscape 	
						• Sanon-aventis"	Cardiology	
						Tobodo*	Slack	
						The Medicines	Publications	
							(Editor)*	
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							• VA Healthcare	
							System [†]	

Appendix 2. Reviewer Relationships With Industry and Other Entities (Comprehensive)-2016 AHA/ACC Guideline on the Management of

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John S. Ikonomidis	Official Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Medical University of South Carolina— Chief	None	None	None	ine	me None None
Amy W. Pollak	Official Reviewer— AHA	Mayo Clinic— Cardiovascular Medicine Physician	None	None	None	ıe	ne None
Michael D. White	Official Reviewer—ACC Board of Governors	Catholic Health Initiatives—Chief Academic Officer	 Anthera Pharmaceuticals[†] 	None	None	ine	ne • AstraZeneca† None
Ehrin J. Armstrong	Organizational Reviewer—SVM	University of Colorado—Director, Interventional Cardiology	 Abbott Medtronic Merck Spectranetics 	None	None	ne	ne None
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reidman	Kevlewel—SCAI	Presbyterian		 Bristol-Myers Squibb† 			 The Medicines Company 	
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, al†	Sciences† • Medinal LTD† • Orexigen Therapeutics† • St. Jude Medical • Stentys†		6	C	Laboratory	C	
† ltific† (line†	 Abbott† AstraZeneca† Boston Scientific† GlaxoSmithKline† Hamilton Health 	None	None	None	Geisinger Medical Center—Staff Physician; Director, Cardiac Catheterization	Content Reviewer	James C. Blankenship
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None	None	None	None	None	Speakers Bureau
none	None	None	None	None	Ownership/ Partnership/ Principal
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<ul> <li>AHA†</li> <li>Tactile Medical*</li> </ul>	None	<ul> <li>Direct Flow Medical†</li> <li>Edwards†</li> </ul>	None	<ul> <li>ACC/AHA†</li> <li>AHA†</li> <li>ASA†</li> <li>Catheterization and Cardiovascular Intervention†</li> <li>Portland Metro Area AHA (President)† Can (President)† Can (Pres</li></ul>	Institutional, Organizational, or Other Financial Benefit
None	None	None	None	None	Expert Witness

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Employment         Constraint         Barcard Partnership Research and Policy.         Forward Francipal         Partnership Professor of Health Research and Policy.         Forward Partnership Research and Policy.         Constraint Partnership Research and Po	Reviewer	Mark A. Hlatky	Michael R. Jaff	José A. Joglar	Glenn N. Levine	Khusrow Niazi	Paul D. Varosy
Current Bureau         Principal Principal         Constraint Principal         Principal         Constraint Principal         Constraint Principal         Principal         Constraint Principal         Principal         Constraint Principal         Principal         Principal <th>Representation</th> <td>Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines</td> <td>Content Reviewer</td> <td>Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines</td> <td>Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines</td> <td>Content Reviewer—ACC Peripheral Vascular Disease Member Section</td> <td>Content Reviewer—Task Force on Performance Measures</td>	Representation	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Content Reviewer	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Content Reviewer— ACC/AHA Task Force on Clinical Practice Guidelines	Content Reviewer—ACC Peripheral Vascular Disease Member Section	Content Reviewer—Task Force on Performance Measures
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### **Appendix 3. Abbreviations**

AAA = abdominal aortic aneurysm ABI = ankle-brachial index ALI = acute limb ischemia CLI = critical limb ischemia GDMT = guideline-directed management and therapy MRA = magnetic resonance angiography PAD = peripheral artery disease RCT = randomized controlled trial SPP = skin perfusion pressure TBI = toe-brachial index TcPO₂ = transcutaneous oxygen pressure QoL = quality of life

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### References

- 1. Committee on Standards for Developing Trustworthy Clinical Practice Guidelines; Institute of Medicine. *Clinical Practice Guidelines We Can Trust.* Washington, DC: The National Academies Press, 2011.
- Committee on Standards for Systematic Reviews of Comparative Effectiveness Research, Institute of Medicine (U.S.), Eden J, Editors. *Finding What Works in Health Care: Standards for Systematic Reviews*. Washington, DC: National Academies Press; 2011.
- 3. ACCF/AHA Task Force on Practice Guidelines. Methodology Manual and Policies From the ACCF/AHA Task Force on Practice Guidelines. American College of Cardiology and American Heart Association. 2010. Available at: <u>http://assets.cardiosource.com/Methodology_Manual_for_ACC_AHA_Writing_Committees.pdf</u> and <u>http://my.americanheart.org/idc/groups/ahamah-</u>

public/@wcm/@sop/documents/downloadable/ucm_319826.pdf. Accessed January 23, 2015.

- Halperin JL, Levine GN, Al-Khatib SM, et al. Further evolution of the ACC/AHA clinical practice guideline recommendation classification system: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. Circulation. 2016;133:1426-28.
- 5. Jacobs AK, Kushner FG, Ettinger SM, et al. ACCF/AHA clinical practice guideline methodology summit report: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Circulation. 2013;127:268-310.
- 6. Jacobs AK, Anderson JL, Halperin JL. The evolution and future of ACC/AHA clinical practice guidelines: a 30-year journey: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines. Circulation. 2014;130:1208–17.
- Anderson JL, Heidenreich PA, Barnett PG, et al. ACC/AHA statement on cost/value methodology in clinical practice guidelines and performance measures: a report of the American College of Cardiology/American Heart Association Task Force on Performance Measures and Task Force on Practice Guidelines. Circulation. 2014;129:2329–45.
- Arnett DK, Goodman RA, Halperin JL, et al. AHA/ACC/HHS strategies to enhance application of clinical practice guidelines in patients with cardiovascular disease and comorbid conditions: from the American Heart Association, American College of Cardiology, and U.S. Department of Health and Human Services. Circulation. 2014;130:1662–7.
- 9. Gerhard-Herman MD, Gornik HL, Barrett C, et al. 2016 AHA/ACC guideline on the management of patients with lower extremity peripheral artery disease: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. Circulation. IN PRESS: 2016.
- 10. Hirsch AT, Haskal ZJ, Hertzer NR, et al. ACC/AHA 2005 guidelines for the management of patients with peripheral arterial disease (lower extremity, renal, mesenteric, and abdominal aortic): executive summary: a collaborative report from the American Association for Vascular Surgery/Society for Vascular Surgery, Society for Cardiovascular Angiography and Interventions, Society for Vascular Medicine and Biology, Society of Interventional Radiology, and the ACC/AHA Task Force on Practice Guidelines (Writing Committee to Develop Guidelines for the Management of Patients With Peripheral Arterial Disease). Circulation. 2006;113:e463–654.
- Rooke TW, Hirsch AT, Misra S, et al. 2011 ACCF/AHA focused update of the guideline for the management of patients with peripheral artery disease (updating the 2005 guideline): a report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. Circulation. 2011;124:2020–45.
- Jones WS, Schmit KM, Vemulapalli S, et al. Treatment Strategies for Patients With Peripheral Artery Disease. Comparative Effectiveness Review No. 118. The Duke Evidence-based Practice Center under Contract No 290-2007-10066-I. 2013;Available at: <u>http://www.effectivehealthcare.ahrq.gov/ehc/products/368/1415/Peripheral-Artery-Disease-Treatment-130301.pdf</u>. Accessed September 25, 2016.
- 13. Vemulapalli S, Dolor RJ, Hasselblad V, et al. Comparative effectiveness of medical therapy, supervised exercise, and revascularization for patients with intermittent claudication: a network meta-analysis. Clin Cardiol. 2015;38:378-86.
- 14. Schmit K, Dolor RJ, Jones WS, et al. Comparative effectiveness review of antiplatelet agents in peripheral artery disease. J Am Heart Assoc. 2014;3:e001330.

- 15. Jones WS, Dolor RJ, Hasselblad V, et al. Comparative effectiveness of endovascular and surgical revascularization for patients with peripheral artery disease and critical limb ischemia: systematic review of revascularization in critical limb ischemia. Am Heart J. 2014;167:489-98.e7.
- 16. Menard MT, Farber A. The BEST-CLI trial: a multidisciplinary effort to assess whether surgical or endovascular therapy is better for patients with critical limb ischemia. Semin Vasc Surg. 2014;27:82-4.
- 17. Popplewell MA, Davies H, Jarrett H, et al. Bypass versus angio plasty in severe ischaemia of the leg 2 (BASIL-2) trial: study protocol for a randomised controlled trial. Trials. 2016;17:11.
- 18. Menard MT, Farber A, Assmann SF, et al. Design and rationale of the Best Endovascular Versus Best Surgical Therapy for Patients With Critical Limb Ischemia (BEST-CLI) Trial. J Am Heart Assoc. 2016;5;e003219.
- Mozaffarian D, Benjamin EJ, Go AS, et al. Heart disease and stroke statistics—2016 update: a report from the American Heart Association [published correction appears in Circulation.2016;133:e599]. Circulation. 2016;133:e38-360.
- 20. Fowkes FG, Rudan D, Rudan I, et al. Comparison of global estimates of prevalence and risk factors for peripheral artery disease in 2000 and 2010: a systematic review and analysis. Lancet. 2013;382:1329-40.
- Creager MA, Belkin M, Bluth EI, et al. 2012 ACCF/AHA/ACR/SCAI/SIR/STS/SVM/SVN/SVS key data elements and definitions for peripheral atherosclerotic vascular disease: a report of the American College of Cardiology Foundation/American Heart Association Task Force on Clinical Data Standards (Writing Committee to develop Clinical Data Standards for Peripheral Atherosclerotic Vascular Disease). Circulation. 2012;125:395–467.
- 22. Rutherford RB, Baker JD, Ernst C, et al. Recommended standards for reports dealing with lower extremity ischemia: revised version. J Vasc Surg. 1997;26:517-38.
- 23. Norgren L, Hiatt WR, Dormandy JA, et al. Inter-society consensus for the management of peripheral arterial disease (TASC II). Eur J Vasc Endovasc Surg. 2007;(33 suppl 1):S1-75.
- 24. Fakhry F, Rouwet EV, den Hoed PT, et al. Long-term clinical effectiveness of supervised exercise therapy on versus endovascular revascularization for intermittent claudication from a randomized clinical trial. Br J Surg. 2013;100:1164-71.
- 25. Murphy TP, Cutlip DE, Regensteiner JG, et al. Supervised exercise versus primary stenting for claudication resulting from aortoiliac peripheral artery disease: six-month outcomes from the claudication: exercise versus endoluminal revascularization (CLEVER) study. Circulation. 2012;125:130-9.
- 26. Murphy TP, Cutlip DE, Regensteiner JG, et al. Supervised exercise, stent revascularization, or medical therapy for claudication due to aortoiliac peripheral artery disease: the CLEVER study. J Am Coll Cardiol. 2015;65:999-1009.
- 27. Treat-Jacobson D, Bronas UG, Leon AS. Efficacy of arm-ergometry versus treadmill exercise training to improve walking distance in patients with claudication. Vasc Med. 2009;14:203-13.
- 28. Hiatt WR, Regensteiner JG, Hargarten ME, et al. Benefit of exercise conditioning for patients with peripheral arterial disease. Circulation. 1990;81:602-9.
- 29. Parmenter BJ, Dieberg G, Smart NA. Exercise training for management of peripheral arterial disease: a systematic review and meta-analysis. Sports Med. 2015;45:231-44.
- 30. Parmenter BJ, Dieberg G, Phipps G, et al. Exercise training for health-related quality of life in peripheral artery disease: a systematic review and meta-analysis. Vasc Med. 2015;20:30-40.
- Pilz M, Kandioler-Honetz E, Wenkstetten-Holub A, et al. Evaluation of 6- and 12-month supervised exercise training on strength and endurance parameters in patients with peripheral arterial disease. Wien Klin Wochenschr. 2014;126:383-9.
- 32. Regensteiner JG, Steiner JF, Hiatt WR. Exercise training improves functional status in patients with peripheral arterial disease. J Vasc Surg. 1996;23:104-15.
- 33. Regensteiner JG. Exercise in the treatment of claudication: assessment and treatment of functional impairment. Vasc Med. 1997;2:238-42.
- 34. Stewart KJ, Hiatt WR, Regensteiner JG, et al. Exercise training for claudication. N Engl J Med. 2002;347:1941-51.
- Collins TC, Lunos S, Carlson T, et al. Effects of a home-based walking intervention on mobility and quality of life in people with diabetes and peripheral arterial disease: a randomized controlled trial. Diabetes Care. 2011;34:2174-9.
- 36. Fakhry F, Spronk S, de Ridder M, et al. Long-term effects of structured home-based exercise program on functional capacity and quality of life in patients with intermittent claudication. Arch Phys Med Rehabil. 2011;92:1066-73.

- 37. Gardner AW, Parker DE, Montgomery PS, et al. Step-monitored home exercise improves ambulation, vascular function, and inflammation in symptomatic patients with peripheral artery disease: a randomized controlled trial. J Am Heart Assoc. 2014;3:e001107.
- 38. Mays RJ, Rogers RK, Hiatt WR, et al. Community walking programs for treatment of peripheral artery disease. J Vasc Surg. 2013;58:1678-87.
- 39. McDermott MM, Domanchuk K, Liu K, et al. The Group Oriented Arterial Leg Study (GOALS) to improve walking performance in patients with peripheral arterial disease. Contemp Clin Trials. 2012;33:1311-20.
- 40. McDermott MM, Mehta S, Greenland P. Exertional leg symptoms other than intermittent claudication are common in peripheral arterial disease. Arch Intern Med. 1999;159:387-92.
- 41. Hirsch AT, Criqui MH, Treat-Jacobson D, et al. Peripheral arterial disease detection, awareness, and treatment in primary care. JAMA. 2001;286:1317-24.
- 42. McDermott MM, Greenland P, Liu K, et al. Leg symptoms in peripheral arterial disease: associated clinical characteristics and functional impairment. JAMA. 2001;286:1599-606.
- 43. Clark CE, Taylor RS, Shore AC, et al. Association of a difference in systolic blood pressure between arms with vascular disease and mortality: a systematic review and meta-analysis. Lancet. 2012;379:905-14.
- 44. Singh S, Sethi A, Singh M, et al. Simultaneously measured inter-arm and inter-leg systolic blood pressure differences and cardiovascular risk stratification: a systemic review and meta-analysis. J Am Soc Hypertens. 2015;9:640-50.e12.
- 45. Shadman R, Criqui MH, Bundens WP, et al. Subclavian artery stenosis: prevalence, risk factors, and association with cardiovascular diseases. J Am Coll Cardiol. 2004;44:618-23.
- 46. Aboyans V, Criqui MH, Abraham P, et al. Measurement and interpretation of the ankle-brachial index: a scientific statement from the American Heart Association. Circulation. 2012;126:2890-909.
- 47. Rose GA. The diagnosis of ischaemic heart pain and intermittent claudication in field surveys. Bull World Health Organ. 1962;27:645-58.
- 48. Khan NA, Rahim SA, Anand SS, et al. Does the clinical examination predict lower extremity peripheral arterial disease? JAMA. 2006;295:536-46.
- 49. Criqui MH, Denenberg JO, Bird CE, et al. The correlation between symptoms and non-invasive test results in patients referred for peripheral arterial disease testing. Vasc Med. 1996;1:65-71.
- 50. Armstrong DWJ, Tobin C, Matangi MF. The accuracy of the physical examination for the detection of lower extremity peripheral arterial disease. Can J Cardiol. 2010;26:e346-50.
- 51. Cournot M, Boccalon H, Cambou JP, et al. Accuracy of the screening physical examination to identify subclinical atherosclerosis and peripheral arterial disease in asymptomatic subjects. J Vasc Surg. 2007;46:1215-21.
- 52. Wassel CL, Loomba R, Ix JH, et al. Family history of peripheral artery disease is associated with prevalence and severity of peripheral artery disease: the San Diego Population Study. J Am Coll Cardiol. 2011;58:1386-92.
- 53. Eslahpazir BA, Allemang MT, Lakin RO, et al. Pulse volume recording does not enhance segmental pressure readings for peripheral arterial disease stratification. Ann Vasc Surg. 2014;28:18-27.
- 54. Raines JK, Darling RC, Buth J, et al. Vascular laboratory criteria for the management of peripheral vascular disease of the lower extremities. Surgery. 1976;79:21-9.
- 55. Mahe G, Pollak AW, Liedl DA, et al. Discordant diagnosis of lower extremity peripheral artery disease using American Heart Association postexercise guidelines. Medicine (Baltimore ). 2015;94:e1277.
- 56. Nicolaï SP, Viechtbauer W, Kruidenier LM, et al. Reliability of treadmill testing in peripheral arterial disease: a meta-regression analysis. J Vasc Surg. 2009;50:322-9.
- 57. Stein R, Hriljac I, Halperin JL, et al. Limitation of the resting ankle-brachial index in symptomatic patients with peripheral arterial disease. Vasc Med. 2006;11:29-33.
- Laing SP, Greenhalgh RM. Standard exercise test to assess peripheral arterial disease. Br Med J. 1980;280:13-6.
- 59. Sumner DS, Strandness DE. The relationship between calf blood flow and ankle blood pressure in patients with intermittent claudication. Surgery. 1969;65:763-71.
- 60. Schröder F, Diehm N, Kareem S, et al. A modified calculation of ankle-brachial pressure index is far more sensitive in the detection of peripheral arterial disease. J Vasc Surg. 2006;44:531-6.
- 61. Premalatha G, Ravikumar R, Sanjay R, et al. Comparison of colour duplex ultrasound and ankle-brachial pressure index measurements in peripheral vascular disease in type 2 diabetic patients with foot infections. J Assoc Physicians India. 2002;50:1240-4.
- 62. Allen J, Oates CP, Henderson J, et al. Comparison of lower limb arterial assessments using color-duplex ultrasound and ankle/brachial pressure index measurements. Angiology. 1996;47:225-32.

- 63. Lijmer JG, Hunink MG, van den Dungen JJ, et al. ROC analysis of noninvasive tests for peripheral arterial disease. Ultrasound Med Biol. 1996;22:391-8.
- 64. Guo X, Li J, Pang W, et al. Sensitivity and specificity of ankle-brachial index for detecting angiographic stenosis of peripheral arteries. Circ J. 2008;72:605-10.
- 65. Niazi K, Khan TH, Easley KA. Diagnostic utility of the two methods of ankle brachial index in the detection of peripheral arterial disease of lower extremities. Catheter Cardiovasc Interv. 2006;68:788-92.
- 66. Aboyans V, Ho E, Denenberg JO, et al. The association between elevated ankle systolic pressures and peripheral occlusive arterial disease in diabetic and nondiabetic subjects. J Vasc Surg. 2008;48:1197-203.
- 67. Belch J, MacCuish A, Campbell I, et al. The Prevention of Progression of Arterial Disease and Diabetes (POPADAD) trial: factorial randomised placebo controlled trial of aspirin and antioxidants in patients with diabetes and asymptomatic peripheral arterial disease. BMJ. 2008;337:a1840.
- 68. Fowkes FG, Price JF, Stewart MC, et al. Aspirin for prevention of cardiovascular events in a general population screened for a low ankle brachial index: a randomized controlled trial. JAMA. 2010;303:841-8.
- 69. Alahdab F, Wang AT, Elraiyah TA, et al. A systematic review for the screening for peripheral arterial disease in asymptomatic patients. J Vasc Surg. 2015;61:42S-53S.
- 70. Lin JS, Olson CM, Johnson ES, et al. The ankle-brachial index for peripheral artery disease screening and cardiovascular disease prediction among asymptomatic adults: a systematic evidence review for the U.S. Preventive Services Task Force. Ann Intern med. 2013;333-41.
- 71. Diehm C, Allenberg JR, Pittrow D, et al. Mortality and vascular morbidity in older adults with asymptomatic versus symptomatic peripheral artery disease. Circulation. 2009;120:2053-61.
- 72. Fowkes FG, Murray GD, Butcher I, et al. Ankle brachial index combined with Framingham Risk Score to predict cardiovascular events and mortality: a meta-analysis. JAMA. 2008;300:197-208.
- 73. Ratanakorn D, Keandoungchun J, Tegeler CH. Prevalence and association between risk factors, stroke subtypes, and abnormal ankle brachial index in acute ischemic stroke. J Stroke Cerebrovasc Dis. 2012;21:498-503.
- 74. Sen S, Lynch DR, Kaltsas E, et al. Association of asymptomatic peripheral arterial disease with vascular events in patients with stroke or transient ischemic attack. Stroke. 2009;40:3472-7.
- 75. Bundó M, Muñoz L, Pérez C, et al. Asymptomatic peripheral arterial disease in type 2 diabetes patients: a 10year follow-up study of the utility of the ankle brachial index as a prognostic marker of cardiovascular disease. Ann Vasc Surg. 2010;24:985-93.
- 76. Bouisset F, Bongard V, Ruidavets JB, et al. Prognostic usefulness of clinical and subclinical peripheral arterial disease in men with stable coronary heart disease. Am J Cardiol. 2012;110:197-202.
- 77. Hiramoto JS, Katz R, Ix JH, et al. Sex differences in the prevalence and clinical outcomes of subclinical peripheral artery disease in the Health, Aging, and Body Composition (Health ABC) study. Vascular. 2014;22:142-8.
- 78. Jiménez M, Dorado L, Hernández-Pérez M, et al. Ankle-brachial index in screening for asymptomatic carotid and intracranial atherosclerosis. Atherosclerosis. 2014;233:72-5.
- 79. Tsivgoulis G, Bogiatzi C, Heliopoulos I, et al. Low ankle-brachial index predicts early risk of recurrent stroke in patients with acute cerebral ischemia. Atherosclerosis. 2012;220:407-12.
- 80. McDermott MM, Liu K, Guralnik JM, et al. Home-based walking exercise intervention in peripheral artery disease: a randomized clinical trial. JAMA. 2013;310:57-65.
- 81. McDermott MM, Fried L, Simonsick E, et al. Asymptomatic peripheral arterial disease is independently associated with impaired lower extremity functioning: the Women's Health and Aging Study. Circulation. 2000;101:1007-12.
- McDermott MM, Applegate WB, Bonds DE, et al. Ankle brachial index values, leg symptoms, and functional performance among community-dwelling older men and women in the Lifestyle Interventions and Independence for Elders Study. J Am Heart Assoc. 2013;2:e000257.
- 83. McDermott MM, Liu K, Greenland P, et al. Functional decline in peripheral arterial disease: associations with the ankle brachial index and leg symptoms. JAMA. 2004;292:453-61.
- 84. McDermott MM, Ferrucci L, Liu K, et al. Leg symptom categories and rates of mobility decline in peripheral arterial disease. J Am Geriatr Soc. 2010;58:1256-62.
- 85. McDermott MM, Liu K, Ferrucci L, et al. Physical performance in peripheral arterial disease: a slower rate of decline in patients who walk more. Ann Intern Med. 2006;144:10-20.
- 86. McDermott MM, Guralnik JM, Criqui MH, et al. Home-based walking exercise in peripheral artery disease: 12-month follow-up of the GOALS randomized trial. J Am Heart Assoc. 2014;3:e000711.
- 87. Selvin E, Erlinger TP. Prevalence of and risk factors for peripheral arterial disease in the United States: results from the National Health and Nutrition Examination Survey, 1999-2000. Circulation. 2004;110:738-43.

- 88. Ramos R, García-Gil M, Comas-Cufi M, et al. Statins for prevention of cardiovascular events in a low-risk population with low ankle brachial index. J Am Coll Cardiol. 2016;67:630-40.
- 89. Fowkes FG, Murray GD, Butcher I, et al. Development and validation of an ankle brachial index risk model for the prediction of cardiovascular events. Eur J Prev Cardiol. 2014;21:310-20.
- 90. Criqui MH, Vargas V, Denenberg JO, et al. Ethnicity and peripheral arterial disease: the San Diego Population Study. Circulation. 2005;112:2703-7.
- 91. Carter SA, Tate RB. Value of toe pulse waves in addition to systolic pressures in the assessment of the severity of peripheral arterial disease and critical limb ischemia. J Vasc Surg. 1996;24:258-65.
- 92. Ramsey DE, Manke DA, Sumner DS. Toe blood pressure. A valuable adjunct to ankle pressure measurement for assessing peripheral arterial disease. J Cardiovasc Surg (Torino). 1983;24:43-8.
- 93. Vincent DG, Salles-Cunha SX, Bernhard VM, et al. Noninvasive assessment of toe systolic pressures with special reference to diabetes mellitus. J Cardiovasc Surg (Torino). 1983;24:22-8.
- 94. Carter SA. Clinical measurement of systolic pressures in limbs with arterial occlusive disease. JAMA. 1969;207:1869-74.
- 95. Biotteau E, Mahe G, Rousseau P, et al. Transcutaneous oxygen pressure measurements in diabetic and nondiabetic patients clinically suspected of severe limb ischemia: a matched paired retrospective analysis. Int Angiol. 2009;28:479-83.
- 96. Yamada T, Ohta T, Ishibashi H, et al. Clinical reliability and utility of skin perfusion pressure measurement in ischemic limbs—comparison with other noninvasive diagnostic methods. J Vasc Surg. 2008;47:318-23.
- 97. Castronuovo JJ, Adera HM, Smiell JM, et al. Skin perfusion pressure measurement is valuable in the diagnosis of critical limb ischemia. J Vasc Surg. 1997;26:629-37.
- 98. Bunte MC, Jacob J, Nudelman B, et al. Validation of the relationship between ankle-brachial and toe-brachial indices and infragenicular arterial patency in critical limb ischemia. Vasc Med. 2015;20:23-9.
- 99. Shishehbor MH, Hammad TA, Zeller T, et al. An analysis of IN.PACT DEEP randomized trial on the ociationlimitations of the societal guidelines-recommended hemodynamic parameters to diagnose critical limb ischemia. J Vasc Surg. 2016;63:1311-7.
- Burbelko M, Augsten M, Kalinowski MO, et al. Comparison of contrast-enhanced multi-station MR angiography and digital subtraction angiography of the lower extremity arterial disease. J Magn Reson Imaging. 2013;37:1427-35.
- 101. Shareghi S, Gopal A, Gul K, et al. Diagnostic accuracy of 64 multidetector computed tomographic angiography in peripheral vascular disease. Catheter Cardiovasc Interv. 2010;75:23-31.
- Ota H, Takase K, Igarashi K, et al. MDCT compared with digital subtraction angiography for assessment of lower extremity arterial occlusive disease: importance of reviewing cross-sectional images. AJR Am J Roentgenol. 2004;182:201-9.
- de Vries SO, Hunink MG, Polak JF. Summary receiver operating characteristic curves as a technique for metaanalysis of the diagnostic performance of duplex ultrasonography in peripheral arterial disease. Acad Radiol. 1996;3:361-9.
- 104. Andreucci M, Solomon R, Tasanarong A. Side effects of radiographic contrast media: pathogenesis, risk factors, and prevention. Biomed Res Int. 2014;2014:741018.
- 105. Stacul F, van der Molen AJ, Reimer P, et al. Contrast induced nephropathy: updated ESUR Contrast Media Safety Committee guidelines. Eur Radiol. 2011;21:2527-41.
- 106. McCullough PA, Capasso P. Patient discomfort associated with the use of intra-arterial iodinated contrast media: a meta-analysis of comparative randomized controlled trials. BMC Med Imaging. 2011;11:12.
- 107. Giugliano G, Laurenzano E, Rengo C, et al. Abdominal aortic aneurysm in patients affected by intermittent claudication: prevalence and clinical predictors. BMC Surg. 2012;12(suppl 1):S17.
- 108. Barba A, Estallo L, Rodríguez L, et al. Detection of abdominal aortic aneurysm in patients with peripheral artery disease. Eur J Vasc Endovasc Surg. 2005;30:504-8.
- 109. Kurvers HA, van der Graaf Y, Blankensteijn JD, et al. Screening for asymptomatic internal carotid artery stenosis and aneurysm of the abdominal aorta: comparing the yield between patients with manifest atherosclerosis and patients with risk factors for atherosclerosis only. J Vasc Surg. 2003;37:1226-33.
- 110. Lee JY, Lee SW, Lee WS, et al. Prevalence and clinical implications of newly revealed, asymptomatic abnormal ankle-brachial index in patients with significant coronary artery disease. JACC Cardiovasc Interv. 2013;6:1303-13.
- 111. McFalls EO, Ward HB, Moritz TE, et al. Coronary-artery revascularization before elective major vascular surgery. N Engl J Med. 2004;351:2795-804.

- 112. Sultan S, Chua BY, Hamada N, et al. Preoperative vascular screening in the presence of aortic, carotid and peripheral pathology for patients undergoing their first arterial intervention: 18 month follow-up. Int Angiol. 2013;32:281-90.
- 113. Hansen KJ, Edwards MS, Craven TE, et al. Prevalence of renovascular disease in the elderly: a populationbased study. J Vasc Surg. 2002;36:443-51.
- 114. Leertouwer TC, Pattynama PM, van den Berg-Huysmans A. Incidental renal artery stenosis in peripheral vascular disease: a case for treatment? Kidney Int. 2001;59:1480-3.
- 115. Olin JW, Melia M, Young JR, et al. Prevalence of atherosclerotic renal artery stenosis in patients with atherosclerosis elsewhere. Am J Med. 1990;88:46N-51N.
- 116. Krishnamurthy V, Munir K, Rectenwald JE, et al. Contemporary outcomes with percutaneous vascular interventions for peripheral critical limb ischemia in those with and without poly-vascular disease. Vasc Med. 2014;19:491-9.
- 117. Selvin E, Hirsch AT. Contemporary risk factor control and walking dysfunction in individuals with peripheral arterial disease: NHANES 1999-2004. Atherosclerosis. 2008;201:425-33.
- 118. Pande RL, Perlstein TS, Beckman JA, et al. Secondary prevention and mortality in peripheral artery disease: National Health and Nutrition Examination Study, 1999 to 2004. Circulation. 2011;124:17-23.
- 119. Dawson DL, Cutler BS, Hiatt WR, et al. A comparison of cilostazol and pentoxifylline for treating intermittent claudication. Am J Med. 2000;109:523-30.
- 120. Lee C, Nelson PR. Effect of cilostazol prescribed in a pragmatic treatment program for intermittent claudication. Vasc Endovascular Surg. 2014;48:224-9.
- 121. Antithrombotic Trialists' Collaboration. Collaborative meta-analysis of randomised trials of antiplatelet therapy for prevention of death, myocardial infarction, and stroke in high risk patients. BMJ. 2002;324:71-86.
- 122. Catalano M, Born G, Peto R. Prevention of serious vascular events by aspirin amongst patients with peripheral arterial disease: randomized, double-blind trial. J Intern Med. 2007;261:276-84.
- 123. Berger JS, Krantz MJ, Kittelson JM, et al. Aspirin for the prevention of cardiovascular events in patients with peripheral artery disease: a meta-analysis of randomized trials. JAMA. 2009;301:1909-19.
- 124. CAPRIE Steering Committee. A randomised, blinded, trial of clopidogrel versus aspirin in patients at risk of ischaemic events (CAPRIE). Lancet. 1996;348:1329-39.
- 125. Cacoub PP, Bhatt DL, Steg PG, et al. Patients with peripheral arterial disease in the CHARISMA trial. Eur Heart J. 2009;30:192-201.
- 126. Bhatt DL, Flather MD, Hacke W, et al. Patients with prior myocardial infarction, stroke, or symptomatic peripheral arterial disease in the CHARISMA trial. J Am Coll Cardiol. 2007;49:1982-8.
- 127. Tepe G, Bantleon R, Brechtel K, et al. Management of peripheral arterial interventions with mono or dual antiplatelet therapy—the MIRROR study: a randomised and double-blinded clinical trial. Eur Radiol. 2012;22:1998-2006.
- Armstrong EJ, Anderson DR, Yeo KK, et al. Association of dual-antiplatelet therapy with reduced major adverse cardiovascular events in patients with symptomatic peripheral arterial disease. J Vasc Surg. 2015;62:157-65.
- Strobl FF, Brechtel K, Schmehl J, et al. Twelve-month results of a randomized trial comparing mono with dual antiplatelet therapy in endovascularly treated patients with peripheral artery disease. J Endovasc Ther. 2013;20:699-706.
- Belch JJ, Dormandy J, CASPAR Writing Committee, et al. Results of the randomized, placebo-controlled Clopidogrel and Acetylsalicylic Acid in Bypass Surgery for Peripheral Arterial Disease (CASPAR) trial. J Vasc Surg. 2010;52:825-33.
- 131. Bonaca MP, Scirica BM, Creager MA, et al. Vorapaxar in patients with peripheral artery disease: results from TRA2°P-TIMI 50. Circulation. 2013;127:1522-9.
- 132. Morrow DA, Braunwald E, Bonaca MP, et al. Vorapaxar in the secondary prevention of atherothrombotic events. N Engl J Med. 2012;366:1404-13.
- 133. Bohula EA, Aylward PE, Bonaca MP, et al. Efficacy and safety of vorapaxar with and without a thienopyridine for secondary prevention in patients with previous myocardial infarction and no history of stroke or transient ischemic attack: results from TRA 2°P-TIMI 50. Circulation. 2015;132:1871-9.
- 134. Bonaca MP, Gutierrez JA, Creager MA, et al. Acute limb ischemia and outcomes with vorapaxar in patients with peripheral artery disease: results from the Trial to Assess the Effects of Vorapaxar in Preventing Heart Attack and Stroke in patients With Atherosclerosis-Thrombolysis in Myocardial Infarction 50 (TRA2°P-TIMI 50). Circulation. 2016;133:997-1005.
- 135. Kumbhani DJ, Steg PG, Cannon CP, et al. Statin therapy and long-term adverse limb outcomes in patients with peripheral artery disease: insights from the REACH registry. Eur Heart J. 2014;35:2864-72.

- 136. Vogel TR, Dombrovskiy VY, Galiñanes EL, et al. Preoperative statins and limb salvage after lower extremity revascularization in the Medicare population. Circ Cardiovasc Interv. 2013;6:694-700.
- 137. Heart Protection Study Collaborative Group. Randomized trial of the effects of cholesterol-lowering with simvastatin on peripheral vascular and other major vascular outcomes in 20,536 people with peripheral arterial disease and other high-risk conditions. J Vasc Surg. 2007;45:645-54.
- 138. Mohler ER3, Hiatt WR, Creager MA. Cholesterol reduction with atorvastatin improves walking distance in patients with peripheral arterial disease. Circulation. 2003;108:1481-6.
- 139. Aung PP, Maxwell HG, Jepson RG, et al. Lipid-lowering for peripheral arterial disease of the lower limb. Cochrane Database Syst Rev. 2007;CD000123.
- 140. Bavry AA, Anderson RD, Gong Y, et al. Outcomes Among hypertensive patients with concomitant peripheral and coronary artery disease: findings from the INternational VErapamil-SR/Trandolapril STudy. Hypertension. 2010;55:48-53.
- 141. Feringa HH, van Waning VH, Bax JJ, et al. Cardioprotective medication is associated with improved survival in patients with peripheral arterial disease. J Am Coll Cardiol. 2006;47:1182-7.
- 142. Sleight P. The HOPE Study (Heart Outcomes Prevention Evaluation). J Renin Angiotensin Aldosterone Syst. 2000;1:18-20.
- 143. Yusuf S, Teo KK, Pogue J, et al. Telmisartan, ramipril, or both in patients at high risk for vascular events. N Engl J Med. 2008;358:1547-59.
- 144. ALLHAT Officers and Coordinators for the ALLHAT Collaborative Research Group. The Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial. Major outcomes in high-risk hypertensive patients randomized to angiotensin-converting enzyme inhibitor or calcium channel blocker vs diuretic: The Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT). JAMA. 2002;288:2981-97.
- 145. Ostergren J, Sleight P, Dagenais G, et al. Impact of ramipril in patients with evidence of clinical or subclinical peripheral arterial disease. Eur Heart J. 2004;25:17-24.
- 146. Yusuf S, Sleight P, Pogue J, et al. Effects of an angiotensin-converting-enzyme inhibitor, ramipril, on cardiovascular events in high-risk patients. The Heart Outcomes Prevention Evaluation Study Investigators. N Engl J Med. 2000;342:145-53.
- 147. Sarac TP, Huber TS, Back MR, et al. Warfarin improves the outcome of infrainguinal vein bypass grafting at high risk for failure. J Vasc Surg. 1998;28:446-57.
- 148. Efficacy of oral anticoagulants compared with aspirin after infrainguinal bypass surgery (The Dutch Bypass Oral Anticoagulants or Aspirin Study): a randomised trial. Lancet. 2000;355:346-51.
- 149. Bedenis R, Lethaby A, Maxwell H, et al. Antiplatelet agents for preventing thrombosis after peripheral arterial bypass surgery. Cochrane Database Syst Rev. 2015;CD000535.
- 150. Alonso-Coello P, Bellmunt S, McGorrian C, et al. Antithrombotic therapy in peripheral artery disease: Antithrombotic Therapy and Prevention of Thrombosis, 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines. Chest. 2012;141:e669S-90S.
- 151. Johnson WC, Williford WO, Department of Veterans Affairs Cooporative Study #362. Benefits, morbidity, and mortality associated with long-term administration of oral anticoagulant therapy to patients with peripheral arterial bypass procedures: a prospective randomized study. J Vasc Surg. 2002;35:413-21.
- Anand S, Yusuf S, Xie C, et al. Oral anticoagulant and antiplatelet therapy and peripheral arterial disease. N Engl J Med. 2007;357:217-27.
- 153. Hennrikus D, Joseph AM, Lando HA, et al. Effectiveness of a smoking cessation program for peripheral artery disease patients: a randomized controlled trial. J Am Coll Cardiol. 2010;56:2105-12.
- 154. Stead LF, Buitrago D, Preciado N, et al. Physician advice for smoking cessation. Cochrane Database Syst Rev. 2013;CD000165.
- 155. Hoel AW, Nolan BW, Goodney PP, et al. Variation in smoking cessation after vascular operations. J Vasc Surg. 2013;57:1338-44.
- 156. Rigotti NA, Pipe AL, Benowitz NL, et al. Efficacy and safety of varenicline for smoking cessation in patients with cardiovascular disease: a randomized trial. Circulation. 2010;121:221-9.
- 157. Rigotti NA, Regan S, Levy DE, et al. Sustained care intervention and postdischarge smoking cessation among hospitalized adults: a randomized clinical trial. JAMA. 2014;312:719-28.
- 158. Tonstad S, Farsang C, Klaene G, et al. Bupropion SR for smoking cessation in smokers with cardiovascular disease: a multicentre, randomised study. Eur Heart J. 2003;24:946-55.
- 159. Tan CE, Glantz SA. Association between smoke-free legislation and hospitalizations for cardiac, cerebrovascular, and respiratory diseases: a meta-analysis. Circulation. 2012;126:2177-83.

- 160. Lu L, Mackay DF, Pell JP. Association between level of exposure to secondhand smoke and peripheral arterial disease: cross-sectional study of 5,686 never smokers. Atherosclerosis. 2013;229:273-6.
- Singh S, Armstrong EJ, Sherif W, et al. Association of elevated fasting glucose with lower patency and increased major adverse limb events among patients with diabetes undergoing infrapopliteal balloon angioplasty. Vasc Med. 2014;19:307-14.
- 162. Takahara M, Kaneto H, Iida O, et al. The influence of glycemic control on the prognosis of Japanese patients undergoing percutaneous transluminal angioplasty for critical limb ischemia. Diabetes Care. 2010;33:2538-42.
- Bedenis R, Stewart M, Cleanthis M, et al. Cilostazol for intermittent claudication. Cochrane Database Syst Rev. 2014;CD003748.
- 164. Salhiyyah K, Senanayake E, Abdel-Hadi M, et al. Pentoxifylline for intermittent claudication. Cochrane Database Syst Rev. 2012;1:CD005262.
- 165. Villarruz MV, Dans A, Tan F. Chelation therapy for atherosclerotic cardiovascular disease. Cochrane Database Syst Rev. 2002;CD002785.
- 166. Lonn E, Held C, Arnold JM, et al. Rationale, design and baseline characteristics of a large, simple, randomized trial of combined folic acid and vitamins B6 and B12 in high-risk patients: the Heart Outcomes Prevention Evaluation (HOPE)-2 trial. Can J Cardiol. 2006;22:47-53.
- 167. Lonn E, Yusuf S, Arnold MJ, et al. Homocysteine lowering with folic acid and B vitamins in vascular disease. N Engl J Med. 2006;354:1567-77.
- 168. Khandanpour N, Loke YK, Meyer FJ, et al. Homocysteine and peripheral arterial disease: systematic review and meta-analysis. Eur J Vasc Endovasc Surg. 2009;38:316-22.
- Brenner I, Parry M, Brown CA. Exercise interventions for patients with peripheral arterial disease: a review of the literature. Phys Sportsmed. 2012;40:41-55.
- 170. Lane R, Ellis B, Watson L, et al. Exercise for intermittent claudication. Cochrane Database Syst Reveart 2014;CD000990.
- 171. Gardner AW, Parker DE, Montgomery PS, et al. Efficacy of quantified home-based exercise and supervised exercise in patients with intermittent claudication: a randomized controlled trial. Circulation. 2011;123:491-8.
- 172. Guidon M, McGee H. One-year effect of a supervised exercise programme on functional capacity and quality of life in peripheral arterial disease. Disabil Rehabil. 2013;35:397-404.
- 173. Saxton JM, Zwierska I, Blagojevic M, et al. Upper- versus lower-limb aerobic exercise training on healthrelated quality of life in patients with symptomatic peripheral arterial disease. J Vasc Surg. 2011;53:1265-73.
- 174. Gommans LN, Fokkenrood HJ, van Dalen HC, et al. Safety of supervised exercise therapy in patients with intermittent claudication. J Vasc Surg. 2015;61:512-8.
- 175. Langbein WE, Collins EG, Orebaugh C, et al. Increasing exercise tolerance of persons limited by claudication pain using polestriding. J Vasc Surg. 2002;35:887-93.
- 176. Walker RD, Nawaz S, Wilkinson CH, et al. Influence of upper- and lower-limb exercise training on cardiovascular function and walking distances in patients with intermittent claudication. J Vasc Surg. 2000;31:662-9.
- 177. Miller JD, Carter E, Shih J, et al. How to do a 3-minute diabetic foot exam. J Fam Pract. 2014;63:646-56.
- 178. Prompers L, Schaper N, Apelqvist J, et al. Prediction of outcome in individuals with diabetic foot ulcers: focus on the differences between individuals with and without peripheral arterial disease. The EURODIALE Study. Diabetologia. 2008;51:747-55.
- 179. Pickwell K, Siersma V, Kars M, et al. Predictors of lower-extremity amputation in patients with an infected diabetic foot ulcer. Diabetes Care. 2015;38:852-7.
- Dorresteijn JA, Kriegsman DM, Assendelft WJ, et al. Patient education for preventing diabetic foot ulceration. Cochrane Database Syst Rev. 2014;CD001488.
- Gardner SE, Hillis SL, Frantz RA. Clinical signs of infection in diabetic foot ulcers with high microbial load. Biol Res Nurs. 2009;11:119-28.
- Lipsky BA, Berendt AR, Cornia PB, et al. 2012 Infectious Diseases Society of America clinical practice guideline for the diagnosis and treatment of diabetic foot infections. Clin Infect Dis. 2012;54:e132-73.
- 183. Dinh MT, Abad CL, Safdar N. Diagnostic accuracy of the physical examination and imaging tests for osteomyelitis underlying diabetic foot ulcers: meta-analysis. Clin Infect Dis. 2008;47:519-27.
- Vartanian SM, Robinson KD, Ofili K, et al. Outcomes of neuroischemic wounds treated by a multidisciplinary amputation prevention service. Ann Vasc Surg. 2015;29:534-42.
- 185. Clerici G, Faglia E. Saving the limb in diabetic patients with ischemic foot lesions complicated by acute infection. Int J Low Extrem Wounds. 2014;13:273-93.
- 186. Leng GC, Lee AJ, Fowkes FG, et al. Incidence, natural history and cardiovascular events in symptomatic and asymptomatic peripheral arterial disease in the general population. Int J Epidemiol. 1996;25:1172-81.

- Dormandy J, Mahir M, Ascady G, et al. Fate of the patient with chronic leg ischaemia. A review article. J Cardiovasc Surg (Torino ). 1989;30:50-7.
- Jelnes R, Gaardsting O, Hougaard Jensen K, et al. Fate in intermittent claudication: outcome and risk factors. Br Med J (Clin Res Ed). 1986;293:1137-40.
- 189. Bloor K. Natural history of arteriosclerosis of the lower extremities: Hunterian lecture delivered at the Royal College of Surgeons of England on 22nd April 1960. Ann R Coll Surg Engl. 1961;28:36-52.
- 190. Nordanstig J, Taft C, Hensäter M, et al. Improved quality of life after 1 year with an invasive versus a noninvasive treatment strategy in claudicants: one-year results of the Invasive Revascularization or Not in Intermittent Claudication (IRONIC) Trial. Circulation. 2014;130:939-47.
- 191. Malgor RD, Alahdab F, Elraiyah TA, et al. A systematic review of treatment of intermittent claudication in the lower extremities. J Vasc Surg. 2015;61:54S-73S.
- 192. Tetteroo E, van Engelen AD, Spithoven JH, et al. Stent placement after iliac angioplasty: comparison of hemodynamic and angiographic criteria. Dutch Iliac Stent Trial Study Group. Radiology. 1996;201:155-9.
- 193. Udoff EJ, Barth KH, Harrington DP, et al. Hemodynamic significance of iliac artery stenosis: pressure measurements during angiography. Radiology. 1979;132:289-93.
- 194. Spronk S, Bosch JL, den Hoed PT, et al. Intermittent claudication: clinical effectiveness of endovascular revascularization versus supervised hospital-based exercise training-randomized controlled trial. Radiology. 2009;250:586-95.
- 195. Gelin J, Jivegård L, Taft C, et al. Treatment efficacy of intermittent claudication by surgical intervention, supervised physical exercise training compared to no treatment in unselected randomised patients, I: one year results of functional and physiological improvements. Eur J Vasc Endovasc Surg. 2001;22:107-13.
- 196. Greenhalgh RM, Belch JJ, Brown LC, et al. The adjuvant benefit of angioplasty in patients with mild to moderate intermittent claudication (MIMIC) managed by supervised exercise, smoking cessation advice and best medical therapy: results from two randomised trials for stenotic femoropopliteal and aortoiliac arterial disease. Eur J Vasc Endovasc Surg. 2008;36:680-8.
- 197. Hobbs SD, Marshall T, Fegan C, et al. The constitutive procoagulant and hypofibrinolytic state in patients with intermittent claudication due to infrainguinal disease significantly improves with percutaneous transluminal balloon angioplasty. J Vasc Surg. 2006;43:40-6.
- 198. Mazari FA, Gulati S, Rahman MN, et al. Early outcomes from a randomized, controlled trial of supervised exercise, angioplasty, and combined therapy in intermittent claudication. Ann Vasc Surg. 2010;24:69-79.
- 199. Mazari FA, Khan JA, Carradice D, et al. Randomized clinical trial of percutaneous transluminal angioplasty, supervised exercise and combined treatment for intermittent claudication due to femoropopliteal arterial disease. Br J Surg. 2012;99:39-48.
- 200. Nordanstig J, Gelin J, Hensäter M, et al. Walking performance and health-related quality of life after surgical or endovascular invasive versus non-invasive treatment for intermittent claudication—a prospective randomised trial. Eur J Vasc Endovasc Surg. 2011;42:220-7.
- 201. Nylaende M, Abdelnoor M, Stranden E, et al. The Oslo Balloon Angioplasty versus Conservative Treatment study (OBACT)—the 2-years results of a single centre, prospective, randomised study in patients with intermittent claudication. Eur J Vasc Endovasc Surg. 2007;33:3-12.
- Perkins JM, Collin J, Creasy TS, et al. Reprinted article "Exercise training versus angioplasty for stable claudication. Long and medium term results of a prospective, randomised trial." Eur J Vasc Endovasc Surg. 2011;42(suppl 1):S41-5.
- 203. Spronk S, Bosch JL, den Hoed PT, et al. Cost-effectiveness of endovascular revascularization compared to supervised hospital-based exercise training in patients with intermittent claudication: a randomized controlled trial. J Vasc Surg. 2008;48:1472-80.
- 204. Taft C, Karlsson J, Gelin J, et al. Treatment efficacy of intermittent claudication by invasive therapy, supervised physical exercise training compared to no treatment in unselected randomised patients, II: one-year results of health-related quality of life. Eur J Vasc Endovasc Surg. 2001;22:114-23.
- 205. Whyman MR, Fowkes FG, Kerracher EM, et al. Randomised controlled trial of percutaneous transluminal angioplasty for intermittent claudication. Eur J Vasc Endovasc Surg. 1996;12:167-72.
- 206. Whyman MR, Fowkes FG, Kerracher EM, et al. Is intermittent claudication improved by percutaneous transluminal angioplasty? A randomized controlled trial. J Vasc Surg. 1997;26:551-7.
- 207. Löfberg AM, Karacagil S, Ljungman C, et al. Percutaneous transluminal angioplasty of the femoropopliteal arteries in limbs with chronic critical lower limb ischemia. J Vasc Surg. 2001;34:114-21.
- 208. Capek P, McLean GK, Berkowitz HD. Femoropopliteal angioplasty. Factors influencing long-term success. Circulation. 1991;83:170-80.

- 209. Clark TW, Groffsky JL, Soulen MC. Predictors of long-term patency after femoropopliteal angioplasty: results from the STAR registry. J Vasc Interv Radiol. 2001;12:923-33.
- 210. Johnston KW, Rae M, Hogg-Johnston SA, et al. 5-year results of a prospective study of percutaneous transluminal angioplasty. Ann Surg. 1987;206:403-13.
- 211. Schulte KL, Pilger E, Schellong S, et al. Primary self-expanding nitinol stenting vs balloon angioplasty with optional bailout stenting for the treatment of infrapopliteal artery disease in patients with severe intermittent claudication or critical limb ischemia (EXPAND Study). J Endovasc Ther. 2015;22:690-7.
- Rastan A, Tepe G, Krankenberg H, et al. Sirolimus-eluting stents vs. bare-metal stents for treatment of focal lesions in infrapopliteal arteries: a double-blind, multi-centre, randomized clinical trial. Eur Heart J. 2011;32:2274-81.
- 213. Siablis D, Kitrou PM, Spiliopoulos S, et al. Paclitaxel-coated balloon angioplasty versus drug-eluting stenting for the treatment of infrapopliteal long-segment arterial occlusive disease: the IDEAS randomized controlled trial. JACC Cardiovasc Interv. 2014;7:1048-56.
- 214. Sachs T, Pomposelli F, Hamdan A, et al. Trends in the national outcomes and costs for claudication and limb threatening ischemia: angioplasty vs bypass graft. J Vasc Surg. 2011;54:1021-31.
- 215. Shammas NW, Shammas GA, Dippel EJ, et al. Predictors of distal embolization in peripheral percutaneous interventions: a report from a large peripheral vascular registry. J Invasive Cardiol. 2009;21:628-31.
- 216. Matsi PJ, Manninen HI. Complications of lower-limb percutaneous transluminal angioplasty: a prospective analysis of 410 procedures on 295 consecutive patients. Cardiovasc Intervent Radiol. 1998;21:361-6.
- 217. Antoniou GA, Chalmers N, Georgiadis GS, et al. A meta-analysis of endovascular versus surgical reconstruction of femoropopliteal arterial disease. J Vasc Surg. 2013;57:242-53.
- 218. Fowkes F, Leng GC. Bypass surgery for chronic lower limb ischaemia. Cochrane Database Syst Rev. 2008;CD002000.
- 219. Aihara H, Soga Y, Mii S, et al. Comparison of long-term outcome after endovascular therapy versus bypassonsurgery in claudication patients with Trans-Atlantic Inter-Society Consensus-II C and D femoropopliteal disease. Circ J. 2014;78:457-64.
- 220. Chiesa R, Marone EM, Tshomba Y, et al. Aortobifemoral bypass grafting using expanded polytetrafluoroethylene stretch grafts in patients with occlusive atherosclerotic disease. Ann Vasc Surg. 2009;23:764-9.
- 221. Goodney PP, Likosky DS, Cronenwett JL, et al. Predicting ambulation status one year after lower extremity bypass. J Vasc Surg. 2009;49:1431-9.
- Lo RC, Bensley RP, Dahlberg SE, et al. Presentation, treatment, and outcome differences between men and women undergoing revascularization or amputation for lower extremity peripheral arterial disease. J Vasc Surg. 2014;59:409-18.
- 223. Nguyen BN, Amdur RL, Abugideiri M, et al. Postoperative complications after common femoral endarterectomy. J Vasc Surg. 2015;61:1489-94.
- 224. Sachwani GR, Hans SS, Khoury MD, et al. Results of iliac stenting and aortofemoral grafting for iliac artery occlusions. J Vasc Surg. 2013;57:1030-7.
- 225. Siracuse JJ, Gill HL, Schneider DB, et al. Assessing the perioperative safety of common femoral endarterectomy in the endovascular era. Vasc Endovascular Surg. 2014;48:27-33.
- 226. Twine CP, McLain AD. Graft type for femoro-popliteal bypass surgery. Cochrane Database Syst Rev. 2010;CD001487.
- 227. Pereira CE, Albers M, Romiti M, et al. Meta-analysis of femoropopliteal bypass grafts for lower extremity arterial insufficiency. J Vasc Surg. 2006;44:510-7.
- 228. AbuRahma AF, Robinson PA, Holt SM. Prospective controlled study of polytetrafluoroethylene versus saphenous vein in claudicant patients with bilateral above knee femoropopliteal bypasses. Surgery. 1999;126:594-602.
- 229. Archie JP. Femoropopliteal bypass with either adequate ipsilateral reversed saphenous vein or obligatory polytetrafluoroethylene. Ann Vasc Surg. 1994;8:475-84.
- 230. Eugster T, Marti R, Gurke L, et al. Ten years after arterial bypass surgery for claudication: venous bypass is the primary procedure for TASC C and D lesions. World J Surg. 2011;35:2328-31.
- 231. Green RM, Abbott WM, Matsumoto T, et al. Prosthetic above-knee femoropopliteal bypass grafting: five-year results of a randomized trial. J Vasc Surg. 2000;31:417-25.
- 232. Hunink MG, Wong JB, Donaldson MC, et al. Patency results of percutaneous and surgical revascularization for femoropopliteal arterial disease. Med Decis Making. 1994;14:71-81.

- 233. Johnson WC, Lee KK. Comparative evaluation of externally supported Dacron and polytetrafluoroethylene prosthetic bypasses for femorofemoral and axillofemoral arterial reconstructions. Veterans Affairs Cooperative Study #141. J Vasc Surg. 1999;30:1077-83.
- 234. Klinkert P, Schepers A, Burger DH, et al. Vein versus polytetrafluoroethylene in above-knee femoropopliteal bypass grafting: five-year results of a randomized controlled trial. J Vasc Surg. 2003;37:149-55.
- 235. Feinglass J, McCarthy WJ, Slavensky R, et al. Functional status and walking ability after lower extremity bypass grafting or angioplasty for intermittent claudication: results from a prospective outcomes study. J Vasc Surg. 2000;31:93-103.
- 236. Koivunen K, Lukkarinen H. One-year prospective health-related quality-of-life outcomes in patients treated with conservative method, endovascular treatment or open surgery for symptomatic lower limb atherosclerotic disease. Eur J Cardiovasc Nurs. 2008;7:247-56.
- 237. Mori E, Komori K, Kume M, et al. Comparison of the long-term results between surgical and conservative treatment in patients with intermittent claudication. Surgery. 2002;131:S269-74.
- 238. Veith FJ, Gupta SK, Ascer E, et al. Six-year prospective multicenter randomized comparison of autologous saphenous vein and expanded polytetrafluoroethylene grafts in infrainguinal arterial reconstructions. J Vasc Surg. 1986;3:104-14.
- 239. Schweiger H, Klein P, Lang W. Tibial bypass grafting for limb salvage with ringed polytetrafluoroethylene prostheses: results of primary and secondary procedures. J Vasc Surg. 1993;18:867-74.
- 240. Baldwin ZK, Pearce BJ, Curi MA, et al. Limb salvage after infrainguinal bypass graft failure. J Vasc Surg. 2004;39:951-7.
- 241. Kannel WB, Skinner JJ, Schwartz MJ, et al. Intermittent claudication. Incidence in the Framingham Study. Circulation. 1970;41:875-83.
- 242. Adam DJ, Beard JD, Cleveland T, et al. Bypass versus angioplasty in severe ischaemia of the leg (BASIL): multicentre, randomised controlled trial. Lancet. 2005;366:1925-34.
- 243. Bradbury AW, Adam DJ, Bell J, et al. Multicentre randomised controlled trial of the clinical and costeffectiveness of a bypass-surgery-first versus a balloon-angioplasty-first revascularisation strategy for severe limb ischaemia due to infrainguinal disease. The Bypass versus Angioplasty in Severe Ischaemia of the Leg (BASIL) trial. Health Technol Assess. 2010;14:1-210, iii-iv.
- 244. Farber A, Rosenfield K, Menard M. The BEST-CLI trial: a multidisciplinary effort to assess which therapy is best for patients with critical limb ischemia. Tech Vasc Interv Radiol. 2014;17:221-4.
- 245. Acín F, Varela C, López de Maturana I, et al. Results of infrapopliteal endovascular procedures performed in diabetic patients with critical limb ischemia and tissue loss from the perspective of an angiosome-oriented revascularization strategy. Int J Vasc Med. 2014;2014:270539.
- 246. Söderstrom M, Albäck A, Biancari F, et al. Angiosome-targeted infrapopliteal endovascular revascularization for treatment of diabetic foot ulcers. J Vasc Surg. 2013;57:427-35.
- 247. Lejay A, Georg Y, Tartaglia E, et al. Long-term outcomes of direct and indirect below-the-knee open revascularization based on the angiosome concept in diabetic patients with critical limb ischemia. Ann Vasc Surg. 2014;28:983-9.
- Kret MR, Cheng D, Azarbal AF, et al. Utility of direct angiosome revascularization and runoff scores in predicting outcomes in patients undergoing revascularization for critical limb ischemia. J Vasc Surg. 2014;59:121-8.
- 249. Osawa S, Terashi H, Tsuji Y, et al. Importance of the six angiosomes concept through arterial-arterial connections in CLI. Int Angiol. 2013;32:375-85.
- 250. Rashid H, Slim H, Zayed H, et al. The impact of arterial pedal arch quality and angiosome revascularization on foot tissue loss healing and infrapopliteal bypass outcome. J Vasc Surg. 2013;57:1219-26.
- 251. Fossaceca R, Guzzardi G, Cerini P, et al. Endovascular treatment of diabetic foot in a selected population of patients with below-the-knee disease: is the angiosome model effective? Cardiovasc Intervent Radiol. 2013;36:637-44.
- 252. Kabra A, Suresh KR, Vivekanand V, et al. Outcomes of angiosome and non-angiosome targeted revascularization in critical lower limb ischemia. J Vasc Surg. 2013;57:44-9.
- 253. Azuma N, Uchida H, Kokubo T, et al. Factors influencing wound healing of critical ischaemic foot after bypass surgery: is the angiosome important in selecting bypass target artery? Eur J Vasc Endovasc Surg. 2012;43:322-8.
- 254. Varela C, Acín F, de Haro J, et al. The role of foot collateral vessels on ulcer healing and limb salvage after successful endovascular and surgical distal procedures according to an angiosome model. Vasc Endovascular Surg. 2010;44:654-60.

- 255. Neville RF, Attinger CE, Bulan EJ, et al. Revascularization of a specific angiosome for limb salvage: does the target artery matter? Ann Vasc Surg. 2009;23:367-73.
- 256. Alexandrescu VA, Hubermont G, Philips Y, et al. Selective primary angioplasty following an angiosome model of reperfusion in the treatment of Wagner 1-4 diabetic foot lesions: practice in a multidisciplinary diabetic limb service. J Endovasc Ther. 2008;15:580-93.
- 257. Iida O, Soga Y, Hirano K, et al. Long-term results of direct and indirect endovascular revascularization based on the angiosome concept in patients with critical limb ischemia presenting with isolated below-the-knee lesions. J Vasc Surg. 2012;55:363-70.
- 258. Bosanquet DC, Glasbey JC, Williams IM, et al. Systematic review and meta-analysis of direct versus indirect angiosomal revascularisation of infrapopliteal arteries. Eur J Vasc Endovasc Surg. 2014;48:88-97.
- 259. Biancari F, Juvonen T. Angiosome-targeted lower limb revascularization for ischemic foot wounds: systematic review and meta-analysis. Eur J Vasc Endovasc Surg. 2014;47:517-22.
- 260. Abu Dabrh AM, Steffen MW, Undavalli C, et al. The natural history of untreated severe or critical limb ischemia. J Vasc Surg. 2015;62:1642-51.
- 261. Gray BH, Laird JR, Ansel GM, et al. Complex endovascular treatment for critical limb ischemia in poor surgical candidates: a pilot study. J Endovasc Ther. 2002;9:599-604.
- 262. Ryer EJ, Trocciola SM, DeRubertis B, et al. Analysis of outcomes following failed endovascular treatment of chronic limb ischemia. Ann Vasc Surg. 2006;20:440-6.
- 263. Krankenberg H, Schlüter M, Steinkamp HJ, et al. Nitinol stent implantation versus percutaneous transluminal angioplasty in superficial femoral artery lesions up to 10 cm in length: the Femoral Artery Stenting Trial (FAST). Circulation. 2007;116:285-92.
- 264. Schillinger M, Sabeti S, Dick P, et al. Sustained benefit at 2 years of primary femoropopliteal stenting compared with balloon angioplasty with optional stenting. Circulation. 2007;115:2745-9.
- 265. Johnson WC, Lee KK. A comparative evaluation of polytetrafluoroethylene, umbilical vein, and saphenous vein bypass grafts for femoral-popliteal above-knee revascularization: a prospective randomized Department of Veterans Affairs cooperative study. J Vasc Surg. 2000;32:268-77.
- 266. Fogle MA, Whittemore AD, Couch NP, et al. A comparison of in situ and reversed saphenous vein grafts for infrainguinal reconstruction. J Vasc Surg. 1987;5:46-52.
- 267. Leather RP, Karmody AM. In-situ saphenous vein arterial bypass for the treatment of limb ischemia. Adv Surg. 1986;19:175-219.
- 268. Taylor LM, Edwards JM, Porter JM. Present status of reversed vein bypass grafting: five-year results of a modern series. J Vasc Surg. 1990;11:193-205.
- Nolan BW, De Martino RR, Stone DH, et al. Prior failed ipsilateral percutaneous endovascular intervention in patients with critical limb ischemia predicts poor outcome after lower extremity bypass. J Vasc Surg. 2011;54:730-5.
- 270. Santo VJ, Dargon P, Azarbal AF, et al. Lower extremity autologous vein bypass for critical limb ischemia is not adversely affected by prior endovascular procedure. J Vasc Surg. 2014;60:129-35.
- Uhl C, Hock C, Betz T, et al. Pedal bypass surgery after crural endovascular intervention. J Vasc Surg. 2014;59:1583-7.
- 272. Okadome J, Matsumoto T, Aoyagi Y, et al. Long-term results of a hybrid revascularization procedure for peripheral arterial disease. Fukuoka Igaku Zasshi. 2015;106:254-61.
- 273. Starodubtsev V, Karpenko A, Ignatenko P. Hybrid and open surgery of Trans-Atlantic Inter-Society II type C and D iliac occlusive disease and concomitant lesion of common femoral artery. Int Angiol. 2016;35:484-91.
- 274. Nishibe T, Maruno K, Iwahori A, et al. The role of common femoral artery endarterectomy in the endovascular era. Ann Vasc Surg. 2015;29:1501-7.
- 275. Kobayashi N, Hirano K, Nakano M, et al. Prognosis of critical limb ischemia patients with tissue loss after achievement of complete wound healing by endovascular therapy. J Vasc Surg. 2015;61:951-9.
- 276. Armstrong DG, Bharara M, White M, et al. The impact and outcomes of establishing an integrated interdisciplinary surgical team to care for the diabetic foot. Diabetes Metab Res Rev. 2012;28:514-8.
- 277. Chung J, Modrall JG, Ahn C, et al. Multidisciplinary care improves amputation-free survival in patients with chronic critical limb ischemia. J Vasc Surg. 2015;61:162-9.
- Moran PS, Teljeur C, Harrington P, et al. A systematic review of intermittent pneumatic compression for critical limb ischaemia. Vasc Med. 2015;20:41-50.
- 279. Abidia A, Laden G, Kuhan G, et al. The role of hyperbaric oxygen therapy in ischaemic diabetic lower extremity ulcers: a double-blind randomised-controlled trial. Eur J Vasc Endovasc Surg. 2003;25:513-8.
- Ruffolo AJ, Romano M, Ciapponi A. Prostanoids for critical limb ischaemia. Cochrane Database Syst Rev. 2010;CD006544.

- 281. Blaisdell FW. The pathophysiology of skeletal muscle ischemia and the reperfusion syndrome: a review. Cardiovasc Surg. 2002;10:620-30.
- 282. Londero LS, Nørgaard B, Houlind K. Patient delay is the main cause of treatment delay in acute limb ischemia: an investigation of pre- and in-hospital time delay. World J Emerg Surg. 2014;9:56.
- Duval S, Keo HH, Oldenburg NC, et al. The impact of prolonged lower limb ischemia on amputation, mortality, and functional status: the FRIENDS registry. Am Heart J. 2014;168:577-87.
- 284. Morris-Stiff G, D'Souza J, Raman S, et al. Update experience of surgery for acute limb ischaemia in a district general hospital—are we getting any better? Ann R Coll Surg Engl. 2009;91:637-40.
- 285. Saarinen E, Vuorisalo S, Kauhanen P, et al. The benefit of revascularization in nonagenarians with lower limb ischemia is limited by high mortality. Eur J Vasc Endovasc Surg. 2015;49:420-5.
- 286. Baril DT, Patel VI, Judelson DR, et al. Outcomes of lower extremity bypass performed for acute limb ischemia. J Vasc Surg. 2013;58:949-56.
- 287. Manojlović V, Popović V, Nikolić D, et al. Analysis of associated diseases in patients with acute critical lower limb ischemia. Med Pregl. 2013;66:41-5.
- 288. Comerota AJ, Weaver FA, Hosking JD, et al. Results of a prospective, randomized trial of surgery versus thrombolysis for occluded lower extremity bypass grafts. Am J Surg. 1996;172:105-12.
- Diffin DC, Kandarpa K. Assessment of peripheral intraarterial thrombolysis versus surgical revascularization in acute lower-limb ischemia: a review of limb-salvage and mortality statistics. J Vasc Interv Radiol. 1996;7:57-63.
- 290. Ouriel K, Veith FJ, Sasahara AA. A comparison of recombinant urokinase with vascular surgery as initial treatment for acute arterial occlusion of the legs. Thrombolysis or Peripheral Arterial Surgery (TOPAS) Investigators. N Engl J Med. 1998;338:1105-11.
- 291. Results of a prospective randomized trial evaluating surgery versus thrombolysis for ischemia of the lower extremity. The STILE trial. Ann Surg. 1994;220:266-8.
- 292. Ouriel K, Shortell CK, DeWeese JA, et al. A comparison of thrombolytic therapy with operative revascularization in the initial treatment of acute peripheral arterial ischemia. J Vasc Surg. 1994;19:1021-30.
- 293. Eliason JL, Wakefield TW. Metabolic consequences of acute limb ischemia and their clinical implications. Semin Vasc Surg. 2009;22:29-33.
- 294. Henke PK. Contemporary management of acute limb ischemia: factors associated with amputation and inhospital mortality. Semin Vasc Surg. 2009;22:34-40.
- 295. Ansel GM, Botti CF, Silver MJ. Treatment of acute limb ischemia with a percutaneous mechanical thrombectomy-based endovascular approach: 5-year limb salvage and survival results from a single center series. Catheter Cardiovasc Interv. 2008;72:325-30.
- 296. Gupta R, Hennebry TA. Percutaneous isolated pharmaco-mechanical thrombolysis-thrombectomy system for the management of acute arterial limb ischemia: 30-day results from a single-center experience. Catheter Cardiovasc Interv. 2012;80:636-43.
- 297. Silva JA, Ramee SR, Collins TJ, et al. Rheolytic thrombectomy in the treatment of acute limb-threatening ischemia: immediate results and six-month follow-up of the multicenter AngioJet registry. Possis Peripheral AngioJet Study AngioJet Investigators. Cathet Cardiovasc Diagn. 1998;45:386-93.
- 298. Taha AG, Byrne RM, Avgerinos ED, et al. Comparative effectiveness of endovascular versus surgical revascularization for acute lower extremity ischemia. J Vasc Surg. 2015;61:147-54.
- 299. Leung DA, Blitz LR, Nelson T, et al. Rheolytic pharmacomechanical thrombectomy for the management of acute limb ischemia: results from the PEARL Registry. J Endovasc Ther. 2015;22:546-57.
- 300. Zaraca F, Ponzoni A, Sbraga P, et al. Factors affecting long-term outcomes after thromboembolectomy for acute lower limb ischemia. Minerva Chir. 2012;67:49-57.
- 301. Ender Topal A, Nesimi Eren M, Celik Y. Management of non-traumatic acute limb ischemia and predictors of outcome in 270 thrombembolectomy cases. Int Angiol. 2011;30:172-80.
- Fogarty TJ, Cranley JJ, Krause RJ, et al. A method for extraction of arterial emboli and thrombi. Surg Gynecol Obstet. 1963;116:241-4.
- 303. Schrijver AM, Reijnen MM, van Oostayen JA, et al. Initial results of catheter-directed ultrasound-accelerated thrombolysis for thromboembolic obstructions of the aortofemoral arteries: a feasibility study. Cardiovasc Intervent Radiol. 2012;35:279-85.
- 304. Schrijver A, Vos J, Hoksbergen AW, et al. Ultrasound-accelerated thrombolysis for lower extremity ischemia: multicenter experience and literature review. J Cardiovasc Surg (Torino ). 2011;52:467-76.
- 305. Schrijver AM, Reijnen MM, van Oostayen JA, et al. Dutch randomized trial comparing standard catheterdirected thrombolysis versus ultrasound-accelerated thrombolysis for thromboembolic infrainguinal disease (DUET): design and rationale. Trials. 2011;12:20.

- 306. Jongsma H, Bekken JA, van Buchem F, et al. Secondary interventions in patients with autologous infrainguinal bypass grafts strongly improve patency rates. J Vasc Surg. 2016;63:385-90.
- 307. Carter A, Murphy MO, Halka AT, et al. The natural history of stenoses within lower limb arterial bypass grafts using a graft surveillance program. Ann Vasc Surg. 2007;21:695-703.
- Ihlberg L, Luther M, Albäck A, et al. Does a completely accomplished duplex-based surveillance prevent veingraft failure? Eur J Vasc Endovasc Surg. 1999;18:395-400.
- Westerband A, Mills JL, Kistler S, et al. Prospective validation of threshold criteria for intervention in infrainguinal vein grafts undergoing duplex surveillance. Ann Vasc Surg. 1997;11:44-8.
- 310. Lundell A, Lindblad B, Bergqvist D, et al. Femoropopliteal-crural graft patency is improved by an intensive surveillance program: a prospective randomized study. J Vasc Surg. 1995;21:26-33.
- 311. Mills JL, Harris EJ, Taylor LM, et al. The importance of routine surveillance of distal bypass grafts with duplex scanning: a study of 379 reversed vein grafts. J Vasc Surg. 1990;12:379-86.
- 312. Bandyk DF, Cato RF, Towne JB. A low flow velocity predicts failure of femoropopliteal and femorotibial bypass grafts. Surgery. 1985;98:799-809.
- 313. Back MR, Novotney M, Roth SM, et al. Utility of duplex surveillance following iliac artery angioplasty and primary stenting. J Endovasc Ther. 2001;8:629-37.
- 314. Baril DT, Marone LK. Duplex evaluation following femoropopliteal angioplasty and stenting: criteria and utility of surveillance. Vasc Endovascular Surg. 2012;46:353-7.
- 315. Troutman DA, Madden NJ, Dougherty MJ, et al. Duplex ultrasound diagnosis of failing stent grafts placed for occlusive disease. J Vasc Surg. 2014;60:1580-4.
- 316. Brumberg RS, Back MR, Armstrong PA, et al. The relative importance of graft surveillance and warfarin therapy in infrainguinal prosthetic bypass failure. J Vasc Surg. 2007;46:1160-6.
- 317. Calligaro KD, Doerr K, McAffee-Bennett S, et al. Should duplex ultrasonography be performed for surveillance of femoropopliteal and femorotibial arterial prosthetic bypasses? Ann Vasc Surg. 2001;15:520-4.
- 318. Stone PA, Armstrong PA, Bandyk DF, et al. Duplex ultrasound criteria for femorofemoral bypass revision. J Vasc Surg. 2006;44:496-502.
- 319. Subherwal S, Patel MR, Chiswell K, et al. Clinical trials in peripheral vascular disease: pipeline and trial designs: an evaluation of the ClinicalTrials.gov database. Circulation. 2014;130:1812-9.
- 320. Darling JD, McCallum JC, Soden PA, et al. Predictive ability of the Society for Vascular Surgery Wound, Ischemia, and foot Infection (WIfI) classification system following infrapopliteal endovascular interventions for critical limb ischemia. J Vasc Surg. 2016;64:616-22.
- 321. Causey MW, Ahmed A, Wu B, et al. Society for Vascular Surgery limb stage and patient risk correlate with outcomes in an amputation program. J Vasc Surg. 2016;63:1563-73.
- 322. Beropoulis E, Stavroulakis K, Schwindt A, et al. Validation of the Wound, Ischemia, foot Infection (WIfI) classification system in nondiabetic patients treated by endovascular means for critical limb ischemia. J Vasc Surg. 2016;64:95-103.
- 323. Zhan LX, Branco BC, Armstrong DG, et al. The Society for Vascular Surgery lower extremity threatened limb classification system based on Wound, Ischemia, and foot Infection (WIfI) correlates with risk of major amputation and time to wound healing. J Vasc Surg. 2015;61:939-44.
- 324. Mills JL, onte MS, rmstrong DG, et al. The Society for Vascular Surgery Lower Extremity Threatened Limb Classification System: risk stratification based on wound, ischemia, and foot infection (WIfI). J Vasc Surg. 2014;59:220-34.





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2016 AHA/ACC Guideline on the Management of Patients With Lower Extremity Peripheral Artery Disease Data Supplements (Section numbers correspond to the full-text guideline.)

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# **Methodology and Evidence Review**

and added to the evidence tables when appropriate. peripheral artery disease/peripheral arterial disease/peripheral vascular disease/lower extremity arterial disease, smoking/smoking cessation, statin, stenting, and diabetes, diagnostic testing, endovascular therapy, exercise rehabilitation/exercise therapy/exercise training/supervised exercise, lower extremity/foot wound/ulcer, were not limited to the following: acute limb ischemia, angioplasty, ankle-brachial index, anticoagulation, antiplatelet therapy, atypical leg symptoms, blood pressure 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 vascular surgery. Additional relevant studies published through September 2016, during the guideline writing process, were also considered by the writing committee. Research and Quality, and other selected databases relevant to this guideline, was conducted from January through September 2015. Key search words included but lowering/hypertension, bypass graft/bypass grafting/surgical bypass, cilostazol, claudication/intermittent claudication, critical limb ischemia/severe limb ischemia,

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<u>9546918</u> Criqui MH, et MM, et al. 1999(4) 10030313 VicDermot 1996(3) without PAD) general medical practice; 105 pts vascular lab; 26 known PAD from Size: n=268 pts (137 known PAD from pts with and without PAD administered Study type: Cross-sectional study of analysis) Size: n=508 pts (980 limbs for Study type: Cross-sectional study of community pts-also did a n=350; 50 vascular clinic pts and 300 ABI assessment San Diego Claudication questionnaire, tibial flow velocity modified WHO/ROSE questionnaire reproducibility study Questionnaire) vs. ABI/TBI/posterior (San Diego Claudication identified from (vascular. lab Pts with and without PAD Mean age 68 y noncompressible ABI >1.50 speakers, life expectancy <6 mo, extremity amputation, non-English chair bound, pts with major lower nursing home residents, wheel-Exclusion criteria: Low MMSE PAD defined as ABI <0.9</li> general medical clinics) Inclusion criteria: Exclusion criteria: N/A Severe PAD (ABI < 0.6) Moderate PAD (ABI 0.61–0.9) Isolated small vessel Optimal (no disease) characterize pts as: Vascular lab studies used to invited to participate Medical Center vascular labs at San Diego VA Hospital or UCSD Pts seen during preceding 10 y Isolated posterior tibial Borderline Normal Inclusion criteria: 4. Pain at rest 3. Atypical exertional leg symptoms clinic Among PAD pts (N=163). factors leg sx: older age, male sex, DM, PAD pt significantly associated absence of exertional Among PAD pts (n=163), factors 30.7% pain at rest. 25.5% atypical exertional leg symptoms 28.5% had IC (classic); vascular lab: Among N=137 PAD pts identified from 2. IC (classic) 1. No exertional leg symptoms based on San Diego Claudication Grouped pts according to 4 categories Rose calf claudication documented PAD, including no sx, pain at spectrum of clinical sx in pts with NPV: 99% community, 81% vascular clinic PPV: 91% community, 100% vascular vascular clinic than vascular lab recruited from general medicine clinic rather 15.3% had no exertional leg symptoms; Questionnaire: rest, noncalf pain, nonRose calf claudication, Results: Questionnaire identified wide Results: as both legs) and included buttock and confirmed PAD lower extremity sx among pts with leg sx in PAD symptoms—early concept of "atypical" Study recognized wider spectrum of at rest, non-calf, non-Rose calf and Questionnaire allows for more Further validated wider spectrum of Rose (calf). can be categorized as: No pain, pain variation of sx and pts leg symptoms left leg symptoms separately (as well Questionnaire accounts for right and San Diego Claudication WHO/Rose and also non-calf leg sx in PAD including leg sx not c/w thigh pain.

Hirsch AT, et al. 2001(6) <u>11560536</u>	McDermott MM, et al. 11585483
<b>Study type:</b> Multi-center cross- sectional study conducted at 350 primary care practices in the US. Pts enrolled underwent San Diego Claudication Questionnaire, medical and CV Hx/risk factor assessment, BP, anthropomorphics, and ABI assessment. Pts. identified as having PAD (and their providers) further asked about awareness of the PAD Dx.	<ul> <li>Study type: Cross-sectional study of pts with and without PAD identified from 3 medical centers in same city. Pts underwent functional capacity assessments (6min walk, 4 M walk, chair raises), assessment of physical activity, ABI, questionnaires</li> <li>Size: n=590 pts (460 with PAD; 130 without PAD)</li> </ul>
<ul> <li>Inclusion criteria:</li> <li>Age ≥70 y; Age 50–69 y with DM or at least 10 pack-year tobacco Hx</li> <li>PAD (lower leg pressure method) defined as ABI ≤0.9 in either leg</li> <li>Exclusion criteria: N/A</li> </ul>	<ul> <li>Inclusion criteria:</li> <li>Pts with and without PAD identified from 3 medical centers (vascular lab, general medical practice)</li> <li>PAD confirmed with study ABI (average leg pressure method) and required ABI &lt;0.9</li> <li>Exclusion criteria:</li> <li>"PAD" pts with normal ABI at study visit</li> <li>Dementia</li> <li>Nursing home residents</li> <li>Wheelchair bound</li> <li>Pts with major lower extremity amputation</li> <li>Recent major surgery</li> <li>Non-English speakers</li> </ul>
<ul> <li><u>Results:</u></li> <li>Prevalence of PAD in this cohort was 29%</li> <li>Among 1865 pts with PAD (mean ABI 0.78):</li> <li>5.5%-15.3% Rose claudication;</li> <li>46.3%-61.7% atypical leg sx;</li> <li>23.3%-48.3% no pain;</li> <li>**rates reported for new Dx/prior Dx and for PAD only and PAD+CVD</li> </ul>	<ul> <li>significantly associated with classical IC lower ABI, PAD recruited from vascular lab rather than general medicine clinic</li> <li>Results: Grouped pts according to 6 types of leg symptoms in 4 overall categories: 1. IC (classic)</li> <li>Atypical exertional leg pain (carry on/stop)</li> <li>No exertional leg pain (active/inactive walk &gt;6 blocks/wk Yes/No)</li> <li>Leg pain on exertion and at rest: 32% had IC;</li> <li>19% leg pain on exertion and at rest; 20% no exertional leg pain.</li> <li>PAD pts in the non-IC groups also demonstrated functional impairment in terms of 6 min walk, 4 meter walk.</li> <li>No exertional and rest pain groups with worse functional capacity than IC group.</li> <li>Atypical exertional leg pain/carry on group with better outcomes on 6 min walk than IC group.</li> </ul>
More data on wide spectrum of leg sx among pts with PAD; only approximately 5%–15% of ABI confirmed PAD pts have classic Rose claudication. Majority have atypical non-Rose leg sx or no leg pain.	<ul> <li>More data on wide spectrum of leg sx among pts with PAD and demonstration that functional impairment is common regardless of type of leg symptoms.</li> </ul>

	<ul> <li>Parental hx of PAD was strongly associated with severe prevalent PAD (OR: 2.91; 95%Cl: 1.33–6.40; p=0.008).</li> </ul>			
	<ul> <li>Parental hx of PAD was significant when adjusting for SBP, DBP, and dyslipidemia (OR: 1 83: 95% CI: 1 00–3 41: n=0 05)</li> </ul>			
	with severe prevalent PAD (OR: 2.42: 95% CI: 1.13–5.23: p=0.02).			
	<ul> <li>(OR: 1.83; 95% CI: 1.03–3.26; p=0.04)</li> <li>Family hx of PAD was strongly associated</li> </ul>		<u>Size</u> : n=2,404 pts	
	<ul> <li>Family nx of PAD was significant, when adjusting for SBP, DBP, and dyslipidemia</li> </ul>	Exclusion criteria: N/A	volunteers and their significant others.	
	Results:	San Diego Population Study	California, San Diego, and their	
	population	completed the baseline visit in the	employees of the University of	2011(9) 21920269
N/A	1° endpoint: Prevalence of PAD in the study	Inclusion criteria: Men and	Study type: Observational population-	Wassel et al.
for AAA also had PAD.	PAD in 10.9%.		screened for AAA. 18,749 attended the screening (uptake 74.7%).	
<ul> <li>4.0% to 3.3%.</li> <li>10.9% of men undergoing screening</li> </ul>	Results: AAA was diagnosed in 3.3% and	Exclusion criteria: N/A	Size: n=25,083 men who were	2010(0) 25923784
The prevalence of AAA in Denmark     has declined in the past decade from	<u>1° endpoint:</u> Prevalence of PAD in pts screened for AAA.	Inclusion criteria: Men age 65–74 y who were screened for AAA.	Study type: Danish intervention arm of screening trial	Grøndal N, et al.
		Exclusion criteria: N/A		
		<ul> <li>17 studies met inclusion criteria (11 on diagnostic accuracy)</li> </ul>		
		Data could be extracted into a 2     x 2 table		
		angiogram		
		<ul> <li>PAD Dx confirmed by reference standard: ABL dunley or</li> </ul>		
	(LR: 0.57; 95% CI: 0.43–0.76))	<ul> <li>Exam maneuvers nad to be described clearly</li> </ul>	מומטוויסטוני מיימו מיץ סומטופס	
	likelihood of any PAD, but lowered likelihood	databases	Size: I otal of 6,272 pts in 11	
moderate to severe PAD.	<ul> <li>Absence of claudication did not lower</li> </ul>	from MEDLINE and Cochrane		
of PAD, but lowers likelihood of	(LR PAD: 3.30; 95% CI: 2.30-4.80)	<ul> <li>51 potential articles identified</li> </ul>	PAD in pts with and without disease	16449619
וגפווווטע טו ראט. אטצפועפ טו claudication does not lower likelihood	<ul> <li>Presence of claudication ↑ likelihood PAD</li> </ul>	Studies published from 1/1900-     Studies published from 1/1900-	and/or physical examination for Dx of	2006(7)
Presence of claudication increases	Results:	Inclusion criteria:	Study type: Systematic review of	Khan NA, et
			Size: n=6,979 (1865 had PAD)	

Ч,	Significant association of PAD at cutoff of 15 mmHg (RR: 1.91; 95% CI: 1.28–2.84;	<ul> <li>Studies that did not report a dichotomous outcome defined by a</li> </ul>					
specificity. 91.9%, 90% CI.	83.1–96.3)	Exclusion criteria:					
= 10 IIIIIII ig and FAD (NN: 2.22, 30 /01. 1.41–3.5; p=0.0006) (sensitivity: 16.6%; 95% OF 6 7 35 // /sensitivity: 01 0%: 05% OF	1.41–3.5; p=0	indices or outcomes.					
Significant association between IASBPD of mmUc and DAD (DD: 2.32: 05% CI:	Significant a	subclavian stenosis, survival or					
	Results:	<ul> <li>Studies reporting CAD, cerebrovascular disease, PAD,</li> </ul>					
stenosis, all-cause, and CV mortality	stenosis, all-ca	simultaneously in arms or legs		<u>26160261</u>			
cerebrovascular disease, subclavian	cerebrovascular	<ul> <li>Studies measuring BP</li> </ul>	studies	2015(11)			
1° endpoint: Prevelance of PAD, CAD,	1° endpoint: Pr	Inclusion criteria:	Study type: Meta-analysis of cohort	Singh S et al.,			
32%; 95%CI: 23-41) (specificity: 91%, 95% CI: 86-94)	32%; 95%Cl: Cl: 86–94)						
• Significant association of $\geq 10$ mmHg and DVD (DD: 2.4:05% CI: 1.5:2.3) (constituity:	Significant						
cause mortality (HR: 1.6; 95% CI: 1.1–2.3	cause morta						
association in noninvasive	<ul> <li>Significant association studies of a difference of</li> </ul>						
- - -	1.1-2.5)						
cardiovascular mortality (HR: 1.7, 95% CI:	cardiovasculi						
association in noninvasive lifference of ≥15 mmHa and	<ul> <li>Significant association studies of a difference of</li> </ul>						
	97)						
95% CI: 2-26) (specificity: 93%; 95% CI: 86-	95% CI: 2–26						
ratio: 1.6, 95% CI: 1.1–2.48) (sensitivity: 8%;	ratio: 1.6, 95% (						
studies of a difference of < to mining and	pre-existing cer						
sociation in noninvasive	Significant association						
%; 95% CI: 9–23) (specificity: 94–98)	(sensitivity: 15%; 95% Cl 96%; 95% Cl: 94–98)						
PVD (risk ratio: 2.5, 95% CI: 1.6–3.8)	PVD (risk ratio: 2.5, 95%	Case reports					
Sociation in noninvasive	<ul> <li>Significant association</li> </ul>						
	3-6-21.2)	PVD, or death					
d SS (risk ratio: 8.8; 95% CI:	≥10 mmHg and SS (risk	<ul> <li>Data for central vascular disease,</li> </ul>					
	<ul> <li>Significant acc</li> </ul>			22293369			
	Results:	of differences in BP between arms	Size: n=20 studies	2012(10)			
		<ul> <li>Cohort or cross-sectional studies</li> </ul>	and the more analysis				
	10 and naint: D\/D	Inclusion criteria:	Study type: Mata-analysis	Clark CE of			
				15358030	Shadman R et al., 2004(12)	Shadman R	
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				(2,975 from 2 free-living cohorts and 1,248 from 2 clinical cohorts)	Study type: Review of cohort studies Size: n=4 cohorts with 4.223 pts	Study type: Review of cohort studies	
	Cohort D: • Pts who visited the San Diego Vererans Administration Medical Centor or UCSD Medical Center vascular laboratories between 1990–1994	Cohort C: • Pt population in Chicago	Cohort B: • Randomly selected from a database of UCSD employees and spouses	<ul> <li>study</li> <li>Part of the Lipid Research Clinics protocol study</li> </ul>	<ul> <li>Inclusion criteria:</li> <li>Cohort A:</li> <li>Geographic defined population</li> </ul>	Inclusion criteria:	specific BP difference cutoff
			<ul> <li>Significant association between higher levels of HDL and SS (OR: 0.87 per 10 mg/dl)</li> <li>Significant association of SS and PAD (OR: 5.11 pc0.001)</li> </ul>	<ul> <li>SS was significantly (p&lt;0.05) associated with past smoking (OR: 1.80), current smoking (OR: 2.61), and higher levels of SRP (OR-1 90 per 20 mm Ho)</li> </ul>	1° endpoint: Prevelance of SS Results:	<ul> <li>Significant association between inter-leg BP difference of ≥15 mmHg and PAD (RR: 11.87; 95% CI: 7.64–18.44).</li> <li>IASBPD of ≥10 mmHg was not associated with carotid-femoral PWV (standardized mean difference: 0.26; 95% CI: 0.15–0.68; p=0.21). One study demonstrated positive association between IASBPD of ≥10 mmHg and brachial ankle PWV (adjusted OR from multivariate model: 1.001; 95% CI: 1.000–1.001; p=0.022).</li> <li>Significant association of inter-leg BP difference of ≥15 mm Hg or more and brachial–ankle PWV (standardized mean difference: 0.68; 95% CI: 0.37–0.99; p=0.0001).</li> </ul>	p=0.001) (sensitivity: 25.1%; 95% Cl 7.9– 56.7) (specificity: 88.2%; 95% Cl: 71.7–95.7).
0			hypertension or PAD diagnosis because of unilateral pressure measurement in an obstructed arm	<ul> <li>bilateral brachial BP measurements should routinely be performed in pts with an elevated risk profile, both to screen for SS and to avoid mission a</li> </ul>	<ul> <li>SS is correlated with current and past smoking histories, SBP, HDL levels (inverselv), and the presence of PAD</li> </ul>	•SS is correlated with current and past	

	Khan NA et al. 2006(7) <u>16449619</u>	Study Acronym; Author; Year Published	ABI indicates anl inter-arm systolic ratio; PAD, perip pressure; SS, su	
	Study type: Systematic review of studies that evaluated element of Hx and/or physical examination for Dx of PAD in pts with and without disease Study size: n=6,272 pts in 11 diagnostic accuracy studies	Study Type/Design; Study Size	kle-brachial index; BP, blc blood pressure; IC, intern heral artery disease; PPV bclavian artery stenosis; [¬] bclavian artery stenosis; [¬]	
A maximum Long to have been and Amazing Calloga of Constitution.	<ul> <li>Inclusion criteria:</li> <li>Studies published from 1/1966– 3/2005</li> <li>51 potential articles identified from MEDLINE and Cochrane databases</li> <li>Exam maneuvers had to be described clearly</li> <li>PAD Dx confirmed by reference standard: ABI, duplex, or angiogram</li> <li>Data could be extracted into a 2 x 2 table</li> <li>17 studies met inclusion criteria</li> </ul>	Patient Population	Cohort A: Missing data Cohort B: N/A Cohort C: • Wheelchair bound • Hx Foot or leg amputations • Nursing home residents • Non-English speaking • Hx dementia • Non-English speaking • Non-English sp	Exclusion criteria:
) }	Results:         Physical Examination         Skin changes         Skin cool to touch in affected leg:         • LR PAD: 5.90; 95% Cl 4.10–8.60         Leg wound/sore:         • LR PAD: 5.90; 95% Cl: 2.60–13.40         Discolored skin:         • LR PAD: 2.80; 95% Cl: 2.40–3.30         Absence of cool skin, wound/sore did not lower         likelihood of PAD         Bruits         Presence of ≥1 bruit	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Cohort A: Missing data         Cohort B: N/A         Cohort C:         • Wheelchair bound         • Nursing home residents         • Non-English speaking         • Hx dementia         • Indicates ankle-brachial index; BP, blood pressure; CI, confidence interval; CV, cardiovascularular; DBP, diastolic blood pressure; GP, general practitioner; HR, hazard ratio; IASBPD, inter-arm systolic blood pressure; IC, intermittent claudication; LR, likelihood ratio; MMSE, Mini-Mental State Examination; N/A, not applicable; NPV, negative predictive value; OR, odds ratio; PAD, peripheral artery disease	
0	<ul> <li>In general, presence of physical findings increases likelihood of PAD</li> <li>Entirely normal pulse exam and absence of any bruits decrease likelihood of PAD</li> <li>Sensitivities/specificities not reported in this review</li> </ul>	Summary/Conclusion Comment(s)	⁷ , general practitioner; HR, hazard ratio; IASBPD, able; NPV, negative predictive value; OR, odds city; RR, relative risk; SBP, systolic blood 40, World Health Organization.	

	Cournot M et al. 2007(13) <u>18154997</u>			
<ul> <li>orbsector underweinten physical examination followed by</li> <li>vascularular studies (carotid, femoral ultrasound, ABI)</li> <li>Physical</li> <li>Physical</li> <li>examination included pulse assessment (present/absent), bruit assessment using the</li> </ul>	<ul> <li>Study type:</li> <li>Part of the EVADEC, prospective cohort study (cross-sectional analysis). Pts with no known vascularular disease underwent</li> </ul>			
	Inclusion criteria: • 18–90 y (mean age 52 y) • No known CVD • Asx Exclusion criteria: CV disease identified by modical record review			(11 on diagnostic accuracy) <u>Exclusion criteria</u> : N/A
<ul> <li>+LR ipsilateral ABI &lt;0.9: 1.00, 95% CI: 1.00–3.01</li> <li>-LR ipsilateral ABI &lt;0.9: 0.94; 95% CI: 0.88–1.01</li> <li>Absent DP pulse</li> <li>+LR ipsilateral ABI &lt;0.9: 2.01; 95% CI: 1.17–3.45</li> <li>-LR ipsilateral ABI &lt;0.9: 0.94; 95% CI: 0.88–1.00</li> <li>Absent DP+PT</li> <li>+LR ipsilateral ABI &lt;0.9: 3.57; 95% CI: 1.93–6.60</li> <li>-LR ipsilateral ABI &lt;0.9: 0.93; 95% CI: 0.97–1.00</li> <li>Interaction term for DM not significant</li> <li>Interobserver agreement 97% for femoral bruit; 92%</li> <li>PT palpation; 92% DP palpation</li> </ul>	Results         14.5% of pts had any bruit or absent PT/DP pulse         Femoral bruit         • +LR ipsilateral ABI <0.9: 2.90; 95% CI: 1.63–5.16	<b>Capillary Refil</b> Abnormal capillary refill time LR PAD: 1.90; 95% CI: 1.20–3.20 Prolonged venous refill LR mod/sev PAD: 3.60; 95% CI: 1.90–6.80 Normal venous refill time not informative to r/o PAD	<ul> <li>Absent/reduced</li> <li>*any=femoral/popliteal/DP/PT</li> <li>Absence of any pulse abnormality:</li> <li>LR PAD: 0.38; 95% CI: 0.23–0.64</li> <li>Abnormal dorsalis pedis pulse less diagnostically useful than abnormal femoral or PT pulse</li> <li>DP not palpable in 8.1% of healthy pts</li> <li>PT not palpable in 2.9% of healthy pts</li> </ul>	<ul> <li>LR PAD: 5.60; 95% CI: 4.70–6.70 Over iliac, femoral, popliteal artery Absence of a bruit over all 3 arteries</li> <li>LR PAD: 0.39; 95% CI: 0.34–0.45 Pulse Palpation Any* pulse abnormality</li> <li>LR PAD: 4.70; 95% CI: 2.20–9.90</li> </ul>
	Both presence of femoral bruit and absent pulses increase likelihood of PAD in asx pts without known PAD/CVD			

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	Armstrong DW et al. 21165366 AE be cli pt	bel pts siz
	Study type: Retrospective database analysis of pts who underwent ABI and had a physical examination documented in the CARDIOfile database between 12.2005– 2.2010 at a single clinic Size: n=1,236 eligible pts with complete data	bell of stethoscope <u>Size:</u> n=2,736 eligible pts <u>Interobserver</u> <u>variability substudy</u> <u>size:</u> 500 pts
<ul> <li>Femoral bruits present/absent</li> <li>Claudication=leg sx with exercise gone within 5 min of rest.</li> </ul>	Inclusion criteria: Pts who had ABI performed for suspected PAD or risk factors for PAD (Age >70 y, DM or smokers ages 50–69 y, intermediate Framingham Risk score) Exclusion criteria: Pts with ABI >1.30 in either leg; incomplete physical examination in the databse Definitions • PAD defined as ABI ≤0.9 • Pulses rated 0-3 scale; analysis absent vs. present	
<ul> <li>Sens 63.9%, Spec 80.6%</li> <li>PPV 43.2%, NPV 90.7%, Accuracy 77.5%</li> <li>+LR PAD 3.3</li> <li>-LR PAD 0.45</li> <li>Absent DP and PT pulses+femoral bruit either side (vs. normal pulses, no femoral bruits)</li> <li>Sens 58.2%, Spec 98.3%</li> <li>PPV 81%, NPV 94.9%, Accuracy 93.8%</li> <li>+LR PAD 34.2</li> <li>-LR PAD 0.43</li> </ul>	Results:           28.1% of pts had an abnormal ABI in at least 1 leg (PAD)           Femoral bruit           • Sens 36.1%, Spec 92.0%           • PPV 51.1%, NPV 86.2%, Accuracy 81.6%           • LR PAD 4.5           • LR PAD 0.69           PT pulse abnl           • Sens 70.0%, Spec 83.4%           • PPV 49.3%, NPV 92.3%, Accuracy 80.9%           • LR PAD 0.36	Also reported on carotid bruit for Dx of carotid stenosis/plaque/increased IMT (did not affect LR)
	<ul> <li>Completely normal exam (all ankle pulses present and no femoral bruits) has high accuracy for normal ABI/no PAD.</li> <li>Pulse abnormalities+femoral bruits makes Dx of PAD likely.</li> <li>Single abnormal physical findings increased likelihood of abnormal ABI (specific findings)</li> <li>Sensitivity of single abnormal physical examination findings lower; not as "reassuring" to rule out PAD/abnormal ABI</li> </ul>	

LR, likelihood ratio; PPV, positive predictive value; PAD, peripheral artery disease; PT, posterior tibial; pt, patient; OR, odds ratio; RR, relative risk; sens, sensitivity; and spec, specificity.

Evidence Table Study Acronym;	Aim of Study;	Evidence Table 3. RCTs of Resting ABI for Diagnosing PAD–Section 3.1.         Study Acronym;       Aim of Study;         Patient Population       Study In	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Autnor; Year Published	Study Type; Study Size (N)		(# patients) / Study Comparator (# patients)	(Absolute Event Rates, P value; OR or RR; & 95% CI)	Study Limitations; Adverse Events
Fowkes FG et al. 2010(15) <u>20197530</u>	<u>Aim</u> : To determine the effectiveness of ASA in preventing events in	Inclusion criteria: Men and women age 50–75 y	Intervention: 100 mg enteric coated ASA	<u>1° endpoint</u> : Composite of initial fatal or nonfatal coronary event, stroke or revascularization. (ASA: 13.7; 95% CI:	<ul> <li>Initial vascular events defined as a composite of a 1° endpoint event or angina, IC, orTIA.</li> </ul>
	people with a low ABI identified on screening	<ul> <li>Exclusion criteria:</li> <li>Previous Hx of vascular</li> </ul>	Comparator:	11.8–15.9 vs. placebo: 13.3; 95% Cl: 11.4–15.4, events per 1,000	ASA: 22.8; 95% Cl: 20.2–25.6 vs. placebo: 22.9; 95% Cl: 20.3–25.7
	the general population	<ul> <li>disease, MI, or stroke;</li> <li>Currently taking ASA or</li> </ul>	Placebo	person-y; HR: 1.03; 95% Cl: 0.84– 1.27	events per 1,000 person-y; HR: 1.00; 95% Cl: 0.85–1.17
	Study type: RCT	warfarin.			<ul> <li>All-cause mortality</li> </ul>
	<u>Size</u> : n=3,350 pts			1° Safety endpoint: Maior Hemorrhage: ASA: 2.5; 95% CI:	ASA group, 176 deaths (12.8; 95% Cl: 11.0–14.8 per 1.000 person-v):
				1.7–3.5 vs. placebo: 1.5; 95% Cl: 0.9– 2.3 ner 1 000 nerson-v [.] HR [.] 1 71 [.] 95%	placebo group, 186 deaths (13.5; 95% Cl: 11 6–15 6 per 1 000
				CI: 0.99–2.97	person-y; HR: 0.95; 95% Cl: 0.77– 1.16)
					<ul> <li>Limitations: higher proportion of women inclusion of nts with DM</li> </ul>
					could have influenced results
POPADAD	Aim: To determine	Inclusion criteria: Age ≥40 y	Intervention and	<u>1° endpoint</u> :	Adverse effect (effect estimates):
Belch J et al.	whether ASA and	ABI of <0.00 but no Sy CVD	100 mg ASA tablet	Death from CHD or stroke, nonfatal	<ul> <li>Malignancy 0.76 (0.52–1.11),</li> <li>Claboding 0.00 (0.52–1.52)</li> </ul>
<u>18927173</u>	combined or alone,		+ antioxidant	ankle for CLI; and death from CHD or	<ul> <li>Dyspepsia 0.77 (0.55–1.08),</li> </ul>
	are more effective	Exclusion criteria: People with:	capsule (n=320);	stroke	• Allergy 1.14 (0.80–1.63)
	reducing CVD events	ASA or antioxidant therapy use	capsule (n=318);	<ul> <li>Tib of 638 T² events in the ASA groups compared with 117 of 638 in</li> </ul>	
	in pts with DM and Asx PAD.	on a regular basis; peptic ulceration. severe dvspepsia. a	placebo tablet + antioxidant capsule	the no ASA groups (18.2% vs. 18.3%)	
	-	bleeding disorder, or intolerance	(n=320); or placebo	deaths from CHD or stroke occurred in	
	Multicenter,	to ASA; suspected serious physical illness (e.g., cancer),	tablet + placebo capsule (n=318).	the ASA groups compared with 35 in the no ASA groups (6 7% vs. 5 5%)	
	randomized, double	which could curtail life		HR: 1.23; 95% CI: 0.79–1.93).	
	placebo controlled	expectancy; psychiatric liness (reported by GP); pts with		<ul> <li>No difference in treatment for ABI</li> <li>&lt;0 90</li> </ul>	
	trial.	congenital heart disease; and			
		Pre missie is give mission			

20					
<ul> <li>Majority of pt with PAD have ≥1</li> </ul>		10.8%-18.2%)			
blacks.	<ul> <li>Prevalence of PAD in adults ≥70 y in U.S. was 14.5% (95% Cl:</li> </ul>	<ul> <li>Prevalence of PAD in</li> </ul>	index		
age and disproportionately affects		3.1%-5.5%)	mean ABI blood pressure		
<ul> <li>PAD prevalence increases with</li> </ul>	adults ≥40 y in U.S. was 4.3% (95% CI:	<ul> <li>Prevalence of PAD in adults ≥40 y in U</li> </ul>	<ul> <li>Participants with valid</li> </ul>		
million adults.		Results:	2000 NHANES	<b><u>Size:</u></b> n=2,1/4 pts	
<ul> <li>In the U.S., PAD affects &gt;5</li> </ul>			<ul> <li>Participants of 1999–</li> </ul>	2	15262830
either leg	nent intensity	of diagnosis, and treatment intensity	• Age ≥40 y	sectional survey	2004(19)
<ul> <li>PAD defined as ABI &lt;0.90 in</li> </ul>	y of detection, pt and physician awareness	1° endpoint: Frequency of detection, pt	Inclusion criteria:	Study type: Cross-	Selvin E, et al.
ערייא (ידיא //)	<ul> <li>Inspanies and Asians has a lower put nonsignificant lower FAD</li> <li>prevalence than Whites</li> </ul>	prevalence than Whites	Exclusion criteria: N/A		
of PAD		(UR: 2.30; p>0.024)	Asian		
Tactors to explain increased risk	<ul> <li>Blacks had a higher PAU prevalence than Non-Hispanic Whites</li> </ul>	Blacks had a higher H	Whites, blacks, Hispanics,		
higher susceptibility to CV risk		• 104 PAD cases (4.4%)	ethnicities: Non-Hispanic	Size: 2,343 pts	
<ul> <li>No evidence of blacks being of</li> </ul>		Results:	<ul> <li>1 of the following</li> </ul>		<u>16246968</u>
risk factor for PAD			• Age 29–91 y	sectional study	2005(18)
<ul> <li>Suggests black ethnicity is a</li> </ul>	ralence	1° endpoint: PAD prevalence	Inclusion criteria:	Study type: Cross-	Criqui MH, et al.
	& 95% CI)				Year Published
Comment(s)	(include P value; OR or RR;	(incl		Study Size	Author;
Summary/Conclusion	Primary Endpoint and Results	Prima	Patient Population	Study Type/Design;	Study Acronym;
D–Section 3.1.	Observational Studies, and/or Registries of Resting ABI for Diagnosing PAD–Section 3.1.	s, and/or Registries	ials, Observational Studie	Evidence Table 4. Nonrandomized Trials,	Evidence Table
schemic attack	ndomized controlled trial; and TIA, transient ischemic attack	x, symptomatic; RCT, ra	physical component summary score; PFWT, pain-free walking time; pt, patient; Sx, symptomatic; RCT, randomized controll	summary score; PFWT, pa	physical component
, cardiovascular disease; CHD, coronary heart disease; GI, , mean walking time; PAD, peripheral artery disease; PCS,	1° indicates primary; ABI, ankle-brachial index; ASA, aspirin; Asx, asymptomatic; CI, confidence interval; BL, baseline; CVD, cardiovascular disease; CHD, coronary heart disease; GI gastrointestimal; HR, hazard ratio, Hx, history; IC, intermittent claudication; MCS, mental component summary score; MWT, mean walking time; PAD, peripheral artery disease; PCS,	CI, confidence interval; f mental component sumi	ASA, aspirin; Asx, asymptomatic; C, intermittent claudication; MCS,	ABI, ankle-brachial index; , hazard ratio, Hx, history; I	1º indicates primary; gastrointestimal; HR,
			• CLI		
	mo		PAD		
	<ul> <li>Intervention: 372 m BL vs. 386 m 6</li> </ul>		<ul> <li>Walking impairment not from</li> </ul>		
	<ul> <li>Control: 347 m BL vs. 329 m 6mo</li> </ul>		sessions	claudication	
	6-MWT:		<ul> <li>Inability to attend weekly</li> </ul>	ŭt	
	Results:		stopping	<u>i</u>	
		Health education	<ul> <li>Inability to walk ≥50 ft without</li> </ul>		
	SF-36	Comparator:	<ul> <li>Lower extremity amputation</li> </ul>	claudication	
	activity; WIQ scores; PCS and MCS of		Exclusion criteria:	PAD with and without	
	treadmill MWT; PFWT; physical	group		vs. control in pts with	
		behavioral walking	ABI 20.9 of 20 % post exercise     dron in ABI	exercise intervention	23821089
vercise in nte with Asy PAD		mediated cognitive	ABI <0 0 or 20% most avarcisa	home-based walking	2013(17)
<ul> <li>wodest induovenient in o-wwwi</li> <li>distance affer 6 mo of home-based</li> </ul>	between baseline and 6 mo	Home-based aroup-	<ul> <li>Age ≥65 v</li> </ul>	testing efficacy of a	et al.
NALI - LI	hongo in G MM/T	Infommation	Inatuaian avitavia:	Ctudu tuna DOT	NNN + NNN

Aboyans V, et al.	Guo X, et al. 2008(20) <u>18362433</u>	Hirsch AT, et al. 2001(6) <u>11560536</u>	
Study type: Scientific	<u>Study type:</u> Observational test comparison <u>Size</u> : n=298 pts	<ul> <li>Study type:</li> <li>Multi-center cross- sectional study conducted at 350 primary care practices in the US.</li> <li>Pts enrolled underwent San Diego Claudication Questionnaire, medical and CV Hx/risk factor assessment. BP, anthropomorphics, and ABI assessment.</li> <li>Pts. identified as having PAD (and their providers) further asked about awareness of the PAD Dx.</li> <li>Size: n=6,979 pts (1,865 had PAD)</li> </ul>	
Inclusion criteria: N/A	Inclusion criteria: • Age ≥35 y • Cardiology clinic: referrals for DSA & ABI Exclusion criteria: Severe DM & hypertension Gold standard: • DSA. • Stenosis ≥50% ABI method: Oscillometry	<ul> <li>Inclusion criteria:</li> <li>Age ≥70 y or age 50–69 y with DM or Hx of ≥10 pack-year tobacco</li> <li>PAD (lower leg pressure method) defined as ABI ≤0.9 in either leg</li> <li>Exclusion criteria: N/A</li> </ul>	<ul> <li>Exclusion criteria:</li> <li>ABI values &gt;1.5</li> <li>Participants with missing variables of interest</li> </ul>
<u>1° endpoint</u> : N/A	<ul> <li><u>1° endpoint</u>: Presence of stenosis below aorto-iliac bifurcation in leg with lower ABI</li> <li><u>Results</u>: <ul> <li>Sensitivity: 76 (N/A)</li> <li>Specificity: 90 (N/A)</li> <li>PPV: 36 (N/A)</li> <li>NPV: 98 (N/A)</li> </ul> </li> </ul>	<ul> <li>Results:</li> <li>Prevalence of PAD in this cohort was 29%</li> <li>Among 1,865 pts with PAD (mean ABI 0.78):</li> <li>5.5–15.3% Rose claudication;</li> <li>46.3–61.7% atypical leg sx;</li> <li>23.3–48.3% no pain</li> <li>**Rates reported for new Dx/prior Dx and for PAD only and PAD+CVD</li> </ul>	<ul> <li>Black race/ethnicity (OR: 2.83; 95% CI: 1.48–5.42); current smoking (OR: 4.46; 95% CI: 2.25–8.84), DM (OR: 2.27; 95% CI: 1.03–7.12), hypertension (OR: 1.74; 95% CI: 0.97–3.13), hypercholesterdemia (OR: 1.68; 95% CI: 1.09–2.57) and low kidney function (OR: 2.00; 95% CI: 1.08–3.70) were positively associated with PAD prevalence.</li> </ul>
AHA Scientific Statement on the	<ul> <li>Moderate sensitivity and good specificity. No indication of % with PAD symptoms but low prevalence of PAD on DSA (7%) suggests it was negligible.</li> <li>53% had coronary heart disease and 13% stroke.</li> </ul>	<ul> <li>More data on wide spectrum of leg sx among pts with PAD; only about 5-15% of ABI confirmed PAD pts have classic Rose claudication. Many majority have atypical non-Rose leg sx or no leg pain.</li> </ul>	CVD risk factor. • Low Prevalence of PAD: 4.3%; 95% Cl: 3.1%–5.5%

		Allen J, et al. 1996(25) <u>8638864</u>		<u>9//789671</u>	Premalatha G, et al. 2002(24)				<u>16950430</u>	Schröder F, et al. 2006(23)		Aboyans V, et al. 2008(22) <u>18692981</u>	2012(21) <u>23159553</u>
	<u>Size</u> : n=200 pts	Study type: Observational test comparison		<u>Size</u> : n=100 pts	Study type: Observational test comparison			Size: n=216 pts	comparison	Study type: Observational test	<u>Size</u> : n=510 pts	<u>Study type</u> : Cross- sectional	statement <u>Size</u> : N/A
Gold standard: Duplex	Exclusion criteria: Previous vascular surgery. DM	Inclusion criteria: Consecutive referrals to a vascular laboratory.	Gold standard: Duplex ultrasound	Exclusion criteria: Calcification of peripheral arteries	Inclusion criteria: Pts with DM with foot lesions	Gold standard: Duplex ultrasound	Exclusion criteria: Previous evidence of PAD, obesity, atrial fibrillation, ABI > 1.3	disease. Age >40 y	medicine clinic "suspected	Inclusion criteria: Attending a vascular	Exclusion criteria: N/A	Inclusion criteria: ambulatory pts presenting to vascular lab	Exclusion criteria: N/A
	<ul> <li>Sensitivity: 0.82</li> <li>Specificity: 0.84</li> <li>PPV: 1.0</li> <li>NPV: 0.83</li> </ul>	<u>1° endpoint</u> : Stenosis >50% <u>Results:</u>		<ul> <li>Sensitivity: 0.71</li> <li>Specificity: 0.89</li> </ul>	<u>1° endpoint</u> : Precise criteria for PAD not stated. Results:		• PPV 0.99;0.93 • NPV: 0.74;0.88	Sensitivity: 0.09;0.93     Dov 0.000	Results: High;Low of post/ant tibial arteries	<u>1° endpoint</u> : Stenosis >70%	Results: In 84.2% of cases, diabetic limbs with ABI ≥1.40 had abnormal results in at least 1 of the 2 noninvasive vascular indicators, suggestive of concomitant occlusive disease.	<u>1° endpoint</u> : Association of risk factors with ABI >1.4 and ABI <0.9 and disease presence by TBI	Results: N/A
	<ul> <li>ABI had good sensitivity and specificity and excellent PPV.</li> </ul>	<ul> <li>Pt symptoms not presented in detail but it would appear that most were sx pts referred for</li> </ul>			Study in pts with DM with clinical suggestion of PAD showing good sensitivity and high specificity.			serisiuvity.	lower ankle pressure improved	ABI had good sensitivity and very high specificity and PPV. Using		<ul><li>50% with DM</li><li>No angiographic correlations</li></ul>	measurement and interpretation of the ABI

Major coronary events:
<ul> <li>CV mortality: Men: 0.71; 95% CI: 0.65–0.76; Women: 0.65; 95% CI: 0.58–0.72</li> <li>Prediction NPI scores:</li> </ul>
<ul> <li>Women: 0.45; 95% Cl: 0.38–0.52.</li> <li>Adding ABI to FRS improves men's scores modestly and women's scores substantially. Major coronary events: Men: 0.69; 95% Cl: 0.61–0.76;</li> <li>Women: 0.069; 95% Cl: 0.61–0.076.</li> </ul>
Men: 0.67, 93% CI: 0.6–0.74; Women: 0.58; 95% CI: 0.49–0.66 • CV mortality: Men: 0.68; 95% CI: 0.63–0.74;
<ul> <li>Results:</li> <li>C index for major coronary events, FRS only:</li> </ul>
and NRI score
<u>1° endpoint</u> : C index (fraction of occasions where the predictor score correctly predicts the earlier event for a pair of individuals)
of women would change risk category with ABI added to FRS.
<ul> <li>• CV mortality HR for different ABI levels: Reference=1.11–1.20; ABI ≤0.60=5.58 for men: 7.04 for women. 19% of men and 36 %</li> </ul>
<ul> <li>Follow-up ranged from 3-6.7 y; 9924 (25% CVD) deaths during 480,325 person-years of follow-up.</li> </ul>
Results:
1° endpoint: Change in FRS CV risk prediction with addition of ARI
Specificity: U.88
Sensitivity: 0.84
Results:
1° endpoint: Stenosis >50%

USPSTF Review Lin JS, et al. 2013(30) 24156115	<b>GETABI study</b> Diehm C, et al. 2009(29) <u>19901192</u>	
<u>Study type</u> : Systematic Evidence Review <u>Size</u> : n=1 meta-analysis, 18 population-based cohorts (52,510 pts)	Study type: Prospective cohort study <u>Size</u> : n=6,880 pts; 5,392 pts=no PAD; 836 pts=asx PAD 593 pts=sx PAD	
Inclusion criteria: 3 mo follow-up; designed to evaluate treatment benefit in screen-detected persons or populations who had Asx or unrecognized PAD Exclusion criteria: Pts with DM	<u>Inclusion criteria</u> : Age ≥65 y, 5 y follow-up data, mentally competent to cooperate and sign consent <u>Exclusion criteria</u> : Life expectancy <6 mo	
<ul> <li>Results:</li> <li>ABI added to other risk predictors increases but questions clinical utility or significance.</li> <li>No randomized studies showing improved outcomes in response to detection of Asx disease.</li> <li>Benefit of reclassification including ABI may be higher and clinically important in older populations at higher risk. May be most useful for pts near the thresholds of risk categories.</li> <li>Acknowledge the evidence demonstrating increased morbidity and mortality in Asx pts.</li> </ul>	<ul> <li><u>1° endpoint</u>: Severe vascular events, CV and all-cause mortality.</li> <li><u>Results:</u> <ul> <li>Mortality (pts /1000):</li> <li>No PAD: 19.5;</li> <li>Asx PAD: 41.7;</li> </ul> </li> <li>HR vs. no PAD: 1.66; 95% CI: 1.38–2.0;</li> <li>Sx PAD: 53.0;</li> <li>HR vs. no PAD: 1.89; 95% CI; 1.55–2.30.</li> <li>No significant differences between asx and sx PAD groups in all-cause mortality.</li> <li>Composite outcome All-cause mortality and Vascular events (pts/1000):</li> <li>No PAD: 27.2, Asx PAD: 60.4;</li> <li>HR vs. no PAD: 1.81; CI: 1,53–2.14;</li> <li>Sx PAD 104.7; HR compared to no PAD: 2.66; 95% CI: 2.25–3.15.</li> <li>Difference between PAD groups also significant (HR: 1.48; 95% CI: 1.21–1.80.</li> <li>No differences between PAD groups in MI, stroke, peripheral amputation. Sig differences in myocardial and peripheral revescularualrizations.</li> </ul>	<ul> <li>Men: 4.3%; 95% CI: 0.0–7.6%; p=0.050;</li> <li>Women: 9.6%; 95% CI: 6.1%–16.4%; p&lt;0.001</li> <li>CV mortality:</li> <li>Men: 5.7%; 95% CI: 2.7%–7.9%; p&lt;0.001);</li> <li>Momen: 15.7%; CI: 11.3–20.2%; p&lt;0.001).</li> <li>Restricting use of prediction model to those at intermediate risk resulted in greater effect (15.9% in men and 23.3% in women)</li> </ul>
<ul> <li>Several studies currently ongoing that could give more definitive answers in the future.</li> </ul>	<ul> <li>1 in 5 elderly pts visiting primary care clinician had PAD.</li> <li>Pts with PD regardless of severity had increased risk of CV events and death compared to those without PAD</li> <li>Sx PAD had greater risk of composite outcome of all-cause death or vascular event than asx PAD pts but no greater risk of all-cause mortality alone, MI, or stroke</li> </ul>	

																23512905	2014(32)	Health ABC Study Hiramoto JS. et al.															25721066	Alandab F, et al. 2015/31)
																	Size: n=2,797 pts	study type: Prospective												reviews, 1 meta-analysis	studies, 2 systematic	Size: n=40 individual		<u>Study type</u> : Systematic Review
									<ul> <li>LEX revascularization</li> </ul>	claudication	<ul> <li>Self-reported Hx of</li> </ul>	Exclusion criteria		measurement	Baseline ABI	<ul> <li>No functional limitation</li> </ul>	<ul> <li>No disability</li> </ul>	<ul> <li>Ane 70–79 v</li> </ul>											on asx pts	original data, did not report	Exclusion criteria: Not		screening for asx pts	reporting results of
Other points:	Women: HR: 2.55; 95% CI: 1.13–5.72	• Incident MI:	Women: HR:2.58; 95% CI: 1.35–4.92;	• Stroke:	Women: HR: 5.56; 95% CI: 2.44-12.67.	Men: HR:7.85; 95% CI: 4.44–13.90;	Incident DAD:	Women HR: 4.96: 95% CI: 1.53–16.01.	• CHD Death:	ABI <0.90	Women had higher rates of incident stroke.	category, where women had higher rates of MI and CHD death.	higher rates of CHD death and incident MI except in the 1.3	PAD compared to women across all categories of ABI. Men had	SBP, prevalent CVD, and DM. Men had higher incident clinical	Results: Baseline low ABI associated with black race, elevated		1° endpoint: Development of CV events/mortality, clinical PAD (assessed every 6 mo) Median follow-up 9.37 v	Discussed notential bleeding risk of ASA with no proven benefit	<ul> <li>Little evidence about notential harm or cost-effectiveness</li> </ul>	<ul> <li>Some evidence that screening can lead to improved morbidity</li> </ul>	<ul> <li>ABI screening can improve FRS in risk prediction.</li> </ul>	for mortality outcomes.	<ul> <li>No studies compared screened vs. non screened populations</li> </ul>	(pooled HR=2.99 for all-cause mortality and 2.35 for CV mortality).	PAD is prevalent (average screening yield=17.2%) and mortal	specificity=86%);	15.33; 95% CI: 9.39–25.02; pooled sensitivity=75%;	<ul> <li>ABI is adequate test (diagnostic accuracy=0.87; diagnostic OR:</li> </ul>	Results:		not harmful	screening leads to reduced morbidity and mortality: screening is	1° endpoint: Multiple that would justify screening for asx pts: Accurate test available: disease sufficiently prevalent and mortal:
											outcomes in women	associated with poorer CV	Category of ABI >1.3:	poor outcomes	ABI in women; associated with	<ul> <li>Higher prevalence of borderline</li> </ul>	compared to men	<ul> <li>Subclinical PAD seems to affect women disproportionately</li> </ul>							risk	already indication to treat their	need screening since there is	<ul> <li>High-risk individuals may not</li> </ul>	individuals	from screening low-risk	supports 'pt-important' benefits	No high quality evidence	traditional risk factors	<ul> <li>Yield of ABI screening text in asy nte depende on prevalence of</li> </ul>

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		Exclusion criteria:		
	<ul> <li>normal ABI Results:</li> <li>Asx PAD (26%) vs. no PAD (74%)</li> </ul>	<ul> <li>Asx PAD vs. normal ABI</li> </ul>	Size: n=102 pts	
		• TIA	based cohort	<u>19713540</u>
including stroke, TIA, MI	1° endpoint: Composite vascular events and vascular death median 2.1 y	<ul> <li>Inclusion criteria:</li> <li>Stroke</li> </ul>	Study type: Prospective longitudinal hospital-	Sen S, et al. 2009(36)
5% CI: 1.28–3.47.	r clinical PAD vs. no PAD: 2.11; 9			
	p=0.01.			
35; 95% CI: 1.11–2.44;	HR for subclinical PAD vs. no PAD: 1.6:			
	Adjusted model:	<ul> <li>Hx cancer</li> </ul>		
	ary episode p<0.001. d	<ul> <li>Acute coronary episode within past 7 d</li> </ul>		
; 95% CI: 1.62–4.07;		Exclusion criteria		
: 1.88; 95% CI: 1.27–2.78;		PAD (n=83)		
ed model:		subclinical PAD (n=181), sx	analysis	
		as no PAD (n=446) or	Size: n=710 in final	
D=87.4%; Subclinical	ed; classified . Median 7.2 y survival rates No PAD=87.4%; Subclinical	<ul> <li>ABI measured; classified</li> </ul>		
	Results:	CHD.	range 5.7–8.6 y).	
ห เษยูเธรรเบเา สเเสเรรเร.	with stable	<ul> <li>Indiconsecutive male pise</li> <li>age 45–74 y, with stable</li> </ul>	(median follow-up 7.2 y;	22513182
prognostic effect of PAD status		Inclusion criteria	Study type: Prospective,	Bouisset, F. et al
		PAD		
v.v, p=v.vv+	analysis nr. 12.40, 50/6 Cr. 2.22-7 0.0, p-0.004			
0.4–6.2. Final multivariate		TIA	Size: n=176 pts	
currence of stroke (19.2%)		ASX FAD     Acute ischemic stroke or	וטווטוושו נטווטו ג אנעטא	22138142
	teria: <u>1° endpoint</u> : 30 d recurrence of stroke	Inclusion criteria	Study type: Prospective	TsivgoulisF, et al.
	CVD: 8.2% vs. 30.6%			
<b>7</b> 0	<ul> <li>Nonfatal CV Events: 10.4% vs. 38.0%</li> </ul>	C		
		diagnosed		
<del>.</del>		PAD or previously	Size: n=262 pts	
	iteria: Sx Results:	Exclusion criteria: Sx	y, mean <i>r</i> .r y)	2000017
al, or 15% decrease in ABI)	progression (from normal to abnormal, or	UM	observational study (10	2010(33)
), CVD, CHD, Disease		Inclusion criteria: Type 2	Study type: Follow-up	Bundó M, et al.
R: 9.31; 95% Cl: 4.01– 1: 2.27−10.30	In women with ABI >1.3, Incident MI HR: 9.31; 95% CI: 4.01– 21.63; Incident stroke HR: 4.81; 95% CI: 2.27–10.30			

8	24529125	/l, et al.	Ramos R, et al 2016(38) 26868687 a a 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5		Ratanakorn D, et s al. 2012(37) 21236702	
sample, n=933 pts	Size: Random population	Study type: Cross- sectional	<u>Study Type:</u> Cohort design for matched pair analysis on the basis of study inclusion date and propensity for statin treatment <u>Size</u> : n=5,480 Spanish pts from the Information System for Development of Research in Primary Care database.		<u>Study type</u> : Cross- sectional <u>Size</u> : n=747 Thai pts	
Exclusion criteria: Hx	risk (REGICOR score >5%	Inclusion criteria: Moderate to high vascular	<ul> <li>severe disability subve, E1 intubation and mechanical ventilation; incomplete ABI data.</li> <li>15–85 y</li> <li>ABI measurement documented</li> <li>ABI</li> <li>ABI</li> <li>Ball (0.95;</li> <li>Exclusion Criteria: Previously hx of sx PAD, CHD, stroke or revascularization procedure.</li> </ul>	CT or MRA; age ≥18 y, Exclusion criteria: Hx of previous or current Sx PAD; severe disabling stroke. ET	Inclusion criteria: Consecutive stroke registry pts with ischemic stroke or TIA within 7 d confirmed by	<ul> <li>&lt;18 y</li> <li>Intercerebral hemorrhage</li> <li>Coma</li> <li>Conditions limiting life</li> <li>expectancy to &lt;12 mo</li> <li>Sx PAD</li> </ul>
SCCA by more than 5-fold. ABI diagnosing SCCA: Sensitivity=0.3;	Results: Prevalence of SCCA higher in those with REGICOR score >10% and in pts with asx PAD. Asx PAD increased risk of	1° endpoint: Presence of carotid stenosis	<ul> <li>4.4.3; p=0.001); CVU (UK: 2.15; 95% CI: 1.3/-3.55; p=0.002).</li> <li>Prevalence in pts ≥60 y =25%; ≥70 y =30%. No significant relationship with atherosclerotic risk factors. Strongest prevalence of abnormal ABI in large artery disease and cardioembolic stroke subtypes.</li> <li><u>1ºendpoint:</u> HR of absolute risk reduction in MACE and all-cause mortality and 1-year number needed to treat for 'new' statin users vs. non-statin users followed 2–7 y.</li> <li><u>Results:</u></li> <li>MACE rates</li> <li>New users: 19.7 (95% CI:17.2 to 22.5) Non-users: 24.7 (95% CI: 21.8 to 27.8) (20% RRR)</li> <li>1 y NNT: 200</li> <li>All-cause mortality rates</li> <li>New users: 24.8 (95% CI: 22.0 to 27.8) (19% RRR)</li> <li>1 y NNT 239</li> <li>NNT decreased with ABI cutpoint</li> </ul>	<ul> <li>Prevalence of abnormal ABI=18/1%; Multivariate analysis abnormal ABI related to female sex (OR: 1.61; 95% CI: 1.09–2.40; p=0.017); Age ≥60 y (OR: 3.54; 95% CI: 2.14–5.85; p&lt;0.001); Previous ischemic events including CAD (OR: 2.55; 95% CI: 1.47– A 2: n=0.001); CVD (OB: 3.15: 05% CI: 1.37, 3.55: n=0.002)</li> </ul>	<u>1° endpoint</u> : Prevalence of PAD among total population and subgroups Results:	• Cumulative event-tree survival: 1.5; 95% CI: 1.2–1.9 y vs. 2.5 y; 95% CI: 2.4–2.6 y; p=0.0001
Helps target screening, 20	subclinical carotid or intracranial atherosclerosis	<ul> <li>ABI emerged as tool to identify pts with high risk of having</li> </ul>	<ul> <li>First study to report the association between statins and both MACE and mortality reduction among individuals free of clinical CVD, but with asx PAD identified by ABI.</li> <li>Reduction observed regardless of CVD risk scores at baseline</li> <li>Absolute reduction in MACE and all-cause mortality similar to that seen in secondary prevention studies.</li> </ul>		<ul> <li>Early detection of PAD may facilitate treatment and identify excess risk of subsequent stroke or other CV events.</li> </ul>	

WALCS Study McDermott MM etStudy type: Prospect cohort study of PAD p with differing types of symptoms (same coh 1528034315280343Size: • n=417 pts with PAD • n=259 pts without P	WALCS Study McDermott MM, etStudy type: Cross- sectional, new pts consecutively identific and pts already identi ith PAD from large general medicine practice.Size: • n=430 men and women with PAD ASX active=63 ASX inactive=28	McDermott MM, etStudy type: Cross- sectional2000(40)Size: • Stratified random sampling of 32,538 • Final sample n=574 asx pts
Study type: Prospective cohort study of PAD pts with differing types of leg symptoms (same cohort as above) 2 yr follow-up <u>Size:</u> • n=417 pts with PAD • n=259 pts without PAD	Study type: Cross- sectional, new pts consecutively identified and pts already identified with PAD from large general medicine practice. <b>Size:</b> • n=430 men and women with PAD • n=130 without PAD. ASX active=63 ASX inactive=28	<u>:</u> : Cross- random f 32,538 ıple n=574
Inclusion criteria • ABI <0.90 • ≥55 y • Non-PAD group identified from internal medicine practice <b>Exclusion criteria:</b> • ABI >1.5 • Normal ABI • Dementia • Amputation • Non-English speaking • Wheelchair bound	Inclusion criteria Diagnosed with PAD (ABI<0.90); ≥55 y Exclusion criteria: • ABI > 1.5; • Normal ABI, • Dementia • Amputation • Non-English speaking • Wheelchair bound • Nursing home resident • Recent surgery	stroke, PAD, CAD Inclusion criteria: Community dwelling disabled women ≥65 y participating in Women's Health and Aging Study Exclusion criteria: Mini-mental score <18
<u>1° endpoint</u> : Decline in 6 MWT, Usual pace and fastest-pace 4- Meter velocity, summary performance score <u>Results</u> : Baseline physical functioning poorer in asx PAD than non-PAD; decline greater on all measures. asx PAD has greater decline in 6 MWT than pts with claudication	<ul> <li><u>1° endpoint</u>: 6 MWT scores, 7 d physical activity, SPPB, Questionnaires</li> <li><u>Results</u>: <ul> <li>PAD sj. Divided into 6 categories. asx 2 categories: active vs. inactive</li> <li>33.3% active and 53.6% inactive PAD pts reported sx during 6MWT</li> <li>All PAD groups had worse functioning that non-PAD group</li> <li>Asx inactive functioning similar to claudication group</li> <li>Asx inactive functioning poorer than claudication group</li> </ul> </li> </ul>	<ul> <li>95% CI: 0.18–0.42; specificity=0.95 (95% CI: 0.93-0.96); PPV=0.26 (95% CI: 0.15–0.37), NPV= 0.95 (95% CI: 0.94–0.97).</li> <li>1° endpoint: Prevalence of Asx PAD; relationship between physical functioning and Asx PAD.</li> <li>Results: • ABI&lt;0.90=198 (34.5%)</li> <li>• ABI&lt;0.50=48 (8.4%)</li> <li>• Subjective and objective measures of mobility and lower extremity function, all statistically lower in Asx PAD compared to non-PAD.</li> </ul>
<ul> <li>Asx pts have &gt;2 y decline in physical functioning compared to asx non-PAD pts. 6 MWT decline greater in asx pts that IC group.</li> </ul>	NA	<ul> <li>Asx PAD is independently associated with impaired lower extremity functioning.</li> </ul>

	<u>17039537</u>	Niazi K, et al. 2006(45)		24222666	al. 2013(44)	LIFE study McDermott MM, et		2010(43) 20550604	WALCS study McDermott MM, et al.		2008(+2) 16389250	WALCS Study McDermott MM, et al.	
	<u>Size</u> : n=107 pts, 208 limbs	Study type: Cross- sectional study	categories of: Definite PAD , borderline PAD, low normal ABI, no PAD	Size: n=1,566 pts categorized into	community-dwelling sedentary older adults	Study type: Cross- sectional study in		<u>Size</u> : n=415 pts followed up to 7 y	<u>Study type</u> : Prospective observational study		Size: n=417 men and women with PAD	Study type: Prospective cohort study with median follow-up of 36 mo	
	prior to DSA <u>Exclusion criteria:</u> • Pts with noncompressible	<ul> <li>ABI performed within 30 d</li> </ul>	Exclusion criteria: N/A	<ul><li>Physical activity/wk</li><li>Functional limitations</li></ul>	<ul> <li>Community-dwelling</li> <li>Sedentary (&lt;125 min of</li> </ul>	<ul> <li>Age 70–89 y</li> </ul>		Exclusion criteria: See above	Inclusion criteria: See above	<ul> <li>1.5; Normal ABI,</li> <li>dementia, amputation,</li> <li>nonEnglish speaking,</li> <li>wheelchair bound, nursing</li> <li>home resident</li> </ul>	Non-PAD group identified from internal medicine practice     Exclusion criteria: ABI	Inclusion criteria: • Age ≥55 y • ABI <0.90	<ul> <li>Nursing home resident</li> <li>Recent surgery</li> </ul>
	<ul> <li><u>Results:</u></li> <li>Sensitivity of the HAP and LAP ABI for diagnosis of PAD was 69% and 84%, respectively</li> <li>Overall accuracy of HAP and LAP ABI was 72% and 80%,</li> </ul>	<u>1° endpoint:</u> N/A		<ul> <li>In asx pts lower ABI values associated with longer 4 meter walk time and slower walking velocity</li> </ul>	<ul> <li>Results:</li> <li>65% of definite PAD pts asx.</li> </ul>	1° endpoint: Physical function measures	claudication (HR: 2.94; 95% CI: 1.39–6.19; p=0.005). Asx pts did not demonstrate as much decline in 6MWT as pts with claudication.	Results: Always asx pts had greater mobility loss than pts with	<u>1° endpoint</u> : 6 MWT, becoming unable to walk up and down a flight of stairs or walk ¼ mile without assistance in pts without mobility loss at baseline	less frequently	<ul> <li>Results:</li> <li>Pts separated into groups based on physical activity level (walk 3 or more times per wk vs. less frequently).</li> <li>Asx PAD pts who walked for exercise 3 or more times per wk had less functional decline than those who walked for exercise</li> </ul>	<u>1° endpoint:</u> Rate of decline in 6 MWT, Usual pace and fastest- pace 4-Meter velocity, summary performance score	
22	comparison to the HAP ABI in diagnosing PAD	<ul> <li>LAP ABI has better sensitifvity and overall accuracy in</li> </ul>	functional impairment.	<ul> <li>Even in individuals who are considered functionally impaired, low ABI is associated with greater</li> </ul>	cause of functional limitations in older persons.	Lower extremity atherosclerosis may be common preventable			N/A		prs.	<ul> <li>Greater physical activity associated with less decline in physical functioning in ASX PAD</li> </ul>	

23	value is in pts with symptoms and normal		limbs) and 25 normal pts	
	Results: 97% and 96% stress testing	Exclusion criteria: N/A	Size: n=218 pts (372	1166101
NA	<u>1° endpoint</u> : Sensitivity and specificity of exercise ABI to detect PAD	Inclusion criteria: Able to have ABI, treadmill ABI and reactive hyperemia	Study type: Observational	Ouriel K, et al. 1982(48)
			Size: n=89 limbs	
	than in waveform interpretation		readers tor each technique)	
	(presence and level of PAD), less variability in interpretation using pressure	incomplete reports	/DWand angiography 2009–2011 (blinded	
	Results: 66% diagnostic accuracy	Exclusion criteria: Those with	pts with SDP/PVR	2014(47) 24200144
<ul> <li>Readings reflecting incompressibility were not utilized</li> </ul>	<u>1° endpoint</u> : Determination of the most accurate diagnostic value	Inclusion criteria: Having both SDP/PVR and angiography	Study type: Single healthcare system,	Eslahpazir BA, et al.
			Size: n=114 pts undergoing SDP/PVR and angiography	
	correct classification using either SDP or PVR	Exclusion criteria: No angiography	angiography for Dx of PAD	
	Results: 97% of normal limbs were correctly classified by SDP/PVR. 86%	angiography	SDP/PVR compared to the gold standard of	1997(46) 9308598
NA	1° endpoint: Correct classification of PAD	Inclusion criteria: 11 normal volunteers and 103 pts having had	<u>Study type:</u> Observational study of	Rutherford RB, et al.
	& 95% CI)			Year Published
Summary/Conclusion Comment(s)	Primary Endpoint and Results (include P value; OR or RR;	Patient Population	Study Type/Design; Study Size	Study Acronym; Author;
cal Testing-Section 3.2.		Evidence Table 5. Nonrandomized Trials, Observational Studies, and/or Registries of Physiologi	5. Nonrandomized Tria	Evidence Table
onnaire.	stenosis >50%; SF, Short Form; Sx, symptomatic; TIA, transient ischemic attack; US, United States; and WIQ, Walking Impairment Questionnaire.	TIA, transient ischemic attack; US, United	Short Form; Sx, symptomatic;	stenosis >50%; SF, {
<ol> <li>number needed to treat OR, odds ratio; PAD, peripheral relative risk; SBP, systolic blood pressure; SCCA, significant</li> </ol>	eclassification improvement; NNT, number nee predictive value; pt, patient; RR, relative risk; S	National Health and Nutrition Examination Survey; NPV, negative predictive value; NRI, net reclassification improvement; NN artery disease; PCS, physical composite score, PFWT, pain free walking time; PPV, positive predictive value; pt, patient; RR,	, physical composite score, PF	National Health and I artery disease; PCS,
FRS, Framingham risk score; HAP, high ankle pressure; HR, hazard ratio; HTN, hypertension; IC, intermittent claudication; LAP, low ankle pressure; MACE, major adverse cardiovascular event; LEX, lower extremity; MCS, mental health composite score; MI, myocardial infarction; MRI, magnetic resonance imaging; MWT, mean walking time; N/A, not applicable; NHANES,	FRS, Framingham risk score; HAP, high ankle pressure; HR, hazard ratio; HTN, hypertension; IC, intermittent claudication; LAP, low ankle pressure; MACE, major adverse cardiovascula event; LEX, lower extremity; MCS, mental health composite score; MI, myocardial infarction; MRI, magnetic resonance imaging; MWT, mean walking time; N/A, not applicable; NHANES	ssure; HR, hazard ratio; HTN, hypertension omposite score; MI, myocardial infarction; I	sk score; HAP, high ankle pres (tremity; MCS, mental health c	FRS, Framingham ris event; LEX, lower ex
CHD, coronary heart disease; CI indicates confidence interval; us; DSA, digital subtraction angiography; ET, endotracheal;	ABI indicates ankle-brachial index; ASA, acetylsalicylic acid; asx, asymptomatic; BL, baseline; CAD, coronary artery disease; CHD, coronary heart disease; CI indicates confidence inter CLI, critical limb ischemia; CT, computed tomography; CVD, cardiovascular disease; CV, cardiovascular; DM, diabetes mellitus; DSA, digital subtraction angiography; ET, endotracheal;	ABI indicates ankle-brachial index; ASA, acetylsalicylic acid; asx, asymptomatic; BL, baseline; CAD, coronary artery disease; CL, critical limb ischemia; CT, computed tomography; CVD, cardiovascular disease; CV, cardiovascular; DM, diabetes mellit	brachial index; ASA, acetylsalii nemia; CT, computed tomograp	ABI indicates ankle-t CLI, critical limb isch

vessels • ABI >1.40

respectively

with variable inspired oxygen	measures	nonsmoking adults	study of repeated measurements of TcPO ₂	Hendricker C 1987 (53)
<ul> <li>Mornings and afternoons over 7 d to 7 mo</li> </ul>	1° endpoint: Variability of repeat	Inclusion criteria: Healthy	Study type: Prospective	Wagener JS and
	occlusive disease.			
	least 1 of the 2 noninvasive vascular			
	Results: In 84.2% of cases, diabetic limbs with ABI ≥1.40 had abnormal results in at			
			Size: n=510 pts	
	presence by TBI	Exclusion criteria: N/A		18692981
S0% with DM     Anniographic correlations	1° endpoint: Association of risk factors with ABI >1.4 and ABI <0.9 and disease	Inclusion criteria: ambulatory pts presenting to vascular lab	sectional	Aboyans V, et al. 2008(22)
	Kesuits: High sensitivity and specificity	•	vascular lab	
		Exclusion criteria: N/A	Size: n=69 pts of the total	<u>18313338</u>
	of disease			2008(52)
abnormal TBI	>1.3, TBI <0.6 and angiographic evidence		ABI >1.3 and angiography	al
<ul> <li>I arger population with normal ABI and</li> </ul>	1° endnoint: Presence of abnormal ABI	Inclusion criteria: TBL ABI and	Study type: Retrospective	Suominen V. et
	Results: 92% of pts with TBI <0.7 had angiographic evidence of PAD	Exclusion criteria: N/A	Size: n=116 limbs	
	PAD with TBI <0.7	angiography and TBI	study	2013(51) 22899598
<ul> <li>67% DM and 19% on hemodialysis</li> </ul>	1° endpoint: Angiographic evidence of	Inclusion criteria: Pts with ABI >1.4,	Study type: Retrospective	Weinberg I, et al.
			Size: n=30 limbs	
	Results: 13 of 30 limbs with abnormal	Exclusion criteria: N/A	ulcers)	
standards)		י. <i>י</i> , חמצפווי פמויפויפי	ABI, TBI (many with	922783531
Studies with normal population and TBI had	<u>1° endpoint:</u> ABI or TBI correlation with	Inclusion criteria: TBI <0.6 or ABI <	Study type: Retrospective	Park SC, et al.
	angiographic severity was worse.			
	ABI could not be determined in 34%. In those with calcifications correlation with	bypass		
	angiographic disease was weak (<0.48).	Exclusion criteria: Distal arterial		
not userui	Results: Correlation between ABI and		extremities	
Biphasic Doppler signals useful, monophasic	clinic	with nonhealing foot ulcer and/ or	<b>Cito:</b> n=107 lowor	<u>21514102</u>
• Anterial calcilication evaluated using prain A- ray	angiography in pts seen in diabetic foot	clinic with angiography and ABI. All	study	2011(49)
		In the indication of the second	Study time. Dependence	
	ABI		and 10 stable claudicants	

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。 Results: The TBI, arm minus toe
1° endpoint: Relationship of toe pressure to healing
controlling is done for the value of the toe pressure and ABI or ankle pressure
significantly related to the occurrence of rest nain skin breakdown or both after
Results: I nw the PW amplitude is
1° endpoint: Clinical correlation
correlation with angiography
<b>Results:</b> Well tolerated and ex
with systolic pressure assessment
ž
wound healing and limb salvage rates compared with indirect revascularization, with no effect on mortality or reintervention rates.
tibial vessels appears to result in improved
<u>1° endpoint:</u> Wound healing salvage, mortality
Best prediction SPP +
Results: Healing more likely a
<u>1° endpoint</u> : Ability of test to p wound healing
Results: Correlation 0.87 (p<0
1° endpoint: Correlation of TB
pulse oximetry

	Results: The pts walked for 1 or 2 min at 4 km/h and 1 or 2 min at 6 km/h, and the fall in pressure was the same when measured immediately after exercise.		<u>Size</u> : n=26 pts	1980 (64) <u>7357254</u>
N/A	1° endpoint: Comparison of 2 protocols	Inclusion criteria: Presentation with claudication	Study type: Observational	Laing SP and Greenhalgh RM
the reliability of the C-protocol was dependent on grade of the treadmill (0%, 10%, and 12%) with a mean ICC of 0.76 (95% CI: 0.54-0.88), 0.89 (95% CI: 0.86-0.91), and 0.91 (95% CI 0.88-0.92), respectively	reliabilities of the C- and G-protocol (as assessed by the ICC) were 0.85 (95% confidence interval [CI]: 0.82-0.88) and 0.83 (95% CI: 0.80-0.85), respectively, without dependency of the reliability on velocity or grade.	reprimed, inclusion ciliteria were me use of a C- or G-protocol, repetition of this protocol, and a retrievable ICC.	<u>Size</u> : n=8 studies, 658 pts	
For ACD, the reliability was significantly better for the G-protocol (0.95, 95% CI: 0.94-0.96)	<u>1° endpoint:</u> Reliability of treadmill testing	Inclusion criteria: Trials assessing reliability offreadmill testing were	Study type: Meta regression analysis	Nicolaï SP, et al. 1990 (63)
	Results: Only small overlap between the 2 populations of PAD identified	Exclusion criteria: Inability to exercise	Size: n=12,312 consecutive pts	
<ul> <li>To determine whether postexercise criteria for PAD diagnosis recommended by the AHA identifies the same group of PAD ats</li> </ul>	<u>1° endpoint:</u> Diagnosis of PAD using the 2 criteria	Inclusion criteria: Consecutive pts underwent exercise ABI	Study type: Retrospective analysis of clinical results	Mahe G, et al. 2015 (62) 26252297
ratios: normal, claudication, limb salvage, claudication/incompressible arteries, and limb salvage/incompressible arteries.	Results: Toe pressure was the most reliable indicator of occlusive disease, and was able to assess disease distal to the ankle	<ul> <li>Both asx volunteers and pts with PAD presenting to the vascular lab were studied</li> </ul>	<u>Size</u> : n=219 limbs	<u>6833348</u>
<ul> <li>5 groups were separated using the ankle- brachial and the toe-ankle systolic pressure</li> </ul>	<u>1° endpoint</u> : Diagnostic accuracy toe pressure and ABI	<ul> <li>Presence of limb</li> </ul>	Study type: Observational	Vincent DG, et al. 1983 (61)
	tissue lesions may improve wound healing and limb salvage rates compared with indirect revascularization	Exclusion criteria: N/A	RCT)	
revascularization to achieve healing of ischemic tissue lesions of the foot and limb salvage is controversial.	Results: Direct revascularization of the foot and isome affected by ischemic	direct revascularization according to the angiosome principle and 575 legs treated by indirect revascularization	analysis Size: n=9 studies (no	Juvonen T 2014 (60) 24491282
<ul> <li>Aim: The efficacy of anglosome-targeted</li> </ul>	1° endpoint: Wound healing	Inclusion criteria: 715 legs treated by	Study type: Meta-	Biancari F and
	pressure, and the absolute toe pressure had an average sensitivity and specificity of 85% and 88% for asx limbs and 89% and 86% for ischemic limbs.		presence or absence of diabetes in 294 limbs <b>Size:</b> n=294 limbs	

Stein R, et al. <u>Study t</u> 2006(70) <u>16669410</u> <u>Size</u> : n:	Bunte MC, et al. <u>Study type</u> : 2015(69) <u>26892836</u> <u>Size</u> : n=89 co pts	Biotteau E, et al. <u>Study t</u> 2009(68) matche <u>20087286</u> <u>Size</u> : n:	Castronuovo JJ, <u>Study type</u> : P et al. double blind st 1997(67) <u>9357464</u> <u>Size</u> : n=53 pts	Sumner DS and Strandness DE 1969 (66) <u>5777227</u>	Raines JK, et al. <u>Study t</u> 1976 (65) <u>1246689</u> <u>Size</u> : n:
<u>Study type</u> : Retrospective review <u>Size</u> : n=396 pts	<u>Study type:</u> Observational <u>Size</u> : n=89 consecutive pts	<u>Study type</u> : Retrospective matched paired study <u>Size</u> : n=120 pts	<u>Study type</u> : Prospective double blind study <u>Size</u> : n=53 pts	Study type: Observation	Study type: Observation Size: n=4,500 procedures
Inclusion criteria: Sx outpatients referred for measurement of segmental blood pressure, the ABI or pulse volume recordings by physicians not specialized in the evaluation and management of pts with PVD	Inclusion criteria: CLI and presentation with rest pain	Inclusion criteria: Pts presenting to the vascular lab with suspected CLI	Inclusion criteria: Vascular lab referrals for CLI Exclusion criteria: Sepsis or need for guillotine amputation	Inclusion criteria: Pts presenting to the vascular lab with claudication	Inclusion criteria: Pts in the vascular lab
<u>1° endpoint</u> : Diagnostic utility of measuring the ABI at rest in pts referred to the vascular laboratory for evaluation of suspected PAD <u>Results</u> : Nearly half of pts referred to the outpatient vascular laboratory because of	<b><u>Results</u>:</b> Among 31 CLI pts with available ABI and TBI results, 19 (61%) had a TBI <0.7 <i>and</i> a non-compressible or resting ABI <0.9. Conversely, no pts with a borderline or normal ABI (0.9–1.4) had a normal TBI (≥0.7)	<u>1° endpoint</u> : Whether a difference can be found for chest and foot TcPo ₂ respectively between pts with and without DM referred for clinically suspected CLI. <u>Results</u> : TcPo ₂ is lower at the chest but not at the foot level in diabetic than in non-diabetic pts with suspected CLI.	<u>1° endpoint</u> : Prediction of wound healing by SPP <u>Results</u> : SPP measurements identified 31 of 32 limbs diagnosed as having CLI by clinical evaluation (i.e., group I, those limbs that required vascular reconstruction or major amputation)	1° endpoint: Relationship between calf blood flow and ankle blood pressure in pts with claudication Results: Close correlation	<u>1° endpoint</u> : Criteria for management <u>Results</u> : Excellent reproducibility for physiologic testing including pulse volume recording and segmental pressures
<ul> <li>Diagnostic accuracy was improved with pulse volume recordings and exercise ABI</li> </ul>	<ul> <li>Among a contemporary, real-world CLI population, 29% had near-normal or normal ABI, despite having significant infragenicular arterial disease.</li> </ul>	<ul> <li>Evenly matched DM and non-DM</li> <li>30 mm Hg threshold applicable to both populations</li> </ul>	<ul> <li>DM and wound size similar in 2 groups</li> <li>The sensitivity of SPP &lt;30 mm Hg as a diagnostic test of CLI was 85%, and the specificity was 73%. The overall diagnostic accuracy of SPP less than 30 mm Hg as a diagnostic test of CLI was 79.3% (p&lt;0.002, Fischer's exact test).</li> </ul>	NA	N/A

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PIVUS study Wilkström J, et al. 2008(72)Study type: Observational test comparisonInclusion criteria: • General population register Sweden • Age 70 y12 endpoint: • Age 70 y12 endpoint: • Age 70 yWilkström J, et al. 2009(73) 19446989Size: n=306 pts Size: n=306 ptsSize: n=306 pts • MBMRA • Bight: 20 (10 • Eft: 15 (7, 2 • Specificity: • 99 (96, 100) • P99 (96, 100) • P99 (96, 100) • PPV: • P10 (P10) • P10 (P10)P10 (P10) 	Study Acronym;       Study Type/Design;       Patient Population         Author;       Study Size         Year Published	Shishehbor MH, 2016(7)         Study type: (Dservational 26860642         Inclusion criteria: (Dservational 2016(7)         Inclusion criteria: (Dservational 26860642         Inclusion criteria: (Dservational 2016(7)         1° endpoint: (Dservational 2016(7)         1° endpoint: (Dservational 2016(7)         Occurrent recommended hemodynamic (PS) and TBI to diagnose lower extremity (Lers and severe disease)         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and ucers and severe disease         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and proven severe disease.         • Current recommended hemodynamic pressures to diagnose CLI are insensitive and pressures to diagnose CLI are insensitive and pressures to diagnose CLI are insensitive and proven severe pressure is more according to cutoffs suggested for CLI.         • Current recommended hemodynamic pressure is more pressure is more discording to cutoffs suggested for CLI.         • Current recording s: pressure is more sensitive	susj
Presence of stenosis in pelvic or leg nt or left legs 0, 34) 27)	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Anatomic A	suspected arterial disease nad a normal resting ABI
<ul> <li>Low sensitivity but good PPV.</li> <li>High specificity. Similar results (not shown) to detect occlusion, except lower PPV</li> </ul>	Summary/Conclusion Comment(s)	urement of extremity       • Current recommended hemodynamic pressures to diagnose CLI are insensitive and lower extremity ulcers and angiographically proven severe disease. Toe pressure is more sensitive in pts with CLI.         a few pts for CLI I for ABI       • Sensitive in pts with cleases. Toe pressure is more sensitive in pts with CLI.         betes mellitus; ICC, intraclass correlation coefficient; ICD, PVR, pulse volume recordings; PW, pulse wave; RCT, ymptomatic; TBI, toe-brachial index; TBP, toe blood pressure;         Anatomic Assessment (Ultrasound, CTA, MRA,	

	Shareghi S, et al. 2010(76) <u>19753637</u>	Burbelko M, et al. 2013(75) <u>23188773</u>	Clairotte R, et al. 2009(74) <u>19366974</u>	Guo X, et al. 2008(20) <u>18362433</u>
	<u>Study type:</u> Observational <u>Size</u> : n=28 pts	<u>Study type:</u> Observational <u>Size:</u> n=152 pts	<u>Study type:</u> Observational test comparison <u>Size:</u> n=63 pts	<u>Study type:</u> Observational test comparison <u>Size</u> : n=298 pts
	Inclusion criteria: consecutive pts with sx lower extremity IC and an abnormal ABI (ABI<0.9) Exclusion criteria: N/A	ABI method: Doppler Inclusion criteria: Underwent MRA and DSA of the lower extremities within 30 d. Exclusion criteria: N/A	Inclusion criteria: Referrals to clinic for duplex Exclusion criteria: DM Gold standard: • Duplex ultrasound • Velocity ratio ≥2 for stenotic:proximal segments	Inclusion criteria:         • Age ≥35 y         • Cardiology clinic: referrals for DSA & ABI         Exclusion criteria: Severe DM & hypertension         Gold standard:         • DSA.         • Stenosis ≥50%         ABI method:
-	<u>1° endpoint</u> : N/A <u>Results:</u> Sensitivity: 99 Specificity: 98	1° endpoint: Evaluation of stenosis grade and image quality Results: Sensitivity: 73–93 Specificity: 64–89	<u>1° endpoint</u> : Presence of stenosis in iliac to ankle arteries <u>Results:</u> Sensitivity: 73 (N/A) Specificity: 98 (N/A) PPV: 98 (N/A) NPV: 78 (N/A)	<ul> <li>80 (74, 84)</li> <li><u>endpoint</u>: Presence of stenosis below aorto-iliac bifurcation in leg with lower ABI</li> <li><u>Results</u>: Sensitivity: 76 (N/A)</li> <li>Specificity: 90 (N/A)</li> <li>PPV: 36 (N/A)</li> <li>NPV: 98 (N/A)</li> </ul>
20	<ul> <li>MDCT demonstrated accurate detection of hemodynamically significate disease of the lower extremities</li> </ul>	<ul> <li>CE-MRA demonstrates good sensitivity and specificity</li> <li>CE-MRA is standardizable and shows good inter-observer agreement</li> <li>Use of CE-MRA as alternative to intra-arterial DSA is well justified</li> </ul>	<ul> <li>Moderate sensitivity &amp; very good specificity. No indication of % pts with PAD symptoms but only 14% had "clinical PAD".</li> <li>Duplex ultrasound not ideal gold standard.</li> <li>Small study.</li> </ul>	<ul> <li>Moderate sensitivity and good specificity. No indication of % with PAD symptoms but low prevalence of PAD on DSA (7%) suggests it was negligible.</li> <li>However 53% had coronary heart disease and 13% stroke.</li> </ul>

	He C, et al. 2014(79) 25252783	Ota H, et al. 2004(78) <u>14684540</u>		De Vries SO, et al. 1996(77) 8706687
טין יסין י <u>אזוט</u> טיק יסין י	Study type: NR (retrospective cohort study)	Study type: Observational Size: n=27 cases in 24 pts		<u>Study type:</u> Meta- analysis
of the arteries in both legs. Exclusion criteria: Allergy to the iodine	Inclusion criteria: Consecutive pts with DM (13 women; mean age, 69.42±11.04 y) and 101 pts without DM (23 women; mean age, 68.50±13.59 y) (23 women; mean age, 68.50±13.59 y)	Inclusion criteria: • Sx lower extremity peripheral arterial occlusive disease • Underwent both MDCT angiography and digital subtraction angiography of the aortoiliac and lower extremity arteries Exclusion criteria: N/A	<ul> <li>Additional references from bibliographies of review articles and original papers.</li> <li>Studies pertaining to diagnostic performance of duplex or color-guided duplex ultrasonography in PAD of the lower extremities</li> <li>Contrast angiography was used as the gold standard Significant lesion defined as an arterial diameter reduction on angiography of 50%–100%</li> <li>The absolute numbers of True- positive, false-negative, true-negative, and false-positive observations were available or derivable.</li> <li>Exclusion criteria: N/A</li> </ul>	Inclusion criteria: • Medline, English-language studies published between January 1984 and hund 1004
included in the analysis. Plaque and stenosis were detected in 681 segments in 60 pts with DM (63.1%) and 854 segments in 101 pts without DM (46.9%;	1° endpoint: Plaque type, distribution, shape and obstructive natures were compared between pts with and without DM	<u>1° endpoint</u> : N/A <u>Results:</u> 99.2 Specificity: • 99.1	<ul> <li>Social Structure</li> <li>Sa (Duplex)</li> <li>Specificity:</li> <li>95</li> <li>95</li> </ul>	<u>1° endpoint</u> : N/A <u>Results:</u> Constitutor
<ul> <li>The advanced and noninvasive</li> <li>MDCT could be used for routine preoperative evaluations of LEA.</li> </ul>	• DM is associated with a higher incidence of plaque, increased incidence of mixed plaques, moderate stenosis and localization primarily in the distribution for promotion	<ul> <li>MDCT angiography is a reliable method for evaluation the aortoiliac and lower extremity arteries</li> </ul>		N/A

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• 40-row MDC IA may be used as a screening tool in pts with mild lower extremity PAOD as it is a noninvasive and more accurate modality when compared to DUS.	<b><u>Results:</u></b> MDCTA detected obstructed or stenotic lesions in 16.8% of arteries, vs. 11.1% compared to DUS. When suprapopliteal arteries alone were considered, MDCTA detected lesions in 15.0% of arteries vs. 11.0% with DUS. When infrapopliteal arteries only were considered, MDCTA detected lesions in 19.6% of arteries, vs. 11.3% with DUS. MDCTA showed 5.7% (95% CI: 3.5%–7.9%) more lesions than DUS when all arteries were considered to together. 8.3% (95% CI: 4.6%–12.0%) more lesions	Exclusion criteria: Pis with to and reg pain, diagnosed as mild PAOD, Exclusion criteria: N/A	Size: n=43 pts	2012(81) 21345629
	<b>Results:</b> In a pt-based analysis, the sensitivity of MDCT for detecting significant proximal IPA disease was 100% and, specificity 74%, positive predictive valve was 66%, and negative predictive value was 100%. In assessing the distal IPA and cavernosal arteries, the sensitivity was 100%, specificity was 64%, positive predictive value 89%, and negative predictive value of 100%. MDCT used significantly more contrast and more radiation than aortography.			
<ul> <li>Studies were read independently and blinded</li> </ul>	<u>1° endpoint</u> : Localize the IPA origin, degree of stenosis (normal: <50% stenosis or abnormal: >50% stenosis or occlusion), normal= and extent of calcification, quantified using a nominal scale (0=no calcification, 1 ≤25%, 2=25%–50%, 3 ≥50% of the IPA length).	Inclusion criteria: MDCT and aortography of the pelvic vascularulature prior to consideration for transcatheter aortic valve replacement Exclusion criteria: N/A	<u>Study type</u> : NR (retrospective cohort study) <u>Size</u> : n=83 pts	Philip F, et al. 2013(80) <u>23553996</u>
	<ul> <li>p&lt;0.05). Regarding these plaques, pts with DM had a higher incidence of mixed plaques (34.2% vs. 27.1% for pts without DM). An increased moderate stenosis rate and decreased occlusion rate were observed in pts with DM relative to pts without DM (35.8% vs. 28.3%; and 6.6% vs. 11.4%; respectively). In pts with DM, 362 (53.2%) plaques were detected in the distal lower leg segments, whereas in pts without DM, 551 (64.5%) plaques were found in the proximal upper leg segments. The type IV plaque shape, in which the full lumen was involved, was detected more frequently in pts with DM than in pts without DM (13.1% vs. 8.2%).</li> </ul>	contrast agent, liver, kidney or HF (Creatinine level ≥120 mol/L), pregnancy and leg amputation. The vascular exclusion criteria included vascular malformations, poor imaging and a lumen diameter <1.5 mm.		

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		Romano M, et al. 2004(84) <u>15145492</u>		Mesurolle B, et al. 2004(83) <u>15246474</u>	Joshi SB, et al. 2009(82) 20083076
	<u>Size</u> : n=42 pts	Study type: NR (prospective)	<u>aize</u> : n- lo pts	Study type: NR (prospective)	<u>Study type</u> : NR (retrospective) <u>Size</u> : n=37 pts
	Exclusion criteria: Pts with previous radiological interventions or surgery for their peripheral vascular occlusive disease	Inclusion criteria: Untreated pts with peripheral vascular occlusive disease	Exclusion criteria: N/A	Inclusion criteria: In the assessment of occlusive arterial disease of abdominal aorta and the lower extremities.	Inclusion criteria: Consecutive pts requiring evaluation of aortoiliofemoral anatomy prior to cardiovascular procedures (pts being considered for percutaneous aortic valve intervention.) Exclusion criteria: N/A
	Results: Overall sensitivity and specificity of MDCTA were 93 and 95%, respectively, with positive and negative predictive values of 90 and 97%. Overall diagnostic accuracy was 94%. Normal arterial	<u>1° endpoint</u> : Sensitivity and specificity of 4 channel MDCTA of the abdominal aorta and lower extremities arteries compared with DSA.	<b>Kesurs:</b> Overall sensitivity of nelical C1 was 91% and specificity 93%. Segmental analysis found a sensitivity of 43% in infrapopliteal arteries, and a specificity of 86%. elical CT was inconclusive in 6.2% of segments whereas angiography was inconclusive in 5%. Overall sensitivity of helical CT was 91% and specificity 93%. Segmental analysis found a sensitivity of 43% in infrapopliteal arteries, and a specificity of 86%.	<u>1° endpoint</u> : Sensitivity and specificity vs. catheter angiography	<ul> <li>when only the infrapopliteal arteries were compared, and 4.0% (95% CI: 1.3%–6.8%) more lesions when only suprapopliteal arteries were compared (p&lt;0.01 for all comparisons).</li> <li>1º endpoint: Conventional angiographic and CT images were analyzed independently to assess suitability for large bore (7 mm diameter) intra-arterial catheter access.</li> <li>Results: Excellent CT image quality was achieved in 34 of 37 pts (92%). The mean contrast dose for CT was 12±2 mL. In 9 pts (24%), CT changed the assessment of femoral access feasibility. Furthermore, in another 7 pts (19%), unfavorable anatomy as shown by CT directed the avoidance of a particular side. Overall, CT findings altered the interventional approach in 16 pts (43%).</li> </ul>
32		NA		<ul> <li>16 pts underwent both transcatheter angiography and helical CT</li> </ul>	<ul> <li>Purpose was to evaluate the feasibility of using ultra-low-dose intra-arterial contrast injection for iliofemoral CT angiography to follow diagnostic cardiac catheterization.</li> <li>0 to 15 mL of contrast diluted with normal saline was injected intra-arterially via the pigtail catheter while a spiral CT of the abdomen and pelvis was acquired</li> <li>There was no significant detected in renal function after coronary and CT angiography (estimated glomerular filtration rate 54.8±3.8 mL/min before 53.3±3.9 mL/min after, p=0.55).</li> </ul>

Andreucci M, et al. 2014(86) <u>24895606</u>	Martin ML, et al. 2003(85) <u>12646460</u>
Study type: A review of the evidence base for the adverse effects associated with radiographic contrast drugs. Size: N/A	<u>Study type</u> : NR (prospective) <u>Size</u> : n=41 pts
Inclusion criteria: N/A Exclusion criteria: N/A	Inclusion criteria: Pts referred for DSA of the lower extremities for investigation of sx atherosclerotic disease of the legs Exclusion criteria: Elevated serum creatinine (>120 micro mol/L) levels, allergy to contrast material, or acute limb-threatening ischemia were excluded. Because pts under- went MDCT angiography and DSA on different days, potential candidates who lived more than 1 H from our hospital were not asked to enroll.
<u>1° endpoint</u> : N/A <u>Results:</u> • Monitor renal functions for contrast-induced nephropathy • Nephrotoxic meds should be discontinued before contrast administration • Either nonionic iso-osmolar contrast media or	segments and 100% occlusions were correctly identified in all cases by MDCTA. Moderately stenotic segments interpretation in the calves appeared to be more controversial, but no statistical difference in accuracy of MDCTA in the infrapopliteal district arteries was noted with respect to accuracy in the interobserver and intraobserver agreement were observed, with k values greater than 0.80. <u>1° endpoint</u> : Sensitivity and specificity of MDCT angiography in showing arterial occlusions and stenoses of ≥75%. Intertechnique agreement was measured for each anatomic segment, and interobserver agreement was calculated for both techniques. Agreement was quantified using the kappa statistic. <u>Results:</u> The sensitivity and specificity of MDCT angiography for depicting arterial occlusions and 92.2% and 96.8%, respectively. Substantial intertechnique agreement (kappa >0.4) was present in 102 (97.1%) of 105 arterial segments. Substantial intertobserver agreement (kappa >0.4) was present in 102 (97.1%) of 105 arterial segments. Substantial intertobserver agreement was present in 104 (99.0%) of 105 comparisons for both MDCT angiography and 0.78 for DSA. MDCT angiography showed more patent segments than DSA (1,192 vs. 1,091). All 9 segments seen on DSA and not seen on MDCT angiography were in the calves. Of 110 segments seen on MDCT angiography and not seen on DSA, 100 (90.9%) were in the calves.
<ul> <li>Important side effects include hypersensitivity reactions, thyroid dysfunction and contrast-induced nephropathy</li> <li>The knowledge and screening of side effects can allow appreciation and then prompt management.</li> </ul>	<ul> <li>MDCT angiography was accurate in showing arterial atheroocclusive disease with reliability similar to DSA.</li> <li>MDCT angiography showed more vascular segments than DSA, particularly within calf vessels.</li> </ul>

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	acute occlusions, was noted in 1 pt in each group. Venous overlay was observed less frequently in the CB group (CB vs. SB: 12 vs. 19 pts, NS;				
	p=0.04). Contrast bolus overriding without pathological reasons, i.e.,			Size: n=83 pts	
	AO in vessels with relevant (50%– 99%) stenoses (CB: 4.54 vs. SB: 4.18;			Study type: prospective RCT	
for run-off CTA.	16 evaluated segments. Patency- based comparison revealed superior			run-off CTA.	
protocol. Ineretore, a CB	was significantly higher (p<0.05) in 7 of	134 mL, 4 mL/sec).		equi-iodine doses for	
compared to the SB	bolus (CB: 279±57HU, SB:	Comparator: The SB protocol	Exclusion criteria: N/A	medium with a SB	
quantitatively and	scale). Overall arterial CD was	100 mL, 4 mL/sec)	40 or 64-slice run-off CTA	high-iodine	22473508
<ul> <li>At equi-iodine doses, the CB protocol led to a</li> </ul>	<u>1° endpoint</u> : Luminal CD values were measured and AO was scored (5-point	(32 pts, iomeprol 400mgl/mL,	Inclusion criteria: 64 pts with suspected PAD who underwent	Aim: Compare a CB injection protocol using	Meyer BC, et al. 2012 (88)
Study Limitations; Adverse Events	or RR; & 95% CI)	Study Comparator (# patients)		Study Size (N)	Year Published
Relevant 2° Endpoint (if any);	Endpoint Results (Absolute Event Rates, P value; OR	Study Intervention (# patients) /	Patient Population	Aim of Study; Study Type;	Study Acronym; Author;
	ography)–Section 3.3.	Evidence Table 7. RCTs of Imaging for Anatomic Assessment (Ultrasound, CTA, MRA, Angiogra	ior Anatomic Assessment (L	7. RCTs of Imaging f	<b>Evidence Table</b>
phy.	artery disease; PAOD, peripheral arterial occlusive disease; PPV, positive predictive value; pt, patient; and WBMRA, whole-body magnetic resonance angiography.	ve value; pt, patient; and WBMRA, v	sive disease; PPV, positive predictiv	D, peripheral arterial occlu	artery disease; PAO
ubtraction angiography; osclerosis; MDCTA, ve value: PAD_peripheral	ABI indicates ankle-brachial index; CE-MRA, contrast-enhanced MRA; CI, confidence interval; CT, computed tomography; DM, diabetes mellitus; DSA, digital subtraction angiography; DSCT, dual source computed tomography; DUS, duplex ultrasonography; IC, intermittent claudication; IPA, internal pudendal artery; LEA, lower extremity atherosclerosis; MDCTA, DSCT, dual source computed tomography; DUS, duplex ultrasonography; IC, intermittent claudication; IPA, internal pudendal artery; LEA, lower extremity atherosclerosis; MDCTA, multidetector computed tomography angiography. MDCT multidetector computed tomography angiography.	nce interval; CT, computed tomogra rmittent claudication; IPA, internal pu tomography: N/A not applicable: N	ontrast-enhanced MRA; Cl, confide S, duplex ultrasonography; IC, inter hv: MDCT_multidetector computed	orachial index; CE-MRA, co computed tomography; DU	ABI indicates ankle-t DSCT, dual source c multidetector comput
and the management of pts receiving metformin	and the matformin				
used to reduce the incidence of CIN,	used to re	• N/A			
medium, the prophylactic measures	medium, t	Results:	Exclusion criteria: N/A	Size: N/A	21866433
<ul> <li>Topics reviewed include the definition of CIN, the choice of contrast</li> </ul>	Topics r      definition	<u>1° endpoint</u> : N/A	Inclusion criteria: N/A	Study type:	Stacul F, et al. 2011(87)
	cysteine may be	<ul> <li>In nign-risk pts N-acetylcysteine may be administered.</li> </ul>			
	Iraged.	<ul> <li>Evidential for the second secon</li></ul>			
	trast media use to be	nonionic low-osmolar contrast media use to be favored			

	Favaretto E, et al. 2007(91) <u>17443099</u>	Met R, et al. 2009(90) <u>19176443</u>	Fraioli F, et al. 2006(89) <u>15988586</u>	
Study type: Prospective series	<b>Aim:</b> Investigate the agreement between DSA in the diagnosis of stenosis	accuracy of low dose MDCT with DSA for the detection of aortoiliac and PAD. <u>Study type</u> : RCT <u>Size</u> : n=75 pts <u>Aim</u> : To determine the accuracy of CTA compared with intra- arterial DSA in differentiating extent of disease in pts with PAD <u>Study type</u> : Meta- analysis CTA vs. DSA <u>Size</u> : n=909 studies	<u>Aim</u> : Compare the influence of radiation dose on image quality and diagnostic	
Exclusion criteria: N/A	Inclusion criteria: IVA artery disease (claudication, critical ischemia, or skin lesions)	MDCT angiography of the aorta and peripheral vessels. <b>Exclusion criteria:</b> Renal insufficiency (serum creatinine >2 mg/dl), contra-indication to iodinated contrast, respiratory failure, congestive heart failure and poor general condition of the pt. <b>Inclusion criteria:</b> • Reviews of effectiveness for studies comparing CTA with intra-arterial DSA for PAD • Compared multidetector CTA with intra-arterial DSA Included at least 10 pts with IC or CLI • Aimed to detect >50% stenosis or arterial occlusion • Presented either 2 x 2 or 3 x 3 contingency tables (≤50% stenosis vs. >50% stenosis or occlusion), or provided data allowing their construction	Inclusion criteria: Onsecutive pts, with a clinical Dx of obstructive arterial disease of the extremities underwent	
Results: Kappa=0.70; 95% CI: 0.588–0.825 for the whole arterial axis. Agreement was	1° endpoint: Diagnostic accuracy of duplex for detected lesion severity of LE PAD	were kept constant, except for the mAs. Comparator: 50 mAs vs. 100 mAs vs. 130 mAs for detecting PAD (>50% stenosis) <u>Results</u> : Sensitivity stenosis >50% (95%CI: 92–9); specificity 96%(95% CI: 93–97)	Intervention: Pt population was randomly divided into three groups of 25 pts. In each group, MDCT scanning parameters	
	The sensitivity and specificity of duplex compared to angiography is modest	degree of stenoses revealed a sensitivity, specificity, accuracy, PPV and NPV of 96%, 94%, 95%, 83% and 99% for Group A (50 mAs), 96%, 96%, 96%, 89% and 99% for Group B (100 mAs) and 98%, 96%, 97%, 91% and 100% for the standard dose protocol, Group C (130 mAs). CTA had adequate sensitivity for detecting PAD	<ul> <li>1° endpoint:</li> <li>The dose reduction was 74% for group A and 40% for group B.</li> <li>The evaluation of the presence and</li> </ul>	29 of 64 legs [45%] vs. 44 of 64 legs [69%]; p=0.01).
	N/A	<ul> <li>Peripheral vessels.</li> <li>This technique provides substantial reduction of the radiation dose delivered to the pt while maintaining optimal diagnostic accuracy.</li> <li>N/A</li> </ul>	<ul> <li>Low-dose scanning is thus a feasible and accurate option for 4-row CT angiography of the</li> </ul>	

McCullough PA, 2011(93) <u>21609484</u>	Kau T, et al. 2011 (92) 21365195	
<u>Aim</u> : To compare discomfort rates in pt- reported outcomes related to IOCM with LOCM	<u>Aim</u> : Evaluate the accuracy of DE-CTA maximum intensity projections <u>Study type</u> : Prospective series DE-CTA vs. angio <u>Size</u> : n=58	Duplex vs. angio <u>Size</u> : n=49 pts
Inclusion criteria: Studies with intra-arterial administration of CM. Exclusion criteria: Studies with intravenous	Inclusion criteria: Pts with sx peripheral arterial occlusive disease Exclusion criteria: in ability to get CTA get CTA	
<u>Intervention</u> : IOCM (Iodixanol) (3,385) <u>Comparator</u> : LOCM (4,796)	1° endpoint: Diagnostic accuracy of DE-CTA to detect stenosis severity <u>Results:</u> In DSA, 52.3% of segments were significantly stenosed or occluded. Agreement of DE-CTA MIPs with DSA was good in the aorto- iliac and femoro-popliteal regions (kappa=0.72; kappa=0.66), moderate in the crural region (kappa=0.72; kappa=0.66), moderate in the crural region (kappa=0.55), slight in pedal arteries (kappa=0.10) and very good in bypass segments (kappa=0.81). Accuracy was 88%, 78%, 74%, 55% and 82% for the respective territories and moderate (75%) overall, with good sensitivity (67%). Sensitivity and specificity was 82% and 76% in claudicants and 84% and 61% in pts with CLI.	good for the aorto-iliac district (kappa=0.63) with a sensitivity of 63% and a specificity of 96%, and for the femoro-popliteal district (kappa=0.70) with a sensitivity of 74% and a specificity of 83%. In infrapopliteal arteries, kappa showed a poor agreement.
<ul> <li>1° endpoint:</li> <li>● Pain:</li> <li>Pts receiving IOCM vs. various LOCMs (RD: -0.049; 95% CI: -0.076 – -0.021; p=0.001). IOCM was favored over all LOCMs combined with a summary RD:</li> </ul>	DE-CTA had good diagnostic accuracy above the knee. Below the knee the diagnostic accuracy was modest at best and worse when arteries were calcified.	
<ul> <li>Cold sensation: NS difference</li> <li>IOCM was found to have less frequent and severe pain and warmth during administration as</li> </ul>	N/A	

Study Acronym; Author; Year Published         Study Type/Design; Study Size         Patient Population         Primary Endpoint and Results (include P value; OR or RR; 2013(94)           Sultan S. et al. 23711680         Study Type: Cross- study         Inclusion criteria: intervention for vascular intervention for PAD, AAA, or carotid         Inclusion criteria: intervention for PAD, AAA, or carotid         1         endpoint: Prevalence of AAA, CAD, and isease defined as disease in 2 territories. Poly vascular vascular intervention for PAD, AAA, or carotid         1         endpoint: Prevalence of AAA, CAD, and isease defined as disease in 2 territories.           Kurvers HA, et al.         Study Type: Cross- study         Inclusion criteria: Enrolled sectional single center study referred to study         Inclusion criteria: Enrolled a vascular center with sx peripheral atherosclerosis in increased prevalence 6.5% in PAD pts vs1% pits         1           Grøndal N, et al.         Study Type: Danish intervention arm of streament for AAA.         Exclusion criteria: N/A 65-74 y who were screened for AAA.         1           25923784         Size: n=25.083 men who uscree correct de for AAA.         Exclusion criteria: N/A in 10.9%.         1	pts     elevated risk factors (e.g. DM)       N, et al.     Study type: Danish intervention arm of screening trial     Exclusion criteria: N/A       Size: n=25,083 men who     Size: n=25,083 men who       Size: n=25,083 men who     Exclusion criteria: N/A
Acronym; utthor; Published       Study Type/Design; Study Size       Patient Population         Published       Study type: Cross- sectional single-center study       Inclusion criteria: Intervention for 1 of the PVD territories. Poly vascular disease defined as disease in ≥2 territories. Poly vascular disease defined as disease in ≥2 territories.         HA, et al.       Study type: Cross- study       Inclusion criteria: N/A disease         HA, et al.       Study type: Cross- study       Inclusion criteria: N/A disease         Study type: n=2,274 vascular pts       Inclusion criteria: Enrolled a vascular center with sx peripheral atherosclerosis in some arterial territory or elevated risk factors (e.g. DM)         N, et al.       Study type: Danish intervention arm of screening trial       Exclusion criteria: Men age for AAA.	Pts     elevated risk factors (e.g. DM)       N, et al.     Study type: Danish intervention arm of screening trial       M     Study type: Danish for AAA.
Acronym;       Study Type/Design;       Patient Population         uthor;       Study type: Cross- sectional single-center       Inclusion criteria:         0       Study type: Cross- sectional single-center       Intervention for 1 of the PVD territories. Poly vascular disease defined as disease         1A, et al.       Study type: Cross- vascular intervention for PAD, AAA, or carotid disease       Inclusion criteria: N/A tisease         1A, et al.       Study type: Cross- study       Inclusion criteria: N/A avascular center         1A, et al.       Study type: Cross- study       Inclusion criteria: N/A avascular center         1A, et al.       Study type: Cross- study       Inclusion criteria: N/A avascular center         1A, et al.       Study type: Cross- study       Inclusion criteria: N/A avascular center with sx peripheral atherosclerosis in some arterial territory or elevated risk factors (e.g. DM)         1D       Study type: Danish       Inclusion criteria: N/A Inclusion criteria: M/A	pts elevated risk factors (e.g. DM) Study type: Danish Inclusion criteria: N/A
Acronym;       Study Type/Design;       Patient Population         uthor;       Study type: Cross- sectional single-center       Inclusion criteria:         et al.       Study type: Cross- sectional single-center       Intervention for 1 of the PVD territories. Poly vascular disease defined as disease in ≥2 territories.         4A, et al.       Study type: Cross- vascular intervention for PAD, AAA, or carotid disease       Inclusion criteria: N/A disease         4A, et al.       Study type: Cross- study       Inclusion criteria: N/A         5       Study type: Cross- study       Inclusion criteria: N/A         9       Size: n=2,274 vascular pts       Inclusion criteria: Enrolled some arterial territory or elevated risk factors (e.g. DM)	elevated risk factors (e.g. DM)
Acronym;Study Type/Design; Study SizePatient PopulationPublishedStudy type: Cross- sectional single-center studyInclusion criteria: Intervention for 1 of the PVD territories. Poly vascular disease defined as disease1A, et al.Study type: Cross- sectional single center PAD, AAA, or carotid diseaseInclusion criteria: territories.1A, et al.Study type: Cross- sectional single center sectional single center studyInclusion criteria: territories.1A, et al.Study type: Cross- sectional single center sectional single center in SMART study referred to a vascular center with sx peripheral atherosclerosis in some arterial territory or	
Acronym;Study Type/Design; Study SizePatient PopulationPublishedStudy type: Cross- sectional single-center studyInclusion criteria: Intervention for 1 of the PVD territories. Poly vascular disease defined as diseaseQSize: 328 pts having a vascular intervention for PAD, AAA, or carotidInclusion criteria: Exclusion criteria: N/A diseaseHA, et al.Study type: Cross- sectional single center sectional single centerInclusion criteria: in SMART study referred to a vascular center with sx	peripheral atherosclerosis in some arterial territory or
Acronym;       Study Type/Design;       Patient Population         uthor;       Study Size       Inclusion criteria:         Published       Study type: Cross-       Inclusion criteria:         et al.       Study type: Cross-       Intervention for 1 of the PVD         sectional single-center       Intervention for 1 of the PVD         study       study       territories. Poly vascular         disease       Size: 328 pts having a       in ≥2 territories.         vascular intervention for       PAD, AAA, or carotid       Exclusion criteria: N/A         disease       Study type: Cross-       Inclusion criteria: Enrolled         sectional single center       in SMART study referred to	a vascular center with sx
Acronym;       Study Type/Design;       Patient Population         uthor;       Study Size       Inclusion criteria:         Published       Study type: Cross-       Inclusion criteria:         et al.       Study type: Cross-       Intervention for 1 of the PVD         sectional single-center       Intervention for 1 of the PVD         study       territories. Poly vascular         disease defined as disease         vascular intervention for         PAD, AAA, or carotid         Exclusion criteria: N/A	HA, et al. Study type: Cross- Inclusion criteria: Enrolled sectional single center in SMART study referred to
Acronym;Study Type/Design; Study SizePatient Populationuthor;Study SizePublishedStudy type: Cross- sectional single-centerInclusion criteria: Intervention for 1 of the PVD territories. Poly vascular disease defined as disease0Size: 328 pts having a vascular intervention for	AA, or carotid
Acronym;       Study Type/Design;       Patient Population         uthor;       Study Size         Published       Inclusion criteria:         et al.       Study type: Cross-         sectional single-center       Intervention for 1 of the PVD         study       territories. Poly vascular         disease defined as disease	in ≥2 territories.
Acronym;     Study Type/Design;     Patient Population       uthor;     Study Size       Published     Inclusion criteria:       et al.     Study type: Cross-       sectional single-center     Intervention for 1 of the PVD       or study     study	disease defined as disease
nym; Study Type/Design; Patient Population ; Study Size shed	et al. <u>Study type</u> : Cross- <u>Inclusion criteria:</u> sectional single-center Intervention for 1 of the PVD
Study Type/Design; Patient Population Study Size	shed
	Study Acronym; Study Type/Design; Patient Population Primary Endpo Author; Study Size (include P va

Study Acronym;	Study Type/Design;	Patient Population	Primary Endpoint and Results	Summary/Conclusion
Author;	Study Size		(include P value; OR or RR;	Comment(s)
Lee JY, et al.	Study type: Cohort	Inclusion criteria: Pts	1° endpoint: Prevalence of abnormal ABI	Doesn't really say the prevalence of CAD in all pts
2013(98)		having coronary	<0.9 or >1.4 and MACE over 3 y.	with abnormal PAD. It looks at a select group who had
24355120	Size: n=2,424 pts with	angiography		cath and then looks at the impact of PAD on outcomes
	CAD and 119 pts without		Results:	over 3 y.
	significant CAD on cath	Exclusion criteria: Pts	• In CAD pts: 14% had ABI <0.9, vs. 4% in	<ul> <li>Shows prognostic value of low ABI for MACE but does</li> </ul>
		with known PAD or prior	pts without CAD. Of the 390 pts with	not provide information on the value of screening for
		ABI	abnormal ABI, 130 (33%) had coronary	CAD in pts with low ABI
			revascularization at time of cath. 3 y	
			MACE significantly higher with abnormal	
			ABI (15.7% vs. 3.3%; p<0.001).	
			<ul> <li>Abnormal ABI HR: 1.87 or 2.40 on</li> </ul>	
			propensity matched analysis.	
Moyer VA and U.S.	Study type: Review of	Inclusion criteria: All	1° endpoint: N/A	<ul> <li>USPSTF summary statement concluding that</li> </ul>
Preventative Services	studies assessing ABI	studies examining the		screening for PAD using the ABI in asx individuals is not
Task Force		prognostic value of	Resulte: See how to right More useful for	

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	1,8749 attended the			
	screening (uptake 74.7%).			
Giugliano G, et al.	Study type: Prospective	Inclusion criteria: 213	1° endpoint: Prevalence of AAA in pts with PAD	<ul> <li>Small study showed that prevalence of AAA</li> </ul>
2012(96)	case series	consecutive pts with PAD		in pts with PAD is much higher than in the
23173942		screened for AAA	Results: AAA was present in 19 pts (9%) with	general population.
	Size: n=213 consecutive		similar prevalence in men and women.	<ul> <li>Prevalence related to age:</li> </ul>
	pts	Exclusion criteria: N/A		<55 y: 0
				55-64 y: 5.1%
				65-74 y: 11.4%
				>75 y: 15.8%
Barba A	Study type:	Inclusion criteria: 1,166	1° endpoint: Prevalence of AAA in pts with PAD	<ul> <li>Prevalence of AAA in pts with PAD is higher</li> </ul>
2005(97)	Observational descriptive	consecutive pts with PAD		than in the general population.
<u>15963741</u>	study	had AAA screening	Results: Prevalence of AAA in men was 13.6%	<ul> <li>As in other studies, the prevalence of AAA in</li> </ul>
			and in women 4.1% but there were only 73	pts with PAD increased with age.
	Size: n=1,166 pts with	Exclusion criteria: None	women.	<ul> <li>The prevalence was much higher in men</li> </ul>
	PAD			than women.
AAA indicates abdomina	al aortic aneurysm; CAD, coron	ary artery disease; CVD, cardio	AAA indicates abdominal aortic aneurysm; CAD, coronary artery disease; CVD, cardiovascular disease; N/A, not applicable; PAD, peripheri	PAD, peripheral artery disease; and PVD, peripheral vascular
disease.				

Evidence Table 11. Nonrandomized Trials, Observational Studies, and/or Registries of Screening in Carotid Artery Disease–Section 4.3 S

Comment(s)	(include P value; OR or RR;		Study Size	Author;	
Summary/Conclus	Primary Endpoint and Results	Patient Population	Study Type/Design;	Study Acronym;	

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an Heart Association, Inc. and American College of Cardiology Foundatior	
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sten sveve	scheduled for vascular ever surgery and <u>Size</u> : n=5,859 pts	15625331and coronaryreparevascularization foroperCAD in high-risk ptsat in	McFalls EO, et al. Study type: RCT of Inclu 2004(100) cardiac catheterization for n	Evidence Table TV. KCTS for CAD Screening in PAD-Section 4.2         Study Acronym;       Aim of Study;       Patient Population         Author;       Study Type;         Year Published       Study Size (N)
Exclusion criteria: Left main stenosis >50%, LVEF <20%, severe aortic stenosis	on	ar	Inclusion criteria: Pts scheduled for major vascular surgery (AAA	Patient Population
	<u>Comparator</u> : No revascularization before elective major vascular surgery	elective major vascular surgery	Intervention: Revascularization before	Study Intervention (# patients) / Study Comparator (# patients)
14% in the no-CAD revascularization group.	revascularization group and 23% in the CAD revascularization group. 30 d postoperative MI=12% in the CAD revascularization group and	<u>Results:</u> No difference in outcomes. Mortality at 2.7 y was 22% in the no-CAD	<u>1° endpoint</u> : Long-term mortality	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% Cl)
perioperative rates of MI.	<ul> <li>Excludes left main disease</li> <li>No advantage to screening for CAD in pts having elective major vascular surgery on mortality or</li> </ul>	revascularization group and 14% in the no-CAD revascularization group.	<ul> <li>No difference in 30 d postoperative MI=12% in the CAD</li> </ul>	Relevant 2° Endpoint (if any); Study Limitations; Adverse Events

HR, hazard ratio; MACE, major adverse cardiovascular events; N/A, not applicable; NRI, net reclassification improvement; PAD, peripheral artery disease, pt, patient; and USPSTF, United	NRI, net reclassification improvement; PAD, pe	events; N/A, not applicable; I	najor adverse cardiovascular	HR, hazard ratio; MACE, major adverse
ABI indicates ankle-brachial index; asx, asymptomatic; CAD, coronary artery disease; CTA, computed tomographic angiography; CT, computed tomography; FRS, Framingham risk score;	e; CTA, computed tomographic angiography; C	CAD, coronary artery disease	al index; asx, asymptomatic;	ABI indicates ankle-brachi
screening ABI in asx individuals		Exclusion criteria: N/A	<u>Size</u> : n=52,510	
<ul> <li>This is more useful for the assessment of the value of</li> </ul>				
NRI small when adding ABI to FRS	FRS	predictor of CAD events	Framingham risk score.	
	Results: NRI small when adding ABI to	value of ABI as a	of ABI in addition to	<u>24156115</u>
above (99)		Studies assessing the	studies assessing value	2013(30)
USPSTF analysis supporting the summary statement	1° endpoint: Test characteristics and NRI	Inclusion criteria:	Study type: Review of	Lin JS, et al.
screening ABI in asx individuals				
This is more useful for the assessment of the value of				
significant				
reported or indicates a change that may not be clinically		Exclusion criteria: N/A		
ABI to CAD events, but that the NRI is often not	ABI		Size: N/A	24026320
an • They find several studies showing a relationship of low	question addressing asx screening with an	screening ABI in asx pts.		2013(99)

States Preventative Services Task Force.

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Year Published			& 95% CI)	
Sultan S, et al.	Study type: Cross-	Inclusion criteria: Intervention for 1 of	1° endpoint: Prevalence of AAA, CAD, and	<ul> <li>Looks at the risk according to multiple</li> </ul>
2013(94)	sectional single-center	the PVD territories. Poly vascular	carotid disease in PAD pts receiving	vascular beds not just PAD
<u>23711680</u>	study	disease defined as disease in ≥2	revascularization	Can't discern the risk of AAA or CVD
		territories.		with PAD alone
	Size: n=328 pts		Results: Poly-vascular bed pts had about 8X	
	having a vascular	Exclusion criteria: N/A	the risk of carotid disease or AAA.	
	intervention for PAD,			
	AAA, or carotid			
	disease			
Kurvers HA, et al.	Study type: Cross-	Inclusion criteria: Enrolled in SMART	1° endpoint: Prevalence of carotid stenosis	<ul> <li>Select sx atherosclerosis population</li> </ul>
2003(95)	sectional single center	study referred to a vascular center with		
<u>12764269</u>	study	sx peripheral atherosclerosis in some	Results: Prevalence 12.5% in PAD pts vs.	
		arterial territory or elevated risk factors	~2% for risk factor only pts. Age >54 y and	
	Size: n=2,274	(e.g. DM)	PAD increased prevalence to 22%.	
	vascular pts			
		Exclusion criteria: N/A		
AAA indicates abdominal pt, patient; PVD, peripher	AAA indicates abdominal aortic aneurysm; ABI, ankle-b pt, patient; PVD, peripheral vascular disease; sx, symp.	e-brachial index; CAD, coronary artery dise np.	AAA indicates abdominal aortic aneurysm; ABI, ankle-brachial index; CAD, coronary artery disease; CVD, cardiovascular disease; DM, diabetes mellitus; PAD, peripheral artery disease; pt, patient; PVD, peripheral vascular disease; sx, symp.	nellitus; PAD, peripheral artery disease;

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Evidence Table 12.	Nonrandomized Trials,	Observational Studies, and/c	Evidence Table 12. Nonrandomized Trials, Observational Studies, and/or Registries for Renal Artery Disease-Section 4.4.	tion 4.4.
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			& 95% CI)	
Olin JW, et al.	Study type: Single	Inclusion criteria: Pts who	1° endpoint: Prevalence of >50% renal artery	<ul> <li>There is a high prevalence of</li> </ul>
1990(101)	center, retrospective	underwent catheter angiography	stenosis	incidental renal artery stenosis in pts
<u>2368764</u>	cohort study	for evaluation of AAA, Aortoiliac		with atherosclerosis in other locations,
		Occlusive Disease and PAD.	Results: Prevalence was 38% in pts with AAA,	even in the absence of clinical clues to
	Size: n=395 consecutive		33% with AOD and 39% with PAD.	suspect RAS.
	pts	Exclusion criteria: N/A		
Leertouwer TC, et al.	Study type: Single	Inclusion criteria: Pts who	1° endpoint: Prevalence of >50% renal artery	<ul> <li>Incidental renal artery stenosis is</li> </ul>
2001 (102)	center, retrospective	underwent catheter based	stenosis	common in pts with PAD
<u>11260411</u>	cohort study	angiography for evaluation of		<ul> <li>Renal replacement therapy did not</li> </ul>
		PAD	Results: 126 (33%) had >50% stenosis.	occur in any of these pts thus
	Size: n=386 consecutive			revascularization to prevent ESRD is
	pts	Exclusion criteria: N/A		not indicated in most pts.
CHS	Study Type: Multicenter,	Inclusion criteria: Free living pts	1° endpoint:	<ul> <li>This is the 1st population based</li> </ul>
Hansen KJ, et al.	longitudinal cohort study	age >65 y were invited to	Prevalence of RAS in a free standing elderly	estimate of the prevalence of RVD
2002(103)		undergo renal artery duplex	population	among free living, elderly black and

		Exclusion criteria: N/A	<ul> <li>834 (96%)</li> <li>Prevalence</li> <li>No differen</li> <li>black pts.</li> </ul>	<ul> <li>834 (96%) were technically adequate to define the presence or absence of RVD</li> <li>Prevalence of RAS was 6.8%.</li> <li>No difference in prevalence between white and black pts.</li> </ul>	
AAA indicates; AOD, a renal vascular disease	, arterial occlusive disea lse.	se; ESRD, end-stage renal disease; N	A, not applicable; PAD	AAA indicates; AOD, arterial occlusive disease; ESRD, end-stage renal disease; N/A, not applicable; PAD, peripheral artery disease; pt, patient; RAS, renal artery stenosis; and RVD renal vascular disease.	l artery stenosis; and RVD,
Evidence Table	13. RCTs Evaluatin	Evidence Table 13. RCTs Evaluating Antiplatelet Agents- Section 5.1	n 5.1.		Deleteration Factoriat (if
Acronym;	Aim of study; Study Type;	Patient Population	(# patients) /	(Absolute Event Rates, P value; OR or RR; &	Re
Year Published			(# patients)	55 /0 <b>C</b> IJ	Adverse Events
POPADAD	Aim: To determine	Inclusion criteria: Aged ≥40 y	Intervention and	<u>1° endpoint:</u>	Adverse effect (effect
Belch J, et al.	whether ASA and	with type 1 or type 2 DM and an	comparator: Daily,	Death from coronary heart disease or stroke,	<u>estimates)</u> :
2008(16)	antioxidant therapy,	ABI of ≤0.99 but no sx	100 mg ASA tablet	nonfatal MI or stroke, or amputation above the	<ul> <li>Malignancy 0.76 (0.52–</li> </ul>
<u>18927173</u>	combined or alone,	cardiovascular disease	+ antioxidant	ankle for CLI; and death from CHD or stroke	1.11),
	are more effective	1 - - -	capsule (n=320),	<ul> <li>116 of 638 primary events occurred in the</li> </ul>	<ul> <li>Gastrointestinal bleeding,</li> </ul>
	than placebo in	<b>Exclusion criteria</b> : People with	ASA tablet +	ASA groups compared with 117 of 638 in the	0.90 (0.53–1.52)
	development of	who use ASA or antioxidant	(n=318), placebo	95% Cl: 0.76–1.26. 43 deaths from coronary	1.08).
	cardiovascular	therapy on a regular basis; those	tablet + antioxidant		<ul> <li>Allergy 1.14 (0.80–1.63)</li> </ul>
	events in pts with DM	with peptic ulceration, severe	capsule (n=320), or	groups compared with 35 in the no ASA groups	
	and asx PAD.	dyspepsia, a bleeding disorder, or intolerance to ASA: those with	placebo tablet + placebo capsule	(6.7% vs. 5.5%): HR: 1.23; 95% CI: 0.79–1.93).	
	Study type:	suspected serious physical illness	(n=318)		
	Multicenter,	(such as cancer), which might have			
	randomized, double	been expected to curtail life			
	hlind Ox0 fantarial	avnortanov: those with nevrohistric			

12218965

Size: n=870 pts

ultrasound

Results:

white Americans

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placebo controlled trial. blind, 2×2 factorial,

illness (reported by their GP); those with congenital heart expectancy; those with psychiatric

Size: n=1,276 pts

informed consent

disease; and those unable to give

	or dSA and a high- dose antioxidant vitamin combination in pts with PAD in terms of reduction of the risk of a first vascular event (MI, stroke, vascular death) and CLI. <b>Study type:</b> Randomized, placebo-controlled, double-blind clinical trial with 2x2 factorial designs. <u>Size</u> : n=366 pts		on screening the cur general population. anti age <u>Study type</u> : hac Randomized Controlled Trial con Size: n=3,350 pts her the	a low ABI identified	a low ABI identified	ASA in preventing events in people with a low ABI identified
	sound, with ankle/brachial x <0.85 or toe index <0.6 ntaine stage III or IV PVD; life pectancy <24 mo; vascular ery or angioplasty in the last 3 egnancy or lactation; ntraindication to ASA; ajor cardiovascular events iring antiplatelet therapy; rticipation in another clinical cooperative pts; eatment with drugs that fere with hemostasis, such as prostanoids, peripheral dilators, ASA and/or	Ð	d d e;	gina, or PAD;		
		Intervention and	Placebo			
	<ul> <li>Incluence of ratal ratio noninatal vasculate events (MI, stroke and pulmonary embolism) and critical leg ischemia</li> <li>7 of 185 ASA and 20 of 181 placebo pts suffered a major vascular event (risk reduction 64%, p=0.022)</li> <li>5 ASA and 8 placebo pts, respectively, suffered critical leg ischemia (total 12 vs. 28, p=0.014)</li> <li>Safety endpoint: Incidence of bleeding 4 in ASA and 0 in placebo (p=0.99)</li> </ul>	1° endpoints:	<ul> <li>between groups (13./ events per 1000 person- years in the ASA group vs. 13.3 in the placebo group; HR: 1.03; 95% Cl; 0.84–1.27)</li> <li>Major hemorrhage</li> <li>Initial event of major hemorrhage requiring admission to hospital occurred in 34 pts (2.5 per 1000 person-years) in the ASA group and 20 (1.5 per 1000 person-years) in the placebo group (HR: 1.71; 95% Cl: 0.99–2.97).</li> </ul>		No statistically significant difference was found	revascularization No statistically significant difference was found
42		<ul> <li>76% with type 2 DM</li> </ul>	ischemic attack; no statistically significant difference between groups (22.8 events per 1000 person-years in the ASA group vs. 22.9 in the placebo group; HR: 1.00; 95% CI: 0.85–1.17) • All-cause mortality no significant difference in all-cause mortality between groups (176 vs. 186 deaths, respectively; HR: 0.95; 95% CI: 0.77–1.16)	c	angina, IC or transient	primary endpoint event or angina, IC or transient

Minar E, et al. 1995(106) <u>7697845</u>	Horrocks M, et al. <u>9257670</u>																																											
<u>Aim</u> : To compare the effects of high-dose (1000 mg/d) and low- dose (100 mg/d) ASA on long-term patency after femoropopliteal angioplasty. <u>Study type</u> : Randomized <u>Size</u> : n=216 pts	<u>Aim</u> : To investigate the effects of 2 platelet inhibitors, ASA and iloprost, on platelet uptake and restenosis at the site of angioplasty in pts undergoing femoral or popliteal angioplasty. <u>Study type:</u> Prospective, randomized <u>Size</u> : n=43 pts																																											
Inclusion criteria: Pts treated successfully by percutaneous transluminal angioplasty for femoropopliteal lesions Exclusion criteria: Failed PTA, recent gastroduodenal ulcer, life expectancy <2 y, severe renal insufficiency, need for ongoing nonsteroidal, unable to consent	supplementary vitamins that could not be discontinued or had to be started. <u>Inclusion criteria</u> : Pts undergoing femoral or popliteal angioplasty <u>Exclusion criteria</u> : Bleeding disorder, ulcer disease																																											
Intervention and <u>Comparator</u> : 1000 or 100 mg ASA daily.	Intervention: ASA (300 mg/d), iloprost (8 H/d IV infusion) or no antiplatelet medication during angioplasty and on the subsequent 2 d.																																											
<u>1° endpoint</u> : Long-term (24 mo) patency 36 pts in the high-dose and 36 in the low-dose ASA group, developed angiographically verified reobstruction within the recanalized segment. By intention-to-treat analysis, the cumulative patency rates at 24 mo were 62.5% in the high- dose and 62.6% in the low-dose ASA group (Wilcoxon, p=0.97; log-rank, p=0.97). The cumulative survival at 24 mo of follow-up was 86.6% in the high-dose and 87.7% in the low- dose ASA group. <u>Safety endpoint</u> : Discontinued therapy for gastrointestinal symptoms, 4 in high dose and 0 in low dose Discontinued therapy 30 high dose and 11 low dose (p<0.01)	<ul> <li><u>1° endpoint:</u></li> <li>Platelet uptake was measured using 111 Indium-labelled platelets. Restenosis was assessed by repeat angiography at 3 mo and clinical symptoms up to 12 mo.</li> <li>Median changes in platelet uptake were similar in the 3 treatment groups, but all platelet radioactivity ratios &gt;2.0 occurred in the control group. Restenosis at 3 mo was observed in 3 control, 5 ASA and 1 iloprost pt.</li> <li>Further surgical intervention was performed in 3 control and 3 ASA pts, but in none of the iloprost pts up to 12 mo after angioplasty</li> </ul>																																											
<ul> <li>100 mg as effective as 1000 mg</li> <li>Treatment started 3 d after PTA</li> </ul>	• Limited utility as iloprost also utilized																																											
																																										8918275	1996 (107)	CAPRIE
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																													-	Size: n=19,185 pts		Randomized, blinded	Study type:		vascular death	stroke, MI, or	cluster of ischemic	a composite outcome	in reducing the risk of	(325 mg once daily)	once daily) and ASA	clopidogrel (75 mg	relative efficacy of	Aim: To assess the
women of childbeating age not	Waman of abildboaring and not	<ul> <li>Hx of ASA sensitivity</li> </ul>	affecting platelet function	anuplatelet drugs of NSAIDS	terni anticoagaiante, non-suay	term anticoaciulants non-study	<ul> <li>Anticipated requirement for long-</li> </ul>	differential, or platelet count	<ul> <li>Known to have abnormal WBC,</li> </ul>	or hepatic abnormalities		Hy of drug-induced bematologic	neutropenia	<ul> <li>Hx of thrombocytopenia or</li> </ul>	systemic bleeding	<ul> <li>Hx of haemostatic disorder or</li> </ul>	bleeding			inclufficiency	<ul> <li>Severe renal or hepatic</li> </ul>	<ul> <li>Contraindications to study drugs:</li> </ul>	<ul> <li>Scheduled for major surgery</li> </ul>	Uncontrolled hypertension	pt's life expectancy to less than 3 y	severe comorbiality likely to limit	alone after qualifying event		Dt unlikely to be discharged	angiography	carotid and arteractomy or	<ul> <li>Dualifying stroke induced by</li> </ul>	after qualifying stroke	demented Carotid endarterectomy	lead to nt heing hedridden or	<ul> <li>Severe cerebral deficit likely to</li> </ul>	• Age <21 y	Exclusion criteria:		PAD	ischemic stroke, recent MI, or sx	manifested as either recent	atherosclerotic vascular disease	Inclusion criteria: Pts with
																																							325 mg per d	Comparator: ASA		per d	Clopidigrel 75 mg	Intervention:
																																grouns	Cafate and maint Blooding similar in the 2		$p = 0.0 \pm 0.7$ . A control (05% CF) 0.3 - 16.5		vascular death lower than 5.83% with ASA	annual 5.32% risk of ischemic stroke. MI, or	showed that pts treated with clopidogrel had an	cluster on which an intention-to-treat analysis	<ul> <li>1960 first events included in the outcome</li> </ul>	stroke, MI, or vascular death	<ul> <li>Composite outcome cluster of ischemic</li> </ul>	<u>1° endpoint:</u>
									ASA.	the use of clopidogrel vs.	from clopidogrel over ASA	subgroup clearly benefited			of ato with roopst stroko	the 3 nredefined subarouns	heing observed between	treatment effect (p=0.042)	statistical heterogeneity of	primary endpoint, with	was observed for the	significant result (p=0.043)	<ul> <li>Iviarginally statistically</li> </ul>			and 16 (0.17%) in the $\Delta C \Delta$	neutrophils (<1.2 x 109/1)	significant reductions in	clonidoarel aroun with	(0.10%) pts in the	respectively There were 10	(0.52% vs. 0.72%).	astrointestinal hemorrhade	(0.33% vs_0.47%) and	intracranial hemorrhade	(0.97% vs. 1.22%).	gastrointestinal discomfort	0.11%), upper	0.10%), diarrhea (0.23% vs.	included rash (0.26% vs.	judged to be severe	clopidogrel and ASA groups	experiences in the	<ul> <li>Reported adverse</li> </ul>

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CHARISMA Bhatt DL, et al. 2007(109) 17498584	<b>CHARISMA</b> Cacoub PP, et 2009(108) <u>19136484</u>
Aim: To determine whether there is benefit of clopidoprel + ASA in a subpopulation of CHARISMA	<u>Aim</u> : To determine whether clopidogrel + ASA provides greater protection against major cardiovascular events than ASA alone in pts with PAD. <u>Study type:</u> Substudy of Bhatt et al., 2007. Post hoc analysis of pt subgroup from a larger randomized trial <u>Size</u> : n=3,096 pts
Inclusion criteria: "CAPRIE-like" if they were enrolled with a documented prior MI, documented prior ischemic stroke, or sx PAD Exclusion criteria:	using reliable contraception Currently receiving investigation drug • Previously entered in other dopidogrel studies Geographic or other factors making study participation impractical Inclusion criteria: Sx (2,838) current IC together with an ABI ≤0.85, or a Hx of IC together with a previous related intervention (amputation, surgical or catheter- based peripheral revascularization) or asx (258) PAD ABI, 0.90 were identified among those with multiple risk factors Exclusion criteria: Taking oral antithrombotic medications or NSAIDs on a long-term basis (although cyclooxygenase-2 inhibitors were permitted). Pts were also excluded if, in the judgment of the investigator, they had established indications for dopidogrel therapy (such as a recent acute coronary syndrome). Pts who were scheduled to undergo a revascularization were not allowed to enroll until the procedure had been completed; such pts were excluded if they were considered to require clopidogrel after revascularization.
Intervention: Clopidogrel + ASA <u>Comparator:</u> Placebo + ASA	Intervention: Clopidogrel + ASA <u>Comparator:</u> Placebo + ASA
<u>1° endpoint</u> : The rate of cardiovascular death, MI, or stroke was significantly lower in the clopidogrel + ASA arm than in the placebo + ASA arm: 7.3% vs. 8.8% (HR 0.83; 95% CI: 0.72–0.96; p=0.01)	<u>1° endpoint</u> : Among the pts with PAD, the primary endpoint occurred in 7.6% in the clopidogrel + ASA group and 8.9% in the placebo + ASA group (HR: 0.85; 95% CI: 0.66– 1.08; p=0.18). In these pts, the rate of MI was lower in the dual antiplatelet arm than the ASA alone arm: 2.3% vs. 3.7% (HR: 0.63; 95% CI: 0.42–0.96; p=0.029), as was the rate of hospitalization for ischemic events: 16.5% vs. 20.1% (HR: 0.81; 95% CI: 0.68–0.95; p=0.011). <u>Safety endpoint</u> : The rates of severe, fatal, or moderate bleeding did not differ between the groups, whereas minor bleeding was increased with clopidogrel: 34.4% vs. 20.8% (OR: 1.99; 95% CI: 1.69–2.34; p<0.001)
<ul> <li>Positive subgroups within negative trials are often the result of confounding or bias, especially post hoc defined subgroups</li> <li>Hospitalizations for</li> </ul>	<ul> <li>Positive subgroups within negative trials are often the result of confounding or bias, especially post-hoc defined subgroups.</li> <li>The rate of the primary safety endpoint (severe bleeding) was 1.7% in each treatment group (p 1/4 0.90).</li> </ul>

	<b>CHARISMA</b> Berger PB, et al. 2010(110) <u>20516378</u>	
	<u>Aim</u> : To determine the frequency and time course of bleeding with DAPT in pts with established vascular disease or risk factors only; identify correlates of bleeding; and determine whether bleeding is associated with mortality. <u>Study type</u> : Post hoc analysis of double- blind, placebo- controlled, randomized trial	(Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization, Management, and Avoidance) trial, where no statistically significant benefit was found in the overall broad population of stable pts studied. <u>Study type</u> : Post hoc analysis of pt subgroup from a larger randomized trial
	<ul> <li>Inclusion criteria: Pts had either established stable vascular disease or multiple risk factors for vascular disease without established disease</li> <li>Exclusion criteria:         <ul> <li>Taking oral antithrombotic medications or NSAIDs on a long- term basis (although cyclooxygenase-2 inhibitors were permitted).</li> <li>In the judgment of the investigator, pts had established indications for clopidogrel therapy (such as a recent acute coronary syndrome).</li> <li>Pts who were scheduled to undergo a revascularization were not allowed to enroll until the</li> </ul> </li> </ul>	<ul> <li>Taking oral antithrombotic medications or NSAIDs on a long- term basis (although cyclooxygenase-2 inhibitors were permitted).</li> <li>In the judgment of the investigator, pts had established indications for clopidogrel therapy (such as a recent acute coronary syndrome).</li> <li>Pts who were scheduled to undergo a revascularization were not allowed to enroll until the procedure had been completed; such pts were excluded if they were considered to require clopidogrel after revascularization.</li> </ul>
	Intervention: Clopdiogrel + ASA <u>Comparator:</u> Placebo + ASA	
	<ul> <li>1° endpoint:</li> <li>Bleeding was assessed with the use of the Global Utilization of Streptokinase and t-PA for Occluded Coronary Arteries (GUSTO) criteria.</li> <li>Severe bleeding occurred in 1.7% of the clopidogrel group vs. 1.3% on placebo (p=0.087); moderate bleeding occurred in 2.1% vs. 1.3%, respectively (p&lt;0.001).</li> <li>Moderate bleeding was strongly associated with increased mortality on multivariable analysis (HR: 2.55; 95% CI: 1.71–3.80; p&lt;0.0001)</li> </ul>	<ul> <li>Safety endpoint:</li> <li>Moderate bleeding was significantly increased: 2.0% vs. 1.3% (HR: 1.60; 95% CI: 1.16–2.20, p=0.004).</li> <li>No significant difference in the rate of severe bleeding: 1.7% vs. 1.5% (HR: 1.12; 95% CI: 0.81–1.53; p=0.50)</li> </ul>
46	• ASA 75 mg to 162 mg	ischemia were significantly decreased in the clopidogrel group, 11.4% vs. 13.2% (HR: 0.86; 95% CI: 0.76–0.96; p=0.008) CI: 0.76–0.96; p=0.008)

$\ensuremath{\textcircled{O}}$ American Heart Association, Inc. and American College of Cardiology Foundation	
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Foundation	

Cassar K, et al. 2005(111) <u>15609386</u>	
<u>Aim</u> : To investigate the antiplatelet effect of a combination of ASA and clopidogrel compared with ASA alone in pts with claudication undergoing endovascular revascularization <u>Study type</u> : Double- blind randomized placebo-controlled <u>Size</u> : n=132 pts	<u>Size</u> : n=15,603 pts
<ul> <li>Inclusion criteria:</li> <li>Pts undergoing lower limb angioplasty</li> <li>Hemoglobin &gt;10 g/L</li> <li>Platelet count &gt;150 × 10⁹ g/L</li> <li>Aspartate aminotransferase, alkaline phosphatase, y-glutamyltransferase &lt;3 times upper normal limit</li> <li>Creatinine &lt;2 times upper normal limit</li> <li>Body mass index &lt;33</li> <li>Age 18–80 y</li> <li>No contraindication to either ASA or clopidogrel</li> <li>Hx of hematological malignancy</li> <li>Acute illness within 14 d of randomization</li> <li>Transfusion of whole blood or red cells within 14 d or randomization Known or suspected drug or alcohol abuse On steroids On warfarin or heparin Hx of bleeding diathesis or coagulopathy Hx of severe neutropenia (neutrophil count &lt;1.8 × 10⁹L) Hx of thrombocytopenia (platelet count &lt;150 × 10⁹L)</li> </ul>	procedure had been completed; such pts were excluded if they were considered to require clopidogrel after revascularization.
Intervention: Clopidogrel 75 mg and ASA 75 mg Placebo and ASA 75 mg	
<ul> <li><u>1° endpoint</u>: Flow cytometric measurements of platelet fibrinogen binding and P-selectin expression were taken as measures of platelet function at baseline, 12 h after the loading dose, platelet activation in the clopidogrel group had decreased (P-selectin by 27.3%, p=0.017; fibrinogen binding by 34.7%, p=0.024; stimulated fibrinogen binding by 34.7%, p=0.024; stimulated fibrinogen binding by 53.9%, 51.7%, and 30 d after endovascular intervention (stimulated fibrinogen binding by 53.9%, 51.7%, and 57.2% respectively; all p&lt;0.001).</li> <li><u>Safety endpoint</u>: 2 pts in each group developed a skin rash and 2 in each group developed a hematoma at the site of radiological access that did not require intervention. The number of pts who developed bruising at and around the site of access was slightly higher in the clopidogrel group between the 2 groups was not statistically significant. 2 pts in the clopidogrel group had an ischemic stroke at d 7 and d 12 after angioplasty. 1 of these pts, however, had stopped taking all medication immediately after intervention. Another pt developed melena secondary to bleeding from multiple small gastric ulcers. Further investigation revealed that the pt had metastatic colonic cancer. 1 pt in the clopidogrel group became hypotensive</li> </ul>	
<ul> <li>Limited to post PTA platelet function</li> </ul>	

	<u>20678878</u>	<b>CASPAR</b> BelchJJ, et al. 2010(112)	
study type: Prospective, multicenter, randomized, double- blind, placebo- controlled <u>Size</u> : n=851 pts	benefit on limb outcomes over ASA alone in pts	<u>Aim</u> : To determine whether clopidogrel + ASA conferred	
<ul> <li>Exclusion criteria:</li> <li>Onset of PAD symptoms before the age of 40 y;</li> <li>Nonatherosclerotic vascular disease;</li> <li>Pts receiving aortobifemoral, iliac-femoral, or crossover (femoral-femoral) grafts, or undergoing peripheral transcutaneous angioplasty during the same surgery;</li> <li>Significant bleeding risk, such as current active bleeding at the surgical site;</li> <li>Withdrawal of an epidural catheter less than 12 hr before randomization;</li> <li>Peptic ulceration within 12 mo of randomization;</li> <li>Previous or current intracranial hemorrhage or hemorrhagic stroke;</li> <li>Any Hx of severe spontaneous bleeding;</li> <li>Current warfarin therapy or anticipated need for warfarin;</li> <li>Concomitant additional antiplatelet agents or thrombolytic agents</li> </ul>	to the trial 2–4 d after bypass surgery. Between 40–80 yr .	Inclusion criteria: Pts undergoing vascular grafting as a treatment for PAD were eligible for recruitment	
Placebo + ASA 75 to 100 mg/d	mg/d Comparator:	Intervention: Clopidogrel 75 mg/d + ASA 75 to 100	
<ul> <li>b) Countred in 143 of 426 pts in the placebo (+ ASA) group vs. 151 of 426 pts in the placebo (+ ASA) group (HR: 0.98; 95% CI: 0.78–1.23). In a prespecified subgroup analysis, the primary endpoint was significantly reduced by clopidogrel in prosthetic graft pts (HR: 0.65; 95% CI: 0.94–1.67; NS). A significant statistical interaction between treatment effect and graft type observed (p=0.008).</li> <li>Safety endpoint: <ul> <li>Although total bleeding (GUSTO)</li> <li>Although total bleeds were more frequent with clopidogrel, there was no significant difference between the rates of severe bleeding in the clopidogrel and placebo (+ ASA) groups (2.1% vs. 1.2%).</li> </ul> </li> </ul>	<ul> <li>In the overall population, the primary endpoint</li> <li>In the overall population, the primary endpoint</li> </ul>	<ul> <li><u>endpoint</u>:</li> <li>Composite of index-graft occlusion or revascularization, above-ankle amputation of</li> </ul>	immediately after intervention and was found to have a retroperitoneal hematoma. This resulted in a delay in discharge from hospital of 7 d but no surgical intervention was necessary
		<ul> <li>Benefit only in prosthetic graft group</li> </ul>	

																																			22569995	2012 (113)	Tepe F, et al.	MIRROR
																							Size: n=80 pts	2	controlled	blind, placebo-	Randomized, double-	Study type:		endovascular therapy	PAD treated with	endpoints in pts with	activation and clinical	local platelet	vs. ASA alone on	antiplatelet therapy	the influence of dual	Aim: To investigate
<ul> <li>Known autoimmune disorders.</li> <li>Known allergy against ASA</li> </ul>	NYHA IV)	limitation (e.g., advanced cancer,	<ul> <li>Other disease with severe life</li> </ul>	retinopathy	<ul> <li>Hypertensive or diabetic</li> </ul>	blood pressure >100 mm hg)	pressure >220 mm hg, diastolic	hypertension (systolic blood	aneurysms Severe uncontrolled	<ul> <li>Arteriovenous malformations or</li> </ul>	<ul> <li>Intracranial neoplasms</li> </ul>	surgery (thrombolysis only)	<ul> <li>Recent (within 2 mo) major</li> </ul>	(thrombolysis only).	or intraspinal surgery or trauma	<ul> <li>Recent (within 2 mo) intracranial</li> </ul>	2 yr (thrombolysis only).	<ul> <li>Cerebrovascular accident within</li> </ul>	platelet count <100,000/mm ³ .	<ul> <li>Hx of bleeding diathesis of</li> </ul>	or serum creatinine >2.5 mg)	disorder (liver cirrhosis, stage B, C	<ul> <li>Known severe hepatic or renal</li> </ul>	genitourinary)	bleeding (e.g. gastrointestinal,	resuscitation, or active internal	<ul> <li>Recent major trauma including</li> </ul>	flow within less than 1 hr.	immediate action and restoration of	threatening ischemia requiring	Exclusion criteria: Acute limb-		Stage Rutherford 3–5	artery and/or popliteal artery)	upper leg (superficial femoral	<ul> <li>Chronic PAD in an artery of the</li> </ul>	<ul> <li>Age &gt;18 y and &lt;90 y.</li> </ul>	Inclusion criteria:
																											replaced by placebo	Clopidogrel	Comparator:		clopidogrel for 6 mo	ASA and 75 mg	dose of 100 mg	followed by a daily	before intervention	mg clopidogrel	mg ASA and 300	Intervention: 500
																							occurred in 1 clopidogrel and 2 placebo pts.	Safety endpoint: Minor bleeding complications		clopidogrel.	needed revascularisation were both resistant to	required TLR (p=0.04). The clopidogrel pts who	were resistant. 2 clopidogrel and 8 placebo pts	(p=0 0.05). 30% of pts who had clopidogrel	concentration of CD40L was 127 and 206.5	clopidogrel and placebo group. The	of β-TG was 224.5 vs. 365.5 (p=0 0.03) in the	<ul> <li>The median peri-interventional concentration</li> </ul>	rate of pt's resistant to clopidogrel	markers β-thromboglobulin and CD40L, and the	<ul> <li>Local concentrations of platelet activation</li> </ul>	<u>1° endpoint:</u>
																																						N/A

Strobl FF, et al. 2013(115) <u>24093324</u>	Bonaca MP, et al. 2013(114) <u>23501976</u>
<u>Aim</u> : Investigating the effects of dual antiplatelet therapy on TLR after balloon angioplasty ± stenting in the femoropopliteal segment <u>Study type:</u> Prospective, randomized, single- center, double- blinded and placebo-	<u>Aim</u> : The effect of vorapaxar on cardiovascular and peripheral vascular outcomes in pts who qualified for TRA2°P- TIMI 50 with sx PAD. <u>Study type:</u> Randomized, double- blind, placebo- controlled trial
Inclusion criteria: PAD pts with TLR after femoropopliteal endovascular intervention Exclusion criteria: N/A	<ul> <li>and/or clopidogrel.</li> <li>Childbearing potential or existing pregnancy.</li> <li>Contraindications to urokinase, reteplase, clopidogrel, heparin and acetylsalicylic acid.</li> <li>Pt who has previously been included in this trial.</li> <li>Pt who requires long-term Cox2 inhibition.</li> <li>Pt who is not able to sign the informed consent form</li> <li>Inclusion criteria: Hx of IC in conjunction with an ABI &lt;0.85 or previous revascularization for limb ischemia</li> <li>Exclusion criteria:</li> <li>A planned revascularization that had not yet been performed;</li> <li>Hx of a bleeding diathesis</li> <li>Were receiving vitamin K antagonist therapy</li> <li>Had active hepatobiliary disease</li> </ul>
Intervention: ASA and clopidogrel <u>Comparator</u> : ASA	Intervention: Vorapaxar <u>Comparator:</u> Placebo
<ol> <li>endpoint: At 6 mo, clopidogrel pts had significantly lower rates of TLR compared to placebo pts [2 (5%) vs. 8 (20%); p=0.04]. After stopping clopidogrel/placebo after 6 mo, there was no significant difference in TLR at 12 mo after treatment [9 (25%) clopidogrel vs. 12 (32.4%) placebo; p=0.35]. Mortality was 0 vs. 1 in the placebo group at 6 mo (p=0.32) and 0 vs. 3 at 12 mo (p=0.08).</li> </ol>	1° endpoint:       Primary efficacy endpoint was cardiovascular death, MI, or stroke. The primary endpoint did not differ significantly with vorapaxar (11.3% vs. 11.9%; HR: 0.94; 95% CI: 0.78–1.14; p=0.53)         Safety endpoint:       Principal safety endpoint was Global Utilization of Streptokinase and t-PA for Occluded Coronary Arteries (GUSTO) bleeding. Bleeding occurred more frequently with vorapaxar compared with placebo (7.4% vs. 4.5%; HR: 1.62; 95% CI: 1.21–2.18; p=0.001).
NA	<ul> <li>Rates of hospitalization for ALI (2.3% vs. 3.9%; HR: 0.58; 95% CI: 0.39–0.86; p=0.006) and peripheral artery revascularization (18.4% vs. 22.2%; HR: 0.84; 95% CI: 0.73–0.97; p=0.017) were significantly lower in pts randomized to vorapaxar.</li> </ul>

Antiplatelet Trialists 2002(117) <u>11786451</u>	Antiplatelet Trialists Collaboration (graft arterial patency) <u>8312766</u>
<u>Aim</u> : To determine the effects of antiplatelet therapy among tps at high risk of occlusive vascular events. <u>Study type</u> : Meta-	size: n=73 pts <u>Aim</u> : To determine the efficacy of antiplatelet therapy in maintaining vascular patency in various categories of pts. <u>Study type</u> : Overviews of 46 RCTs of antiplatelet therapy vs. control and 14 RCTs comparing one antiplatelet regimen with another. <u>Size</u> : n=12,000 pts
Inclusion criteria: PAD includes those with claudication and/or peripheral revascularization Exclusion criteria: N/A	Inclusion criteria: Pts at varying degrees of risk of vascular occlusion (by virtue of disease or of having some vascular procedure) were in trials of antiplatelet therapy vs. control or trials comparing different antiplatelet regimens Exclusion criteria: 39 trials of antiplatelet therapy vs. control were identified among pts having peripheral vascular procedures or with PVD (see part I) but vascular occlusion was monitored systematically in only 14 of them
Intervention: Antiplatelet therapy <u>Comparator:</u> Control	Intervention: Antiplatelt therapy <u>Comparator</u> : No antiplatelet therapy
<b>1° endpoint:</b> Allocation to antiplatelet therapy reduced the combined outcome of any serious vascular event by about one quarter; nonfatal MI was reduced by one third, nonfatal stroke by one quarter, and vascular mortality by one sixth (with no apparent adverse effect on other deaths)	<ul> <li><u>1° endpoint</u>: Antiplatelet therapy produced a highly significant (2p &lt;0.0001) reduction in vascular occlusion, with similar proportional reductions in several different types of pts As well as preventing subclinical occlusion, antiplatelet therapy produced a significant (2p=0.002) reduction of about one quarter in the odds of suffering a "vascular event" (nonfatal MI, nonfatal stroke, or vascular death).</li> <li><u>Safety endpoint</u>: No clear excess bleeding</li> </ul>
<ul> <li>Among 9,214 pts with PAD in 42 trials (compared with 4,939 such pts in 33 trials previously evaluated there was a proportional reduction of 23% (8%) in serious vascular events (p=0.004), with similar</li> </ul>	<ul> <li>Allocation to antiplatelet therapy in the 14 trials with pts with PAD was associated with a proportional reduction of 43% (SD 8%) in vascular occlusion, which was highly significant. Studies of pts with saphenous vein grafts or prosthetic implants for lower limb disease contributed most of the data; of the 3 other studies, 1 assessed the patency of native vessels in pts with IC and 2 concerned pts who had had peripheral angioplasty.</li> <li>allocation to a mean scheduled duration of 19 mo of antiplatelet therapy produced a substantial absolute reduction of 92 (SD 15) per 1,000 in the risk of peripheral artery occlusion (15.7% of antiplatelet allocated pts vs. 24.9% of corresponding controls</li> </ul>

Bonaca MP, et <u>Aim:</u> al. effect 2013 on CV <u>23501976</u> <u>Stud</u> <u>Size:</u>	Morrow DA, et <u>Aim</u> : D al. 2012(118) on sec <u>22443427</u> atherot events <u>Study</u> <u>Size</u> : n	analysis o antiplatele for preven death, MI, in high risi <u>Size:</u> n=2 involving in compar antiplatele vs. contro 77,000 in comparise different a regimens
<u>Aim</u> : Determine the effect of vorapaxar on CV and peripheral vascular outcomes <u>Study type</u> : RCT <u>Size</u> : n=26,449 pts	<u>Aim</u> : Determine the impact of vorapaxar on secondry prevention of atherothrombotic events <u>Study type</u> : RCT <u>Size</u> : n=26,449 pts	analysis of RCTs of antiplatelet therapy for prevention of death, MI, and stroke in high risk pts <u>Size</u> : n=287 studies involving 135,000 pts in comparisons of antiplatelet therapy vs. control and 77,000 in comparisons of different antiplatelet different antiplatelet
Inclusion criteria: Pts who qualified for TRA 2°P-TIMI 50 pts with a with stable atherosclerotic vascular disease and a prior MI, ischemic stroke, or PAD Exclusion criteria: N/A	Inclusion criteria: Pts who had a hx of MI, ischemic stroke, or PAD Exclusion criteria: Pts were ineligible if they were planning to undergo a revascularization procedure, had a hx of bleeding diathesis, had recent active abnormal bleeding, were receiving ongoing treatment with warfarin, or had active hepatobiliary disease.	
Intervention: Vorapaxar. Thienopyridine was planned at randomization in 12,410 pts <u>Comparator</u> : Placebo	Intervention: Vorapaxar <u>Comparator</u> : Placebo	
<u>1° endpoint</u> : CV death, MI, or stroke <u>Safety endpoint</u> : Global Utilization of Streptokinase and t-PA for Occluded Coronary Arteries bleeding.	<u>1° endpoint</u> : Composite of death from cardiovascular causes, MI, or stroke in 1,028 pts (9.3%) in the vorapaxar group and in 1,176 pts (10.5%) in the placebo group (HR for the vorapaxar group: 0.87; 95% CI: 0.80–0.94; p<0.001). <u>Safety endpoint</u> : There was an increase in the rate of intracranial hemorrhage in the vorapaxar group (1.0%, vs. 0.5% in the placebo group; P<0.001).	Safety endpoint: The proportional increase in risk of a major extracranial bleed with antiplatelet therapy was about one half (OR: 1.6; 95% CI: 1.4–1.8), with no significant difference between the proportional increases observed in each of the 5 high risk categories of pts
<ul> <li>In the PAD Cohort:</li> <li>No significant difference between vorapaxar and comparator for CV death, MI, or stroke (11.3% vs. 11.9%; HR: 0.94; 95% CI: 0.78–1.14; p=0.53)</li> <li>Significantly lower rates of hospitalization for ALI for vorapaxar group (2.3% vs. 3.9%; HR: 0.58; 95% CI: 0.39–0.86; p=0.006)</li> <li>Significant increase in bleeding in vorapaxar group</li> </ul>	•3,787 PAD pts	benefits among pts with IC, those having peripheral grafting, and those having peripheral angioplasty • Much of the data was from the picotamide trial

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				al. 2016(120)	Bonaca MP, et	Bohula EA, et al 2015(119) <u>26338971</u>	
<b>Size:</b> n=3,787 pts	<u>Study type:</u> Subgroup of a randomized trial	and by etiology.	PAD and whether	the causes, sequelae and predictors of ALI	Aim: Evaluate	<ol> <li><u>Aim</u>: To determine whether the efficacy and safety of antiplatelet therapy with vorapaxar was modified by concurrent thienopyridine use.</li> <li><u>Study type:</u> Randomized, double- blind, placebo- controlled trial</li> <li><u>Size</u>: n=16,897 pts</li> </ol>	
			absence of PAD	50 pts with PAD	Inclusion criteria: TRA 2°P-TIMI	Inclusion criteria: TRA 2°P-TIMI 50 pts who qualified with a MI in the preceding 2 weeks to 12 months and was restricted to. Exclusion criteria: Pts without a hx of stroke or transient ischemic attack given its contraindication in that population	
			Placebo	Vorapaxar	Intervention:	Intervention: Vorapaxar. Thienopyridine was planned at randomization in 12,410 pts <u>Comparator</u> : Placebo	
		Safety endpoint: Bleeding (see TRA 2°P-TIMI 50)	as total ALT events by 41% (94 events vs. 30 events, risk ratio: 0.59; 95% CI: 0.38– 0.93,p=0.022)	Vorapaxar reduced first ALI events by 41% (HR: 0.58; 95%CI: 0.39–0.86; p=0.006), as well	1° endpoint: ALII	<ul> <li><u>1° endpoint</u>: Vorapaxar significantly reduced the composite of cardiovascular death, MI, and stroke in comparison with placebo regardless of planned thienopyridine therapy (planned thienopyridine, HR: 0.80; 95% Cl: 0.70–0.91; p&lt;0.001; no planned thienopyridine, HR: 0.75; 95% Cl: 0.60–0.94; p=0.011; p- interaction=0.67).</li> <li><u>Safety endpoint</u>: Consistent with the findings in the overall cohort, these rates reveal an increased RR of GUSTO moderate to severe bleeding in pts treated with vorapaxar in comparison with placebo; however, there was no significant modification by planned thienopyridine use (planned thienopyridine HR: 1.50; 95% Cl: 1.18–1.89, p&lt;0.001; no planned thienopyridine HR: 1.90; 95% Cl: 1.17–3.07; p=0.009; p-interaction=0.37</li> </ul>	
			Enect consistent across     all etiologies	graft thrombosis or in situ native vessel thrombosis	<ul> <li>Most ALI events were</li> </ul>	N/A	compared with placebo (.4% vs. 4.5%; HR: 1.63; 95% Cl: 1.21–2.18; p=0.001).

	<ul> <li>0.89; p&lt;0.001).</li> <li>3 y KM event rate of CV death, MI, stroke, or</li> </ul>		either an ABI <0.85 or previous revascularization for limb ischemia	TIMI 50 trial	
nx of stroke).	was 7.9% in vorapaxar compared with 9.5% in placebo (HR: 0.80; 95% CI: 0.73–	Placebo	<ul> <li>Those with symptomatic PAD had hx of IC in conjunction with</li> </ul>	blinded, placebo-	
population (not those with a	• 3 y KM event rate of CV death, MI, or stroke		2 wk to 12 mo	Study type:	
and FDA approved	to urgent coronary revascularization	2.08 mg) daily	Hx of spontaneous MI within prior		25792124
stroke in the intended use	death, MI, stroke, or recurrent ischemia leading	2.5 mg (vorapaxar	criteria	vorapaxar	2015(123)
reduce CV death, MI, or	death, MI, or stroke, and CV	Vorapaxar sulfate	Met TRA 2°P-TIMI 50 inclusion	safety and efficacy of	a
<ul> <li>Vorapaxar was shown to</li> </ul>	1° endpoint: Composite endpoints of CV	Intervention:	Inclusion criteria:	Aim: To observe the	Magnani G, et
	Clopidogrei plus ASA (RR: 1.48; 95%Cri: 1.05- 2.10; NNT=215)				
	(RR: 1.80; 95%Crl: 1.22–2.69; NNH=130), and				
	5.03; 95%Crl: 1.23–39.6; NNH=25), Vorapaxar				
	was significantly higher with Ticlopidine (RR:				
	Safety endpoint: The risk of severe bleeding				
	95%Cri: 0.46–0.99 compared to ASA, NN I≡94)			<b>Size:</b> n=34 518 pts	
	following leg revascularization (RR: 0.68;			analysis	
	aspirin significantly reduced major amputations			Study type: Meta-	
	Dual antiolatelet therany with Clonidogral plus				
	0.61–0.99; NNT=98).			Amputations in pts	
	Clopidogrel plus aspirin (RR: 0.78; 95%Crl:			Events and Leg	
	0.75; 95%Crl: 0.58–0.96; NN =87), and			Major Cardiovascular	
	95%Crl: 0.58–0.91; NNT=80), Ticlopidine (RR:	Placebo		for Prevention of	
	0.46-0.96: NNT=66). Clonidoarel (RR: 0.72)	Comparator:	Exclusion criteria: N/A	Antiplatelet Agents	26274912
	Ticarrelor plus aspirin /PP: 0.67: 05%/0rl:			of Different	2015 (122)
N/A	1° endpoint: MACE and leg amputations	<u>Intervention</u> : Antinlatelet therany	antinlatelet drugs in the with PAD	AIM: Comparative	Aatsanos K, et
		•	- - - - -	Size: n=936 pts	
				randomized trial	
	similar magnitude of risk.			Subgroup of large	
	bleeding in both pts with and without PAD at a			Study type:	
	Safety endnoint: Voranaxar increased				
	statistical significance."	Placebo	without PAD	documented PAD	2014(121) 25262270
	amputation with vorapaxar did not reach			safety of vorapaxar in	Jones WS, et al.
	points, peripheral revascularization, and	Vorapxar	with a hx of PAD	the efficacy and	TRACER
N/A	1° endpoint: Lower rates of ischemic end	Intervention:	Inclusion criteria: TRACER pts	Aim: Investigate	PAD from

ran; Connormany, construction nce interval; CLI, critical limb ; IC, intermittent treat; NS, not significant; uminal angioplasty; pt,	Size: n=18 trials,       minitian of the second secon	<ul> <li>ischemia; ASA, aspirin;</li> <li>ischemia; ASA, aspirin;</li> <li>of Streptokinase and t-P</li> <li>ent; MI, myocardial infarc</li> <li>ent; MI, myocardial, infarc</li> <li>s; OR, odds ratio; PAD, p</li> <li>s; OR, odds ratio; PAD, p</li> <li>ative risk; and TLR, tarc</li> </ul>	Size:         n=18 trials,           5,269 pts         5,269 pts           ABI indicates ankle-brachial index; ACS, acute coronary syndrome; ALI, acute limb ischemia; ASA, aspirin; CHD, coronary heart disea ischemia; CV, cardiovascular; GP, general practitioner; GUSTO, Global Utilization of Streptokinase and t-PA for Occluded; Coronary A claudication; IV, intravenous; KM, Kaplan-Meier; MACE, major adverse cardiac event; MI, myocardial infarction; N/A, not applicable; N NYHA, New York Heart Association; NSAIDs, nonsteroidal anti-inflammatory drugs; OR, odds ratio; PAD, peripheral artery disease; P patient; PVD, peripheral vascular disease; RCT, randomized controlled trial; RR, relative risk; and TLR, target lesion revascularization.	Size: n=18 trials, 5,269 pts -brachial index; ACS, acu iovascular; GP, general p ravenous; KM, Kaplan-M leart Association; NSAID, heral vascular disease; R	ABI indicates ankle ischemia; CV, card claudication; IV, int NYHA, New York H patient; PVD, perip
<ul> <li>events</li> <li>ASA did have significant reduction in nonfatal stroke</li> <li>No significant outcome for MI, CV mortality, or all-</li> </ul>	Safety endpoint: Major bleeding		Data on all-cause mortality, CV death, MI, stroke, and major bleeding     Exclusion criteria: N/A	<u>Study type</u> : Meta- analysis of prospective RCTs	
<ul> <li>ASA therapy, alone or in combination with dipyridomole, had no significant effect on CV</li> </ul>	<ul> <li><u>1° endpoint:</u></li> <li>Nonfatal MI, nonfatal stroke, CV death</li> <li>Secondary outcomes were all-cause mortality</li> </ul>	Intervention: ASA <u>Comparator:</u> Placebo/control	<ul> <li>Inclusion criteria:</li> <li>Prospective RCTs</li> <li>PAD pts assigned to aspirin or placebo/control group</li> </ul>	<u>Aim</u> : To determine the effect of ASA on CV event rates in pts with PAD	Berger JS et al, 2009 (124)
	<ul> <li>Combined bleeding criteria was 3.7% with vorapaxar and 2.4% in placebo (HR, 1.55; 95% Cl: 1.30–1.86, p&lt;0.001).</li> <li>Severe bleeding was 1.3% with vorapaxar vs. 1.0% with placebo (HR 1.24; 95% Cl: 0.92–1.66, P=0.16</li> </ul>				
	<ul> <li>7.2% in vorapaxar and 8.3% in placebo; HR: 0.83; 95% Cl: 0.75–0.93, p&lt;0.001).</li> <li>3 y KM event rate of MI was 5.4% in vorapaxar and 6.4% in placebo (p&lt;0.001)</li> <li>3 y KM event rate of stroke was 1.2% in vorapaxar and 1.6% in placebo (p=0.002) individually.</li> </ul>				
	urgent coronary revascularization was 10.1% in vorapaxar and 11.8% in placebo (HR: 0.83; 95% CI: 0.76–0.90; p<0.001). • 3 y KM event rate of CV death or MI was		Exclusion criteria: N/A	<u>Size</u> : n=16,897 pts	

ALI indicates acute										<u>25864042</u>	2015(125)	et al.	Armstrong EJ	Year	Author	Acronym	Study	Evidence Table
imb ischemia; ASA, ace	Size: n=629 pts	Observational cohort	Study type:	disease.	peripheral arterial	among pts with sx	with ASA monotherapy	clopidogrel compared	DAPT with ASA and	additive benefit of	whether there is	conducted to determine	Aim: This study was		Study Size (N)	Study Type;	Aim of Study;	e 14. Nonrandomize
ylsalicylic acid; Cl, confidenc	stenosis	stenosis, or renal artery	carotid artery stenosis,	<ul> <li>In registry for ALI,</li> </ul>	(28)	No antiplatelet therapy	• Warfarin use (96 nts)	Exclusion criteria:		<ul> <li>All had angiography</li> </ul>	<ul> <li>Claudication or CLI</li> </ul>	<ul> <li>UC Davis PAD registry</li> </ul>	Inclusion criteria:				Patient Population	d Trials, Observational
ce interval; CLI, critical limb i								median follow 3.2 y	Record review with		only	DAPT, 281 with ASA	Groups: 348 with	(include # patients)	Study Comparator	(include # patients) /	Study Intervention	Studies, and/or Regis
ALI indicates acute limb ischemia; ASA, acetylsalicylic acid; CI, confidence interval; CLI, critical limb ischemia; DAPT, dual antiplatelet therapy; HR, hazard ratio; MACE, major adverse		ASA or clopidogrel and adverse events at 1 y.	55% had a decreased response to clopidogrel. No association was found between a reduced response to	function testing, 21% had decreased response to ASA and	subgroup of 94 pts who underwent point-of-care platelet	amputation (adjusted HR: 0.69; 95% Cl: 0.37-1.29). In a	was found between DAPT use and the risk of major	(adjusted HR: 0.55; 95% CI: 0.35–0.89). No association	HR: 0.65; 95% CI: 0.44–0.96) and overall mortality	was associated with a decreased risk of MACEs (adjusted	monotherapy group. After propensity weighting, DAPT use	occurred in the DAPT group vs. 59 (29%) in the ASA	1° endpoint: During 3 y of follow-up, 50 events (20%)		95% CI)	(include Absolute Event Rates, P value; OR or RR; and	Endpoint Results	Evidence Table 14. Nonrandomized Trials, Observational Studies, and/or Registries of Antiplatelet Agents–Section 5.2.
1ACE, major adverse													N/A	Adverse Events	Study Limitations;	Endpoint (if any);	Relevant 2°	

cardiac event; PAD, peripheral artery disease; and pt, patient. Ĵ Ē 3 чру, і . -<u>ר</u> שייי

Evidence Table 15. R
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	I J. Naliuvilizeu II	Evidence rable 13. Nandonnized mais comparing statim Agents-Section 3.2.			
Study Acronym	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2 [°] Endpoint (if any);
Author	Study Type;		(include # patients) /	(include Absolute Event	Study Limitations;
Year	Study Size (N)		Study Comparator	Rates, P value; OR or RR; and	Adverse Events
			(include # patients)	95% CI)	
HPS	Aim: Assess impact of	Inclusion criteria:	Intervention:	1° endpoint: 24% (95% CI: 19–	<ul> <li>Comparable proportional reduction in first</li> </ul>
HPS	cholesterol-lowering	• Age 40–80 y	Simvastatin 40 mg	28; p<0.0001) proportional	major coronary event, stroke, and
Collaborative	therapy on major	<ul> <li>Chol &gt;135mg/dL</li> </ul>	(10,269)	reduction in the first occurrence	revascularization (considered separately)
Group	adverse vascular	<ul> <li>PAD, CVD, DM, or HTN (if</li> </ul>		of a major vascular event	<ul> <li>16% reduction in peripheral vascular</li> </ul>
2007(126)	events in pts with PAD	male and >65)	Comparator: Placebo	Those with LEPAD: 22% (95%	events (5%–25%; p=0.006), primarily
<u>17398372</u>			(10,267)	Cl: 15–29; p<0.0001)	through reduction in noncoronary
	Study type:	Exclusion criteria: If PCP		proportional reduction	revascularizations
	Prospective, blinded,	feels statin clearly indicated or			<ul> <li>Statin group: 85% compliant with statin</li> </ul>
	RCT.	contraindicated; prior MI,		1° Safety endpoint (if	<ul> <li>Non-statin group: 17% non-study statin</li> </ul>
		stroke, or admission with		<u>relevant)</u> :	
	Size: n=20,536 pts	angina in previous 6 mo; liver		<ul> <li>CPK elevation &gt;10x ULN in 1</li> </ul>	
		dysfunction; renal dysfunction;		out of 10,000 pts/y.	

	Safety endpoint: 2/3 of pts in each treatment group reported drug-related adverse event (pruritis, diarrhea, elevated blood sugar). Flushing in 54%. Serious adverse events were		Exclusion criteria: Pts with CAD or other indication for lipid lowering therapy.	<u>Size</u> : n=387	
	<ul> <li>High Niacin/Lova: 37.8%; 95%</li> <li>CI: 26.6%–50.1%, p=0.137</li> </ul>		• <20% variability in 2 assessments.	Study type: RCT	
	L• ow Niacin/Lova: 38.6%; 95% CI: 27.6%–50.6%, p=0.096	Comparator: Diet	exercise (sarurier) ● LDL ≤160 ● PWT 1–20 min	with diet intervention.	
	<ul> <li>20 wk</li> <li>Diet: 26.5%; 95% Cl: 16.4%–</li> <li>27.6%</li> </ul>	2000 mg plus	ABI SU.90     20% reduction in ABI post-	exercise performance	<u>1021207</u> 0
Composite of CV events	and in claudication onset time at	plus lovastatin 40 mg	• Stable IC	Niacin plus lovastatin	2010(128)
Change in ABI     Walking Impairment Questionnaire	<ul> <li><u>1° endpoint:</u></li> <li>Change from baseline in PWT</li> </ul>	dose Niacin 1000 mg	<ul> <li>Age &gt;40 y</li> </ul>	Aim: lest the hypothesis that ER	Hiatt WR, et al.
			exercise rehab program.		
			DVI/PE within 3 mo.		
			Stroke or TIA within 6 mo.		
			<ul> <li>USA within 3 mo.</li> </ul>		
			revascularization within 6 mo.	Size: n=354 pts	
			<ul> <li>IVII, coronary</li> <li>revascularization nerinheral</li> </ul>		
			Exclusion criteria:	Prospective, blinded, RCT	
		(114 pts)		Study type:	
		Comparator: Placebo	● LDL ≤160.		
	<ul> <li>Atorva 80: 90±18</li> <li>(p=0.37)</li> </ul>	(120 pts)	<ul> <li>20% reduction in ABI post exercise (Gardner)</li> </ul>	in pts with IC	
• Atorva 80: 81±15 (p=0.025)	<ul> <li>Atorva 10: 90±18</li> </ul>	atorvastatin 80 g daily	• ABI ≤0.90	atorvastatin improves	
• Atorva 10: 74±14 (p=0.13)	<ul> <li>Placebo: 50±12 s</li> </ul>	daily (120 pts) or	Stable IC for 6 mo	lowering with	12952839
	12 mo.	Atorvastatin 10 mg	• Age >25 y	whether cholesterol	2003(127)
		lutan satis			
			Cluck limitations to		
			Rx (cyclosporine, fibrates,		
	<ul> <li>Mean follow-up 5.0 y</li> </ul>		muscle disease; concurrent		

Stoekenbroek <u>Ai</u> RM, et al.         wh           2015(131)         state           25595417         ind           ind         C/	West AM, et al. <u>2011(130)</u> <u>21570685</u> ez th re att th St R( Si	Giri J, et al. 2006(129) <u>16516084</u> ar Fr U Iat Iat St St St St	
<u>Aim</u> : Determine whether high-dose statin vs. usual dose statin reduces incidence of PAD and CAD outcomes in pts	<u>Aim</u> : LDL-C cholesterol by adding ezetimibe to statin therapy would regress atherosclerosis measured by MRI in the SFA in PAD. <u>Study type</u> : Single center, prospective, RCT, double-blinded <u>Size</u> : n=87 pts	<u>Aim</u> : To determine whether statin use is associated with less annual decline in LE functioning with/without LEPAD <u>Study type:</u> Prospective cohort study (identified in noninvasive vascular lab between 1998- 2000 at 3 Chicago institutions). <u>Size</u> : n=544	
<ul> <li>Inclusion criteria:</li> <li>Age ≤80 y</li> <li>Confirmed prior MI</li> <li>Exclusion criteria: N/A</li> </ul>	Inclusion criteria: 30–85 y, PAD (ABI 0.4–0.9) Exclusion criteria: Rest pain, CLI, contraindication to MRI, pregnancy.	Inclusion criteria: • PAD group: ABI <0.90. • Non-PAD: 1.50 ≥ABI ≥0.90 Exclusion criteria: • SNF resident • Wheelchair bound • Foot or leg amputation • Non-English speaking • Recent major surgery • Prior vasc surgery • Normal ABI	
Intervention: Atorvastatin 80mg <u>Comparator:</u> Simvastatin 20–40mg	Intervention: Statin- naive (randomized to simvastatin or simvastatin plus ezetimibe) or previously on statin given open label ezetimibe Comparator: Simvastatin alone	Intervention: On statin <u>Comparator:</u> Not on statin	
<ul> <li>1° endpoint:</li> <li>No PAD at baseline: new clinical Dx of PAD requiring diagnostic procedures or interventions.</li> <li>2.2% in atorvastatin</li> </ul>	1° endpoint:• Atherosclerotic plaque volumein the proximal 15–20 cm of SFAat baseline and annually x 2.• Baseline and y 2 volumes:• S + E (11.5 $\pm$ 1.4 vs.10.5 $\pm$ 1.3 cm ³ ; p=NS) or• S (11.0 $\pm$ 1.5 vs.10.5 $\pm$ 1.4 cm ³ , p=NS)• E (10.0 $\pm$ 0.8–10.8 $\pm$ 0.9; p<0.01)	<ul> <li>1^o endpoint:</li> <li>Pts with PAD using statins had less annual decline in:</li> <li>Usual-pace walking velocity (0.002 vs0.024 m/s/y; p=0.013)</li> <li>Rapid-pace walking velocity (-0.006 vs0.042 m/s/y; p=0.006)</li> <li>6 min walk performance (-34.5 vs57.9 ft/y; p=0.088)</li> <li>Summary performance score (-0.152 vs0.376; p=0.067) compared with non-users.</li> <li>Among pts without-PAD, there were no significant associations between statin use and functional decline.</li> </ul>	similar in all 3 groups (11.2%, 11.2%. 10.3%)
<ul> <li>Post-hoc evaluation of CAD outcomes in pts with PAD at baseline</li> <li>Baseline PAD in 374 pts (4.2%)</li> <li>Major coronary events nonsignificantly lower in the atorvastatin group (14.4%) compared with the simvastatin group</li> </ul>	<ul> <li>Only 72 pts at follow-up (2 died, 11 lost to follow-up, 2 withdrew prior to baseline imaging)</li> <li>Statin initiation with or without ezetimibe in statin-naive pts halted plaque progression</li> <li>Ezetimibe added to existing statin still resulted in progression of plaque volume; ezetimibe's effect on PAD may depend on relative timing of statin therapy.</li> <li>LDL-C was lowered by the addition of ezetimibe in both groups, but did not translate to change in plaque volume. Study was underpowered to detect a difference between S and S + E</li> </ul>	N/A	

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Feringa HH, et al. 2007(136) <u>17360142</u>	Westin GG, et al. 2014(135) <u>24315911</u>	Vogel TR, et al. 2013(134) <u>24300135</u>
<u>Aim</u> : To determine whether higher-dose LDL are independently associated with better outcomes in PAD <u>Study type</u> : Single center, prospective, observational, cohort study <u>Size</u> : n=1,374 pts	<u>Aim</u> : To determine the associations between statin use and MACCE and amputation-free survival in CLI pts. <u>Study type</u> : Single center registry (retrospective cohort) <u>Size</u> : n=380 (between 2006–2012)	<u>Aim</u> : To evaluate preoperative administration of statins and longitudinal limb salvage after LE endovascularization and LE open surgery. <u>Study type</u> : Medicare Claims Database Review Size: n=22,954
Inclusion criteria: • Age ≥18 • ABI ≤0.90 Exclusion criteria: • MI or coronary revascularization in past 6 mo • Liver disease (Cirrhosis or hepatitis)	Inclusion criteria: ≥1 presentation with CLI (Rutherford 4–6). "On statin" if hospitalization data or most recent pre- procedure clinic note had statin listed (65% of pts enrolled) Exclusion criteria: N/A	Inclusion criteria: Age ≥65 y with a diagnosis of atherosclerosis of LE arteries who were hospitalized during 2007–2008 for LE revascularization Exclusion criteria: N/A
Intervention: Statin therapy (propensity analysis applied to control for confounders)	<u>Intervention</u> : On statin (246 or 65%) <u>Comparator</u> : No statin	Intervention: On statin at time of revascularization (11,687) <u>Comparator</u> : No statin
<ul> <li><u>1° endpoint</u>: All-cause mortality and cardiac death</li> <li><u>Results</u>:</li> <li>6 mo LDL:</li> <li>&lt;100 in 30.8%</li> <li>&lt;70 in 9.7%</li> <li>Lowest all-cause and cardiac mortality (18% and 13%) in pts with lowest cholesterol (&lt;70), p&lt;0.001; gradually increasing with increasing cholesterol levels</li> </ul>	<u>1° endpoint</u> : Composite MACCE (death, MI, stroke) within 1 y of procedure. <u>Results:</u> Statin: 18%, no statin: 23% (HR: 0.53; 95% CI: 0.28– 0.99; p=0.048) Propensity score to control for confounding variables	<u>1° endpoint</u> : 1 y limb salvage rates Statin: RR=0.82; 95% CI: 0.78– 0.86; p<0.0001
Secondary endpoint: progression to kidney failure Conclude: pts with ABI <0.90 benefit from LDL <70 Mean follow-up 6 y	<ul> <li>Secondary outcomes (1 y): death, MI, stroke, ipsilateral LE bypass, ipsilateral major amputation, amputation-free survival, vessel patency (primary, primary assisted, secondary)</li> <li>Amputation-free survival HR: 0.59; 95% CI: 0.35–0.98; p=0.04</li> <li>Improved vessel patency</li> <li>Pts on statin had higher rates of DM, HTN, CAD, CVD, prior MI</li> </ul>	NA

	KC IS for Antinype	Evidence Table 17. RCTs for Antihypertensive Agents- Section 5.3.	•	- - -	
Study Acronym; Author; Year Published	Study Type; Study Size (N)		Study intervention (# patients) / Study Comparator (# patients)	(Absolute Event Rates, P value; OR or RR; & 95% Cl)	Kelevant 2° Encipoint (it any); Study Limitations; Adverse Events
HOPE Study ABI subgroup	Aim: Impact of ramipril on CVD events	Inclusion criteria: Age ≥55 y with CVD (CAD, stroke, PAD) or DM+RF	Intervention: Ramipril vs. placebo	<u>1° endpoint</u> : ● MACE	N/A
Ostergren J, et al. 2004(137) 14683738	Study type: RCT	HE or IV divefinition (EE on A)	PAD group (N=1996 ramipril vs. N=2085 placebo)	• Asx PAD: ABI 0.6–0.9 15.7 vs. 21.6 0.72 (0.56,	
	Size: n=9,297 pts overall, 4,051 with			<0.52/ <0.6 16.4 vs. 22.0 0.77 (0.55. 1.09)	
	PAD			• Clinical PAD 20.1 vs. 25.8	
	8,986 pts with ABI measured. 3,099 pts			0.75 (0.61, 0.92)	
HOPE	Aim: To investigate	Inclusion criteria: Pts ≥55 y with	Intervention: Ramipril	1° endpoint: Composite of	Death from cardiac causes
Yusuf S, et al. 2000/138\	effect of ACEI (Raminril-10mm) on	hx of CAD, stroke, PVD or DM with	(10mg) (4,645)	MI, stroke, or mortality from	reduced (6.1% vs. 8.1%; p<0.001)
<u>10639539</u>	CV events in high risk	cholesterol, low LDL, smoking, or	Comparator: Placebo		12.3%; p<0.001)
	pts ≥55 y with a mean	micro albuminuria.	(4,652)	Results: Endpoint	• Death from any cause (10.4 %
	mmHg in both groups	Exclusion criteria:		vs. Placebo (14% vs.	Ramipril was found to be
	Study type: RCT, 2x2	• <0.40 EF		0.86; p<0.001)	
	factorial design	On ACE-I or Vitamin E			
	<u>Size</u> : n=9,297 pts	<ul> <li>Uncontrolled hypertension or overt nephropathy</li> <li>Had MI or stroke&lt;4 wk</li> </ul>			
	Aim: Impact of	Inclusion criteria:	Intervention: Telmisartan	1° endpoint:	<ul> <li>Increased risk of hypotension,</li> </ul>
2008(139)	vs. combination on	cerebrovascular disease. PAD) or	combo	● Overall trial 16 5% in	combination group
18378520	CVD events in pts with	DM+end-organ damage	PAD group (N=1136	Ramipril, 16.7%	
	vascular disease or		ramipril vs. N=1161	telmisartan, 16.3%	
	nign-risk Divi	Exclusion criteria:	combo)	combination group.	

Table 17 DCTe fo vnertensive Agents- Section 5.3. CAD indicates coronary artery disease; CLI, critical limb ischemia; CVD, cardiovascular disease; CV, cardiovascular; DM, diabetes mellitus; HR, hazard ratio; HTN, hypertension; LDL, low-density lipoprotein; LE, lower extremity; MACCE, major adverse cardiovascular and cerebrovascular events; MI, myocardial infarction; N/A, not applicable; pt, patient; and RR, relative risk.

				Size: n=Parallel in 177	
			Inability to exercise Poorly controlled DM	Study type: RCT	
			Exclusion criteria:		
No adverse effects BB	Increase 28% vs. 26%.			capacity in IC	21602713
No difference in ABI change     hetween groups	<u>1º endpoint:</u> Initial	Intervention: Nebivolol 5	Inclusion criteria: PAD with IC	Aim: Nebivolol vs.	Diehm C, et al. 2011/142)
			Requiring BB use	toilow-up 4.∠ y.	
			●ĦF	procedure. Mean	
			<ul> <li>Severe liver or kidney disease</li> </ul>		
			Valvular heart disease	nospitalization, MI,	
			stroke within 3 mo		
			<ul> <li>Coronary revascularization or</li> </ul>	CVD events: cardiac	
			<ul> <li>Renal artery stenosis</li> </ul>	Size: n=15,245 pts	
			Exclusion criteria:		
		amlodipine		analysis of PAD	
		valsartan, N=1062	Clinical PAD=2114	Study type: Subgroup	
	vs. 13.6 p=0.63	I● n PAD subaroup N=1052	HTN, CVDRF or CVD.	-	<u>17053536</u>
decrease.	subaroup: Event rates 13.4	<ul> <li>Valsartan vs. amlodipine</li> </ul>	<ul> <li>Age ≥50 v</li> </ul>	amlodipine	2006(141)
<ul> <li>Amlodipine with greater BP</li> </ul>	1° endpoint: In PAD	Intervention:	Inclusion criteria: Overall trial:	Aim: Valsartan vs.	Zanchetti A, et al.
				death, MI, stroke.	
				Primary outcome:	
			Liver failure	Mean follow-up 2.7 y	
			Creatinine ≥4	(total trial: 22,576) pts.	
			Class IV HF	Size: n=2,699 pts	
	CI- 0 74–1 07· n=0 21)		syndrome, AVB >1st degree		
	strategies (HR: 0.89: 95%		Sinus bradycardia, sick sinus	analysis of RCT	
	types of medication		within 1 mo	Prespecified post hoc	
	<ul> <li>No difference between 2</li> </ul>		angina, angioplasty, CABG, stroke	Study type:	
CI 0 77–0 99 n=0 03)	relationshin		<b>Exclusion criteria:</b> Unstable		19996066
● Poor/Fair Ool(HR: 0.87: 95%	<ul> <li>Least trequently SBP</li> <li>135-145 with i-shaped</li> </ul>	atenolol±hctz	<ul> <li>Age ≥50 ywith HTN+stable CAD</li> </ul>	HTN in older with CAD	2010(140)
procedures (HK: 0.94; 95% CI:	• 16.2% in PAD pts	inerapy with	PAU+CAU pts (clinician defined)	VS. BB Dased	PAD subgroup
No difference in vascular	<u>1° endpoint:</u>	Intervention: Intensive	Inclusion criteria:	Aim: Compare CCB	INVEST
	0.39, 93% 01. 0.32-1.07			PAD	
				overall 3 168 with	
	1.09) - Combo vo Dominuil DD:			<b>Cito:</b> n=8 576 nto	
	RR: 1.01; 95% CI: 0.94–			Study type: RCI	

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ABI indicates ankle-bra converting-enzyme inhi blockers; CI, confidence ejection fraction; hctz, t events; MI, myocardial systolic blood pressure	ALLHAT 2002(145) <u>12479763</u>	Paravastu SC, et al. Cochrane Review 2013(144) <u>24027118</u>	NORMA trial Espinola-Klein C, et al. 2011(143) <u>21646599</u>	
<b>Size:</b> n=33,357 pts brachial index; ACEI, angio inhibitor; AE, adverse event ince interval; CLI, critical lin z, hydrochlorothiazide; HF, lial infarction; PAD, periphe liar infarction; PAD, periphe	<u>Aim</u> : Comparison of an alpha blocker, ACE inhibitor, or CCB, each compared to a thiazide-type diuretic on non-fatal or fatal CHD <u>Study type</u> : RCT	<u>Aim</u> : BB Safety in PAD <u>Study type</u> : Update of a review <u>Size</u> : n=119 pts	<u>Aim</u> : Compare BB on walking parameters <u>Study type</u> : RCT <u>Size</u> : n=128 pts	completers
Size: n=33,357 pts           ABI indicates ankle-brachial index; ACEI, angiotensin converting enzyme inhibitor; AE, adverse event; AVB, atrioventricular block; ACD, absolute claudication distance; ACEi, angiotens converting-enzyme inhibitor; AE, adverse event; BP, blood pressure; CABG, coronary artery bypass grafting; CAD, coronary arterial disease; CCB, calcium channel blockers; CI, confidence interval; CLI, critical limb ischemia; CVD, cardiovascular disease; CVDRF, cardiovascular disease risk factors; CV, cardiovascular; DM, diabetes mellitus; EF, ejection fraction; hcz, hydrochlorothiazide; HF, heart failure; HR, hazard ratio; HTN, hypertension; IC, intermittent claudication; LV, left ventricular; MACE, major adverse cardiovascular disease; PVD, peripheral vascular disease; QoL, quality of life; RCT, randomized controlled trial; RR, relative risk; and SBP, systolic blood pressure.	Inclusion Criteria: • Age >50 y • African American15,085 (35.5) • White 19,977 (47.0) • Hispanics 5,299 (12.5) <u>Exclusion criteria</u> : N/A	Inclusion criteria: 6 RCT comparing BB to placebo.	Inclusion criteria: IC+HTN Exclusion criteria: • CLI • Inability to exercise • Contraindications BB • MI within 6 mo • Uncontrolled DM	
, adverse event; AVB, atrioventri e; CABG, coronary artery bypass ase; CVDRF, cardiovascular dise ypertension; IC, intermittent clau cular disease; QoL, quality of life	Intervention: Chlorthalidone vs. Doxazosin, Amlopdipine, or Lisinopril	Intervention: BB vs. placebo	Intervention: Nebivolol 5mg vs. metoprolol 95mg	
cular block; ACD, absolute clauc s grafting; CAD, coronary arteria ase risk factors; CV, cardiovasc dication; LV, left ventricular; MA ; RCT, randomized controlled tr	1° endpoint: Nonfatal MI and fatal CHD	<u>1° endpoint</u> : None of the trials showed worsening of walking measures with BB	1° endpoint: ICD and ACD increased in both groups. No difference between groups.	
block; ACD, absolute claudication distance; ACEi, angiotensin- ing; CAD, coronary arterial disease; CCB, calcium channel isk factors; CV, cardiovascular; DM, diabetes mellitus; EF, on; LV, left ventricular; MACE, major adverse cardiovascular , randomized controlled trial; RR, relative risk; and SBP,	<ul> <li>No difference in primary outcome (nonfatal MI and fatal CHD)</li> </ul>	<ul> <li>No evidence that BB adversely affect walking parameters in IC</li> </ul>	<ul> <li>No difference in ABI change between treatments.</li> <li>7 pts with AE bradycardia Re-enforces safety BB in IC</li> </ul>	

Study Acronym:	Study	Patient Population	Study Acronym: Study Patient Population Primary Endpoint and Results Summar	Summary/Conclusion
Author;	Type/Design;		(include P value; OR or RR;	Comment(s)
Year Published	Study Size		& 95% CI)	
Feringa HH, et al.	Study type:	Inclusion criteria:	All-cause mortality: 44% at median follow-up	<ul> <li>Potential for residual confounding</li> </ul>
2006(146) 0	Observation Cohort	<ul> <li>Referred for Evaluation of PAD</li> </ul>	time of 8 y. MV and propensity score adjusted	<ul> <li>Supports use of BB, ACEi in clinical PAD</li> </ul>
<u>16545650</u>		● ABI ≤0.9	BB HR: 0.68; 95% CI: 0.58–0.80; p<0.001	
	Size: 2,420 PAD pts	● 77% with ABI ≤0.7	ACEi HR: 0.80; 95% CI: 0.69–0.94; p=0.005	
			Nonsignificant: diuretics, CCB	
		Exclusion criteria: N/A		
	Study type:	Inclusion criteria: N/A	<u>1° endpoint:</u> N/A	<ul> <li>Significant benefits in mortality and morbidity</li> </ul>
Sleight P, et al. E	Editorial review			from use of Ramipril in subjects at high risk of
2000(147)		Exclusion criteria: N/A	Results: N/A	future CV events (ACEi could be offered to wider
11967789	<u>Size</u> : n=9,297 pts			group of pts. including those on Aspirin
				prophylaxis).
				<ul> <li>ACEi found to be highly cost effective in a</li> </ul>
				preliminary analysis
				preliminary analysis

pt, patient; and RR, relative risk.

## Evidence Table 19. RCTs for Smoking Cessation–Section 5.4.

	9. RUIS IOF SITION	EVIDENCE TADIE 13. NOTS TO STITUTING CESSALION-SECTION 3.4.			
Study Acronym;	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint
Author;	Study Type;		(# patients) /	(Absolute Event Rates, P value;	(if any);
Year Published	Study Size (N)		Study Comparator	OR or RR; &	Study Limitations;
			(# patients)	95% CI)	Adverse Events
Rigotti NA, et al.	Aim: To compare	Inclusion criteria:	Intervention: Automated	<u>1° endpoint:</u>	<ul> <li>Single-center</li> </ul>
Helping HAND Trial	post discharge	• Age >18 y	voice response calls, free	<ul> <li>Biochemically confirmed tobacco</li> </ul>	<ul> <li>20% lost to follow-up at</li> </ul>
2014(148)	tobacco cessation	<ul> <li>Current smoker</li> </ul>	smoking cessation	abstinence at 6 mo	6 mo
<u>25138333</u>	intervention with	<ul> <li>Plan to quit</li> </ul>	medication for 90 d	• 26% vs. 15% (RR: 1.71; 95% CI:	
	standard care in	<ul> <li>Agree to accept medication</li> </ul>		1.14–2.56; p=0.009) NNT 9.4	
	hospitalized adult	<ul> <li>38% (N=151) with Circulatory Dx:</li> </ul>	Comparator: Printed	<ul> <li>Subgroup analysis in Circulatory</li> </ul>	
	to quit	cardiovascular, peripheral vascular, cerebrovascular	recommendations	disorders showed similar results	
	Study type:	Exclusion criteria: LOS <24 H, no			
	Size: n=397	telephone, substance use (other than tobacco, alcohol, marijuana), admitted for			
	hospitalized adult				

Hennrikus D, et al. 2010(150) <u>21144971</u>	Rigotti NA, et al. 2010(149) <u>20048210</u>	
<u>Aim</u> : To evaluate intensive tailored counseling intervention for smoking cessation in PAD pts <u>Study type</u> : RCT <u>Size</u> : n=124 pts	<u>Aim</u> : To evaluate effect of smoking cessation rates in pts with stable cardiovascular disease. <u>Study type</u> : Multi- center RCT <u>Size</u> : n=714 pts	smokers
<ul> <li>Inclusion criteria:</li> <li>Primary inclusion criteria were a Dx of lower extremity PAD (defined as at least 1 of the following:</li> <li>An ABI of &lt;0.90 in at least 1 lower extremity;</li> <li>A TBI of &lt;0.60.</li> <li>Objective evidence of arterial occlusive disease in 1 lower extremity by duplex ultrasonography, MRA, or CTA</li> <li>Prior leg arterial revascularization or amputation due to PAD</li> <li>Current smoking (defined as smoking ≥1 cigarette a day ≥6 d per wk).</li> <li>Additional inclusion criteria included a desire to quit within the next 30 d</li> </ul>	<ul> <li>Inclusion criteria:</li> <li>Age 35–75 y</li> <li>Want to quit smoking but had not tried in past 3 mo</li> <li>Stable CVD (CAD, PAD, Cerebrovascular disease). PAD=179, 25%</li> <li>Cardiovascular intervention within 2 mo</li> <li>Uncontrolled hypertension</li> <li>Prior amputation</li> <li>Class III/IV CHF</li> <li>Moderate/severe COPD</li> <li>Uncontrolled Gl/hepatic/endocrine disease</li> <li>Severe renal impairment</li> <li>Cancer, depression, psychosis, drug or alcohol use/abuse</li> </ul>	instability, admitted to obstetric or psychiatric units, life expectancy <12 mo
Intervention: Clinician advice, smoking counselor, individualized letter, motivational interview, info about pharmacologic intervention <u>Comparator</u> : Verbal advice, list of programs	Intervention: Varenicline (0.5 once daily for 3 d, 0.5 twice a day for 4 d, 1 mg twice a day for 12 wk) Comparator: Placebo	
<u>1° endpoint</u> : 6 mo biologically confirmed smoking cessation 21.3% vs. 6.8%; chi-square: 5.21; p=0.023	<ul> <li>1° endpoint:</li> <li>4 wk continuous abstinence rate</li> <li>9–12 wk CAR:</li> <li>47% vs. 13.9% (OR: 6.11; 95% CI: 4.18–8.93; p&lt;0.0001)</li> <li>Safety endpoint:</li> <li>SAE 6.5% varenicline vs. 6.0 placebo</li> <li>No difference in psychiatric AEs</li> <li>Non-statistically different but higher rate CV events in varenicline 25 vs. 20</li> </ul>	
NA	<ul> <li>9–52 wk abstinence rate: 19.2 vs. 7.2% (OR: 3.14; 95% CI: 1.93–5.11; p&lt;0.0001)</li> <li>FDA advisory: may increase risk of adverse cardiovascular events</li> </ul>	

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		2014(154) 24323793	Mills EJ et al.		2012(133) 22563098	Hilton JF	) - -		23728631	2013(152)	Stead LF. et al.				<u>12714026</u>	Tonstad S et al. 2003(151)		
	<u>Size</u> : n=63 RCT	analysis	Study type: Meta-		Size: n=22 trials	<u>Study type</u> : Meta- analysis		31,000 pts	2	analysis	Study type: Meta-		Size: n=629 pts	Study type: RCT	CVD	<u>Aim</u> : Buproprion SR in established		
	Exclusion criteria: N/A	and varenicline that reported CVD outcome	Inclusion criteria: RCT of NRT, bupropion,	Exclusion criteria: N/A	• 2 with active CVD, 11 with Hx CVD	<ul> <li>RCT adults with varenicline vs. placebo</li> </ul>	Exclusion criteria: N/A	dinicians	<ul> <li>Trials of smoking interventions involving</li> </ul>	Trials between 1972–2012	Inclusion criteria:	Exclusion criteria: • Seizure • Renal/hepatic/heme/pulmonary neurologic disease • Psychosis • Depression	cigarettes/d during previous 12 mo without quit attempt in previous 3 mo.	<ul> <li>HF (Class I or II)</li> <li>Adults who smoke average ≥10</li> </ul>	• PAD (33%)	Inclusion criteria:	Exclusion criteria: N/A	<ul> <li>Age &lt; Io y</li> <li>Ability to speak and write English</li> <li>No participation in a smoking cessation program in the past 30 d</li> <li>Consumption of &lt;21 alcoholic drinks/wk.</li> </ul>
	<u>Comparator</u> : N/A	bupropion, or varenicline	Intervention: NRT,		Comparator: Placebo	Intervention: Varenicline		<u>Comparator</u> : N/A		cessation advice	Intervention: Smokina			Comparator: Placebo	then 150bid	Intervention: 7 wk buproprion 150/d 1–2,		
	<ul> <li>Buproprion: RR: 1.03; 95% CI:</li> <li>0.71–1.50</li> <li>Varenicline: RR: 1.24; 95% CI:</li> </ul>	<ul> <li>All CVD and MACE</li> <li>NRT: RR 1.81: 95% Cl: 1.35–2.43</li> </ul>	1° endpoint:	Results: RR: 1.40; 95% CI: 0.82– 2.39; p=0.22	discontinuation	<u>1° endpoint</u> : CV events during drug treatment or within 30 d of		<ul> <li>Intensive RR: 1.84; 95% CI: 1.60–</li> <li>2.13</li> </ul>	1.42–1.94	Brief advice RR: 1.66; 95% CI:	1º endnoint:			2.24-4.84)	43% vs. 19% (OR: 3.27; 95% CI:	1° endpoint: 4 wk smoking cessation		
66			N/A		meta-analysis	<ul> <li>Risk of cardiovascular</li> <li>SAE with varenicline use:</li> </ul>			rates	small effect on cessation	<ul> <li>Simple advice has a</li> </ul>					N/A		

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$^{\odot}$ American Heart Association, Inc. and American College of Cardiology Found:
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	1.21; 95% CI: 1.02–1.43; p=0.03)			<u>24502815</u>
with early gratt failure.	Results. Higher early graft failure in active smokers (OR:	<ul> <li>Intrainguinal bypass surgery</li> <li>Pre-operative smoking status</li> </ul>	<b>Size:</b> n=16 534 nts	Selvarajan S, et al. 2014(157)
<ul> <li>Active smoking associated</li> </ul>	1° endpoint: 30 d graft failure	Inclusion criteria:	Study type: Registry	ACS/NSQIP
	<ul> <li>For a surgeon's onered phannacongic interapy of referral to smoking cessation program. Rates of cessation higher in these surgeons 48% vs. 33%</li> </ul>	<ul> <li>Exclusion criteria:</li> <li>Lost to follow-up at 1 y</li> <li>Lack of smoking status at 1 y</li> </ul>		
cessation after surgical procedures	Variability across treatment center in smoking cessation rates 28%–62%	AAA repair		
<ul> <li>High rates of smoking</li> </ul>	<ul> <li>46% pts post LE bypass quit at 1 y</li> </ul>	• LE bypass		23375433
smoking cessation in pts with vascular disease	Results:	CEA     Carotid stent	<u>Size</u> : n=7,807 pts	2013(156)
Systems of care promote	1° endpoint: Self-reported smoking cessation at 1 y	Inclusion criteria:	Study type: Registry	VSGNE
	• Long-term Quitters RR: 0.42; 95% CI: 0.16–1.10			
	<ul> <li>M:</li> <li>Recent Quitters RR: 0.36; 95% CI: 0.04–2.97</li> </ul>	CVD.		
	• Long-term Quitters RR: 0.29; 95% CI: 0.16–0.52		evenis.	
	<ul> <li>Recent Quitters RR: 0.61; 95% Cl: 0.21–1.78</li> </ul>	<ul> <li>Stratified by DM</li> </ul>	tollow-up 25 y, 631 CVD	
	No DM:	nonsmoker	Size: n=3,251 pts, mean	
	Results:	long-term quitter >4 y,		
weignt gain.	PAU events=/3	<ul> <li>self-reported smoking status: smoker, recent quitter (&lt;4 y),</li> </ul>	on cardiovascular events	
PAD) even when adjusting for	disease, PAD, congestive heart failure).	1984–2011.	impact of weight gain on the	234831/6
with lower CVD rates (including	CVD events (coronary heart disease, cerebrovascular	<ul> <li>Longitudinal cohort study</li> </ul>	cohort. To investigate the	2013(155)
<ul> <li>Smoking cessation associated</li> </ul>	<u>1° endpoint:</u>	Inclusion criteria:	Study type: Prospective	Clair C, et al.
	& 95% CI)			Year Published
Comment(s)	(include P value; OR or RR;	-	Study Size	Author;
Summary/Conclusion	Primary Endpoint and Results	Patient Population	Study Type/Design;	Study Acronym;
	Evidence Table 20. Nonrandomized Trials, Observational Studies, and/or Registries of Smoking Cessation–Section 5.4.	bservational Studies, and/or	Nonrandomized Trials, O	Evidence Table 20.
	ratio; PAD, peripheral artery disease; pt, patient; RCT, randomized controlled trial; RR, relative risk; and SAE, serious adverse event.	ndomized controlled trial; RR, relativ	ery disease; pt, patient; RCT, ra	ratio; PAD, peripheral art
needed to treat; NRT, nicotine replacement therapy; OR, odds	v, not applicable; NNT, number needed to treat; NRT, nicotine	stay; MACE, major adverse cardiovascular event; MRA, magnetic resonance angiogram; N/A, not applicable; NNT, number i	se cardiovascular event; MRA, I	stay; MAČE, major adver
and Drug Administration; GI, gastrointestinal; LOS, length of	CV, cardiovascular; FDA, Food and Drug Administration; GI,	pulmonary disease; CTA, computed tomography angiography; CVD, cardiovascular disease; CV, cardiovascular; FDA, Food	, computed tomography angiogr	pulmonary disease; CTA,
D. chronic obstructive	AE indicates adverse event: CAD. coronary arterial disease: CAR. continuous abstinence rate: CHF. concestive heart failure: CI. confidence interval: COPD. chronic obstructive	se: CAR. continuous abstinence rate	ent: CAD. coronarv arterial disea	AE indicates adverse eve

0.85-1.81

		Exclusion criteria: N/A		
UCSD	Study type: Retrospective	Inclusion criteria:	1° endpoint: Amputation-free survival	<ul> <li>Smoking cessation</li> </ul>
Armstrong EJ, et al.	cohort	<ul> <li>Peripheral angiography for</li> </ul>		associated with better
2014(158)		claudication or CLI	Results:	outcomes in PAD.
<u>25282696</u>	<u>Size</u> : n=204 pts	<ul> <li>Active smoking at time of angingraphy</li> </ul>	<ul> <li>Smoking cessation associated with lower mortality 14%</li> <li>vs 31% (HR: 0.40: 95% CI: 0.18–0.90</li> </ul>	
		30% quit for 1 y	<ul> <li>Higher amputation-free survival 81% vs. 60% (HR: 0.43; 95% Cl: 0.2–0.86)</li> </ul>	
		Exclusion criteria: N/A		
Scottish Family	Study Type: Cross-	Inclusion criteria:	Results: Second-hand smoke exposure (≥40 hrs/wk)	No cotinine levels available,
Health Study	sectional cohort study	<ul> <li>Never smokers</li> </ul>	higher prevalence PAD (OR: 5.56; 95% CI: 1.82–17.06;	cross-sectional
Lu L, et al		• Age ≥18 y	p=0.003)	
2013(159)	Size: n=5,686 pts, 134			
23880175	(2.4% with PAD defined by	Exclusion criteria: N/A		
Tan CE and Glantz SA 2012(160)	<u>Study Type</u> : Meta-analysis of impact of smoke-free	Inclusion criteria: Studies published before November 30,	Results: Smoke-free legislation associated with lower hospital admission or death for: coronary events (RR:	Did not ascertain PAD events
<u>23109514</u>	laws with coronary, heart	2011	0.84; 95% CI: 0.82–0.88), other heart disease (RR: 0.61;	
	disease, cerebrovascular		95% Cl: 0.44–0.85), cerebrovascular events (RR: 0.84;	
	events	Exclusion criteria: N/A	95% CI: 0.75–0.94)	
	<u>Size</u> : n=45 studies of 33 smoke-free laws			
AAA indicates abdominal a	aortic aneurysm; ABI, ankle-bra	chial index; CEA, carotid endartere	AAA indicates abdominal aortic aneurysm; ABI, ankle-brachial index; CEA, carotid endarterectomy; CLI, critical limb ischemia; CI, confidence interval; CVD, cardiovascular disease; DM, diabetes mellitus: HR hazard ratio: LE lower extremity: N/A not annlicable: OR odds ratio: PAD nerioheral artery disease: and RR relative risk	D, cardiovascular disease; DM,
diabetes mellitus; HR, haz	zard ratio; LE, lower extremity; I	I/A, not applicable; UR, odds ratio; I	diabetes mellitus; HR, hazard ratio; LE, lower extremity; N/A, not applicable; OR, odds ratio; PAD, peripheral artery disease; and RR, relative risk.	

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## Evidence Table 21. RCTs Evaluating Glycemic Control in Patients with PAD and Diabetes Mellitus–Section 5.5.

	Year Published	Author;	Study Acronym;	
	Study Size (N)	Study Type;	Aim of Study;	
			Patient Population	
(# patients)	Study Comparator	(# patients) /	Study Intervention	
95% CI)	P value; OR or RR; &	(Absolute Event Rates,	Endpoint Results	
	Adverse Events	Study Limitations;	Relevant 2° Endpoint (if any);	

	● PAD n=1,0⁄ 1,274 subset	randor <u>Size</u> : • n=5,	DM Study		Dormandy JA et whether
	<ul> <li>● PAD subgroup ~20% n=1,043 (reported as 1,274 in 2009 PAD subset publication)</li> </ul>		DM <u>Study type</u> : Double blind. placebo controlled	reduces macrovascular morbidty and mortality in high-risk pts with type 2	<u>Aim</u> : To ascertain whether pioglitazone
	Exclusion criteria: • Type I DM • Pt only on insulin • Planned coronary/peripheral revascularization	<ul> <li>or surve or objective alternation of surve or objective alternation of PAD</li> <li>PAD defined as major amputation or claudication+ABI &lt;0.9</li> </ul>	<ul> <li>agents (with or without insulin).</li> <li>Evidence of "extensive macrovascular disease" CAD or stroke or "objective arterial</li> </ul>	<ul> <li>Age 35–75 y</li> <li>HgB A1c &gt;6.5% despite treatment with diet or oral</li> </ul>	Pts with DM
			adjusted if needed) <u>Comparator</u> : Placebo	qd mo 1; 30 mg mo 2; 45 qd mo 3-end; medication could be	pioglitazone (15 mg
	<b>Safety endpoint:</b> No difference in CHF admissions or death due to CHF between pioglitazone and placebo	<u>1° endpoint</u> : HR: 0.90; 95% Cl: 0.80−1.02; p=0.095	amputation Average follow-up 34.5 mo.	nonfatal MI, stroke, ACS, coronary or peripheral revascularization, major	<u>1° endpoint</u> : Composite all-cause mortality,
rnere was a beriericial effect of progritazone seen.	subset of pts enrolled with PAD (N=1,274 reported), there was no benefit of pioglitazone on the primary or secondary endpoint with increased rate of LE revascularization in the pioglitazone vs. placebo groups (p=0.0077). In the subgroup of pts randomized WITHOUT PAD,	<ul> <li>vs. placebo; no PAD specific data presented,</li> <li>vs. placebo; no PAD specific data presented,</li> <li>though 20% of pt population had sx PAD</li> <li>PAD substudy (2009 publication): PAD subset</li> <li>had higher event rates than non-PAD subset. In</li> </ul>	<ul> <li>Summary:</li> <li>Primary endpoint was negative, but secondary endpoint (primary for most studies of MACE) notifice for reduction in events with ninclitazone</li> </ul>	0.84; 95% CI: 0.72–0.98; p=0.027 • Subgroup analysis for PAD not reported.	<ul> <li><u>2° endpoint</u>:</li> <li>All-cause mortality, non–fatal MI, stroke HR:</li> </ul>

Evidence Table 22. Nonrandomized Trials, Observational Studies, and/or Registries of Glycemic Control-Section 5.5.

Study Acronym; Author;	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR;	Summary/Conclusion Comment(s)
Year Published			& 95% CI)	
PAD-UCD	Study type: Observational	Inclusion criteria: Pts	1° endpoint: Patency of the target lesion	<ul> <li>Observational study provides</li> </ul>
Singh S, et al.	registry of pts undergoing	with PAD within a		some support for adequate peri-
2014(162)	interventional procedures	peripheral interventional	Results: Pts with peri-procedural FBG values below	procedural glycemic control with
24939930	for CLI or ALI at a single	registry with DM with CLI	the median value of 144 mg/dL had improved primary	revascularization for infrapopliteal
	center	or ALI who underwent	patency at 1 yr (46% vs. 16%; HR: 1.82; p=0.005);	lesions in pts with DM with ALI/CLI
		infrapopliteal intervention	association robust after adjustment for insulin use and	to prevent MALE, possibly patency

Strong Heart Study Resnick HE, et al. 2004(164) <u>14970108</u> cc wi		∞ ۱ <u>۵</u>	Takahara M, et al. <u>Si</u> 2010(163)         cc <u>20843974</u> re           (s)         (s)	<u>N</u>
Study type: Observational cohort study <u>Size:</u> n=4,549 in entire cohort; 1,974 with DM without prior lower extremity amputation		a single center <mark>Size:</mark> n=278 pts; 197 pts with DM	Study type: Observational cohort study vs. retrospective chart review (study design not clear) at	<u>Size:</u> n=149 pts, 309 PTA procedures
Inclusion criteria: Native Americans age 45–74 y seen for baseline examination 1989–1992 and subsequent follow-up visits Exclusion criteria: Pts without DM; those with prior LE amputation excluded		<b>Exclusion criteria</b> : Pts with CLI who were not candidates for PTA and treated by other means	Inclusion criteria: Pts with PAD undergoing PTA for CLI including pts with and without DMs	during the study period <u>Exclusion criteria</u> : No FBG on day of angiogram procedure or within 2 d of the procedure
<u>1° endpoint</u> : Incident lower extremity amputation <u>Results</u> : After average 8 yr follow-up. Among pts with PAD (ABI <0.9), higher HgB A1c increased odds of lower extremity amputation. Relationship also seen among pts with normal ABI and those with non- compressible vessels (ABI >1.4). Odds of incident LE amputation among pts with DM and PAD (ABI <0.9) or non-compressible vessels (ABI ≤1.4); reference pts with DM with normal ABI and HgB A1c <6.5%* (OR=1)	Increased quartiles of HgB A1C had stepwise increase in risk for major amputation, adjust HRs (for Fontaine Stage IV, dialysis, infection) Quartile Adjusted HR Q1 $\leq$ 5.9% - Q2 6-6.7% 2.030 (0.657-6.266, p NS) Q3 6.8–7.6% 3.398 (1.227-9.412, p=0.019) Q4 $\geq$ 7.7% 3.983 (1.398-11.35, p=0.010)	Among 287 CLI pts with DM: HgB A1c level not associated with increased mortality HgBA1c level associated with major amputation, adjusted HR: 1.349 per 1% increment; 95% CI: 1.103– 1.650; p=0.004) Association was robust after MV adjustment for other factors.	<u>1° endpoint</u> : Major amputation, mortality (all-cause) <u>Results</u> : Average follow-up 90±72 wk.	lesion characteristics One yr major adverse limb events lower for pts with FBG below median (23% vs. 35%; p=0.05)
• Epidemiological cohort study providing evidence of an association between HgBA1c/glycemic control and risk of LE amputation among pts with DM with PAD and also those with non compressible vessels (most of whom have PAD when assessed by other means)		undergo revascularization (pre- procedural HgB A1c) to reduce risk of amputationassociation more pronounced for highest quartile of HgB A1c vs. lowest quartile. • No mortality benefit seen over a relatively short period of follow-up	<ul> <li>Another observational study providing some support for adequate glycemic control among PAD pts with DM with CLI who will</li> </ul>	of PTA sites

Higb Alic Age adjusted OK LE amp         <6.5%       1.7         <6.5%       1.7         <6.5%       5.6 (p<0.05)         >9.5%       8.7 (p<0.05)         ABI indicates ankle-brachial index; ALI, acute limb ischemia; Cl indicates confidence interval; CLI, critical limb ischemia; DM, diabetes mellitus; FBG, fasting blood glucose; HgbA1c, brachial index; ALE, major adverse limb event; MV, multivariate; NS, non-significant; OR, odds ratio; PAD, peripheral artery disease; PTA,	Pts with DM with PAD ABI <0.9
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percutaneous transluminal angioplasty; pt, patient; and RR, relative risk. ( 1..... ,

## Evidence Table 23. RCTs Evaluating Oral Anticoagulation–Section 5.6.

	J. NO IS EVALUALING O	Evidence Table 23. No 13 Evaluating Otal Anticoagulation-Section 3.0.	JII J.U.		
Study Acronym;	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Year Published	Study Size (N)		Study Comparator	value; OR or RR; &	Adverse Events
			(# patients)	95% CI)	
WAVE TRIAL	Aim: Evaluate	Inclusion criteria:	Intervention:	1° endpoint: MI, stroke, or	<ul> <li>Mean follow-up 35 mo</li> </ul>
Anand S, et al.	anticoagulant agents in	• Age 35–85 y	Anticoagulation and	death no difference (12.2% vs.	
2007(165)	prevention of	<ul> <li>PAD defined as</li> </ul>	antiplatelet	13.3%, p=0.48)	Summary:
17634457	cardiovascular	atherosclerosis of the arteries			<ul> <li>Combination of an anticoagulant and</li> </ul>
	complications in pts	of the lower extremities, the	<u>Comparator:</u>	1° Safety endpoint: Life	antiplatelet therapy not more effective
	with PAD	carotid arteries, or the	Antiplatelet alone	threatening bleeding significantly	than antiplatelet therapy alone in
		subclavian arteries		increased (4.0% vs. 1.2%,	preventing major cardiovascular
	Study type: RCT			p<0.0001)	complications and associated with
		Exclusion criteria:			increase in life-threatening bleeding
	Size: n=2,161 pts	<ul> <li>Indication for oral</li> </ul>			
		anticoagulant treatment			
		<ul> <li>Actively bleeding or at high</li> </ul>			
		risk for bleeding			
		<ul> <li>Stroke within 6 mo before</li> </ul>			
		enrollment			
		<ul> <li>Dialysis</li> </ul>			

	No difference in RBC transfusions				
	(32% vs. 3.7%, p=0.004)				
	<ul> <li>Postop hematoma increased</li> </ul>			Size: n=64 pts	
and limb salvage.	Safety endpoint:			2	
hematomas, but improves patency rate				Study type: RCT	
<ul> <li>Anticoagulation atter vein bypass</li> </ul>	(01% VS. 01%; p=0.01)				
Summary:				rates, and complication	
	56%, p=0.2)	alone	Exclusion criteria: N/A	patency, limb salvage	
<ul> <li>Did not evaluate stroke, MI</li> </ul>	vs. 56%, p=0.5; SP: 81% vs.	Comparator: ASA		bypass on duration of	
unclear			risk for graft occlusion	after autogenous vein	<u>9737454</u>
<ul> <li>Small study</li> <li>Definition of high risk for bypass failure</li> </ul>	<ul> <li>3 y patency improved (PP:</li> </ul>	and ASA	Inclusion criteria: Infrainguinal vein bypass high	AIM: Effects of anticoagulation therapy	5arac 1P, et al. 1998(168)
-	(p=0.02)				
	<ul> <li>Maior hemorrhane increased</li> </ul>				
<ul> <li>Benefit in subgroup analysis of natency for 6 mm PTEF</li> </ul>	<ul> <li>Mortality increased (32% vs.</li> <li>23%: n=0 0001)</li> </ul>			end 100-111-021	
hemorrhage.	Safety endpoint:			<b>Ciro:</b> n=921 nto	
increased mortality and major				Study type: RCT	
ASA no difference in overall patency but	vs. 58%; p=0.02)	Antiplatelet alone	warfarin		
<ul> <li>Anticoagulation + ASA compared to</li> </ul>	analysis significant benefit (71%	Comparator:	Contraindication to ASA or	patency	
Summary:	<ul> <li>6 mm PTFE bypass subgroup</li> </ul>	-	Exclusion criteria:	morbidity and bypass	<u>11877686</u>
c	difference	antiplatelet	:	ASA alone on mortality,	2002(167)
anticoagulation	<ul> <li>Bypass patency no significant</li> </ul>	Anticoagulation and	bypass for PAD	+ ASA therapy) vs.	Williford WO
<ul> <li>1/3 of anticoagulation pts stopped</li> </ul>	1° endpoint:	Intervention:	Inclusion criteria: Any	Aim: Evaluate warfarin	Johnson WC and
			• Anemia	Size: n=2,690 pts	
			<ul> <li>Abnormalities of platelets</li> </ul>		
a construction of the second			surgerv	Study type: RCT	
bleeding complications	1.42-2.71)		MI or stroke 1 mo before		
subgroup analysis and increased	increased (HR: 1.96; 95% CI:		<ul> <li>Shortened life expectancy</li> </ul>	clinical events	
<ul> <li>No difference other than in vein graft</li> </ul>	Safetv endpoint: Bleeding		medications	graft occlusion and	
Summary:			<ul> <li>Contraindication to trial</li> </ul>	infrainguinal bypass-	
	amputation no difference		Exclusion criteria:	ASA in prevention	
anticoaculation	Vascular death. MI stroke or	Comparator: ASA		anticoagulants with	10665553
<ul> <li>Wean rollow-up 2 into</li> <li>Vein graft subset-benefit to</li> </ul>	<ul> <li>Graft occlusion no difference</li> </ul>		Infrainguinal bypass for PAD	effectiveness of oral	2000(166)
Mean follow-up 21 mo	1º endnoint:	Intervention: Warfarin	Inclusion criteria	Aim: Compare	ROA TRIAI

Evidence Table 24.	Nonrandomized Trials,	Observational Studies,	Evidence Table 24. Nonrandomized Trials, Observational Studies, and/or Registries of Oral Anticoagulation–Section 5.6	n-Section 5.6.
Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Summary/Conclusion Comment(s)
Alonso-Coello P, et al. 2012(170)	Study type: Clinical practice guidelines based	Inclusion criteria: • Asx PAD	<ul> <li><u>1° endpoint</u>:</li> <li>Prevention of cardiovascular disease</li> </ul>	<ul> <li>Recommend against the use of warfarin + ASA in pts with asx or sx PAD (Grade 1B)</li> </ul>
<u>22315275</u>	on meta-analysis of 3 RCTs evaluating warfarin + ASA vs. ASA alone.	<ul> <li>Sx PAD</li> <li>ALI</li> <li>Post peripheral arterial</li> </ul>	<ul> <li>Relief of lower extremity symptoms and critical ischemia</li> </ul>	
		revascularization	Results: Results failed to demonstrate or	
		Carotid stenosis	alone on mortality, nonfatal MI, or nonfatal	
		Exclusion criteria: N/A	stroke. However, there was a significant increase in major bleeding events with warfarin.	
Bedenis R, et al. 2015(171)	Study type: Cochrane Review	Inclusion criteria: Lower extremity bypass for PAD	1° endpoint: Bypass primary patency	<ul> <li>No patency benefit with use of anticoagulation</li> </ul>
<u>25695213</u>	<b><u>Size</u>:</b> n=1,381 pts in the 3	Exclusion criteria: N/A	Results: No difference in primary graft patency when ASA or ASA with dipyridamole was	
	studies included for the analysis of anticoagulants.		compared to a vitamin K antagonist	
Cosmi B, et al.	Study type: Cochrane	Inclusion criteria: IC,	<u>1° endpoint:</u>	<ul> <li>No significant difference was observed</li> </ul>
2001(172)	Review	RCT data	Maximum walking distance	for pain-free walking distance or maximum
	Size: n=3 studies in the	Exclusion criteria: N/A		walking distance at the end of treatment
	primary analysis; 4		Results: No benefit of heparin, LMWHs or oral	<ul> <li>Major and minor bleeding events were</li> </ul>

Antonicelli R, et al. 1999(169) <u>10492316</u>

<u>Aim</u>: Evaluate the efficacy of low-dose,

Inclusion criteria:

Intervention: Subcutaneous heparin and ASA

Maximum walking time 40% in

the study

132 of 201 randomized pts completed

1° endpoint:

Size: n=201 pts

Study type: RCT

Exclusion criteria: N/A

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examination

alone

Comparator: ASA

Pain-free walking time 39% in

subcutaneous heparin is safe and

Treatment with low-dose

Summary:

effective in improving walking

performance

heparin group and 23% in placebo group (p=0.09).

in placebo group (p=0.05)

heparin group and 16%

Willingness to use parenteral therapy
≥6 mo Hx of IC who had PAD confirmed by Doppler

subcutaneous calciumheparin in comparison with placebo in pts with

additional studies were included in the sensitivity analysis been observed especially with oral anticoagulants. The use of anticoagulants for IC cannot be recommended at this stage. increased risk of major bleeding events has anticoagulants has been established for IC. An significantly more frequent in the group treated with oral anticoagulants compared to control, with a nonsignificant increase in fatal bleeding events.

ALI indicates acute limb ischemia; ASA, aspirin; IC, intermittent claudication; LMWH, low molecular weight heparin; N/A, not applicable; PAD, peripheral arterial disease; pt, patient; and RCT, randomized controlled trial.

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Cilostazol–Section 5.7.
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Study Author; Year Published	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if
	Study Size (N)		(# patients) / Study Comparator (# patients)	(Absolute Event Rates, P value; OR or RR; & 95% Cl)	any); Study Limitations; Adverse Events
Bedenis R, et al. 2014 (173)	Aim: To determine Cilostazol's impact on	Inclusion criteria: Cilostazol with	All included studies compared cilostazol	For 8 studies data were compatible for comparison by meta-analysis, but data for 7 studies were too	There was no association between treatment type and
25358850	claudication walking	placebo, or	100mg 2x/d with placebo.	heterogeneous to be pooled. For the studies	all-cause mortality for any of
	distances, mortality,	medications	In addition, 2 studies	included in the meta-analysis, for ICD there was an	the treatment comparisons,
	and vascular events in	currently known to	compared cilostazol 50	improvement in the cilostazol group for the 100 mg	but there were very few
	pts with stable IC.	increase walking	mg 2x/d with placebo, and	and 50 mg 2x/d, compared with placebo (WMD:	events, and therefore
		distance e.g.	1 study compared	31.41 meters; 95% CI: 22.38–40.45 meters;	inadequately powered. In
	Study type: Meta-	pentoxifylline. All pts	cilostazol 150 mg 2x/d	p<0.00001) and (WMD: 19.89 meters; 95% CI:	general cilostazol was
	analysis: Double-blind,	had IC secondary to	with placebo. 3 studies	9.44–30.34 meters; p=0.0002), respectively. ICD	associated with a higher odds
	RCTs of cilostazol vs.	PAD.	compared cilostazol 100		of headache, diarrhea,
	placebo, or vs. other		mg 2x/d with pentoxifylline	comparison of cilostazol 150 mg vs. placebo and	abnormal stool, dizziness and
	antiplatelet agents in		400 mg 3x/d. 1 study	cilostazol 100 mg vs. pentoxifylline, but only single	palpitations
	pts with stable IC.		compared cilostazol 100	studies were used for these analyses. ACD was	
			mg 2x/d with pentoxifylline	significantly increased in pts taking cilostazol 100	
	<u>Size</u> :		600 mg 2x/d and 1 study	mg and 50 mg 2x/d, compared with placebo (WMD:	
	<ul> <li>n=15 studies.</li> </ul>		compared cilostazol 100	43.12 meters; 95% CI: 18.28–67.96 meters;	
	<ul> <li>n=3,718 pts</li> </ul>		mg 2x/d with the	p=0.0007) and (WMD: 32.00 meters; 95% CI:	
			antiplateletik-134 ou mg	14.1/–49.83 meters; p=0.0004), respectively. As	
			and 100mg 2x/d	with ICD, ACD was increased in pts taking cilostazol	
				150 mg vs. placebo, but with only 1 study an	
				association cannot be clearly determined. 2 studies	
				comparing cilostazol to pentoxitylline had opposing	
				tindings, resulting in an imprecise CI (WMD: 13.42	
				meters (95% Cl: -43.51 – 70.35 meters; p=0.64).	
				ABI was lowered in the cilostazol 100 mg group	
				compared with placebo (WMD: 0.06; 95% CI: 0.04–	
				0.00, p<0.0001). The single study evaluating Add	

Goldenberger NA, et al. 2012 (175) <u>22615190</u>	Dawson DL, et al. 2000 (174) <u>11063952</u>
<u>Aim</u> : To investigate the effect of cilostazol + I-carnitine vs. cilostazol alone on treadmill performance in IC. <u>Secondary</u> <u>objectives</u> : To evaluate QoL measures and safety indices with the drug	<u>Aim</u> : To determine evaluate the relative efficacy and safety of cilostazol and pentoxifylline. Randomized, double- blind, placebo- controlled, multicenter trial. <u>Size</u> : n=698 pts
Inclusion criteria: PAD pts with stable IC were randomized to either I-carnitine 1 g or matching placebo 2x/d, on a background of cilostazol.	Inclusion criteria: • Moderate-to- severe claudication • Baseline pain-free walking distance ≥53.6 m • Baseline maximal walking distance ≤537.6 m Exclusion criteria: • Buerger's disease • Critial ischemia (category II or III chronis lower extremity ischemia) • Lower extremity arterial reconstruction (surgical or endovascular) or sympathectomy within 3 mo • Prior use of cilostrazol
145 pts met criteria for the mITT population and 120 pts for the per-protocol population. 74 L- carnitine/71 placebo.	Study intervention: Pentoxifylline or cilostazol <u>Comparator</u> : Placebo
In the mITT (n=145), the mean In ratio in PWT was 0.241 for cilostazol/I-carnitine vs. 0.134 for cilostazol/placebo (p=0.076), corresponding to mean increases of 1.99 and 1.36 min, respectively. In the per-protocol population (n=120), the mean <i>/n</i> ratio in PWT was 0.267 for cilostazol/I-carnitine vs. 0.145 for cilostazol/placebo (p=0.048).	<ul> <li>for the comparison of cilostazol vs. pentoxifylline found no change in ABI.</li> <li>Primary endpoint: Walking ability, measured by MWD.</li> <li>Cilostazol treatment resulted in greater MWD than both pentoxifylline treatment resulted in no improvement in MWD compared to placebo</li> <li>Secondary endpoints: PFWD and resting Doppler limb pressures</li> <li>At wk 4 and after, there was a greater improvement in PFWD with cilostazol treatment than placebo (p&lt;0.01)</li> <li>There was no difference in PFWD with pentoxifylline treatment compared with placebo (p&lt;0.05).</li> </ul>
The per-protocol population, the mean <i>In</i> ratio in PWT was significantly increased in the cilostazol/l-carnitine group vs. the cilostazol/placebo group (0.267 vs. 0.145, respectively; p=0.048). This represented an arithmetic mean increase in PWT of 39.2% from baseline to d 180 for cilostazol/l- carnitine, as compared to 21.5% for cilostazol/placebo.	<ul> <li>Withdrawal rates due to adverse effects were similar among the cilostazol (16%) and the pentoxifylline treatments (19%)</li> <li>Adverse events were higher in the active treatment groups than in placebo (27% for placebo; p=0.006)</li> <li>Overall results have not shown clear evidence of an improvement in walking performance with pentoxifylline treatment.</li> </ul>

Warner CJ, et al. 2014 (176) 24468286	
<u>Aim</u> : MEDLINE (1946- 2012), and Cochrane CENTRAL (1996- 2012), and trial registries searched for studies comparing cilostazol in combination with antiplatelet therapy alone after PVI. <u>Study type</u> : Meta- analysis: <u>Size</u> : n=1,522 pts	combination. <u>Study type</u> : A multicenter, randomized, double- blind, placebo- controlled trial <b>Size:</b> n=164 pts
<ul> <li>Inclusion criteria:</li> <li>Pts undergoing endovascular treatment (angioplasty or stenting) for infrainguinal LE PVD.</li> <li>The intervention must be cilostazol in the periprocedural setting.</li> <li>The comparison group may be no cilostazol, an antiplatelet medication, or placebo.</li> <li>≥6 mo follow-up</li> <li>The study reported at ≥1 pre- specified outcome of interest (restenosis, freedom from amputation, mortality).</li> </ul>	
<ul> <li>2 RCTs and 4</li> <li>retrospective cohorts met inclusion criteria. 1,522 pts included in the review.</li> <li>A majority (87%) were from retrospective cohort studies.</li> <li>All studies were conducted in Japan and published between 2008– 2012.</li> <li>All compared cilostazol (n=4) or an alternative antiplatelet medication (n=2), with both groups receiving various co- interventions (ASA with or without an adjunct antiplatelet medication).</li> </ul>	
The addition of cilostazol was associated with decreased restenosis (RR: 0.71; 95% CI: 0.60–0.84; p<0.001), improved amputation-free survival (HR: 0.63; 95% CI: 0.42–0.66; p<0.001), and improved freedom from target lesion revascularization (RR: 1.36; 95% CI: 1.14–1.61; p<0.001).	
There was no significant reduction in mortality among those receiving cilostazol (RR: 0.73; 95% CI: 0.45–1.19; p=0.21). p=0.21).	In the cilostazol/l-carnitine group, the mean increase in physical functioning on the SF-36v2 was also nearly double that of the cilostazol/placebo group (6.77 [16.379] vs. 3.73 [17.566], respectively; p=0.066).

ABI indicates ankle- intent-to-treat; LE, lo transluminal angiopi polytetrafluoroethyle					<u>23652861</u>	2013 (177)	lida O, et al.	
brachial index; ACD, abs wer extremity; mITT, mo asty; PVD, peripheral vas me; pt, patient; QoL, qual	Size: n=152 pts: 75 in cilostazol/77 placebo	Study type:	stenting for femoropopliteal disease	after PTA with	cilostazol reduces restenosis at 12 mo	whether treatment with	angiographic follow-up	<b>Aim:</b> To determine by
olute claudication distanc dified intent-to-treat; MW scular disease; PVI, perip ity of life; RCT, randomiz				provisional nitinol stenting.	pt was admitted and underwent PTA with	randomization, each	Within 1 wk after	Inclusion criteria:
e; Cl, confidence interval; HF D, maximal walking distance; heral vascular intervention; F d controlled trial; RR, relativ					Study comparator: 77		cilostazol	Ctudy intervention: 75 in
ABI indicates ankle-brachial index; ACD, absolute claudication distance; CI, confidence interval; HR, hazard ratio; IC, intermittent claudication; ICD, initial claudication distance; ITT, intent-to-treat; LE, lower extremity; mITT, modified intent-to-treat; MWD, maximal walking distance; PAD, peripheral artery disease; PFWD, pain free walking distance; PTA, percutaneous transluminal angioplasty; PVD, peripheral vascular disease; PVI, peripheral vascular intervention; PWT, peak walking time; RCT, randomized controlled trial; and PTFE, polytetrafluoroethylene; pt, patient; QoL, quality of life; RCT, randomized controlled trial; RR, relative risk; and WMD, walking maximal distance.				by ITT analysis.	12 mo was 20% (15/75) in the cilostazol group vs. 49% (38/77) in the noncilostazol group (p=0.0001)	data at 12 mo. The angiographic restenosis rate at	died and 152 pts (80%) had evaluable angiographic	Describe. During the 19 ms follow-up pariod 11 nte
slaudication distance; ITT, ng distance; PTA, percutaneous al; and PTFE,				were similar in both groups.	vs. 71%, p=0.02), although cardiovascular event rates	free survival at 12 mo (83%	a significantly higher event-	The electron area and a

Evidence Table 26. Nonrandomized Trials, Observational Studies, and/or Registries of Pentoxifylline–Section 5.8

LAINCHCC I NAIC		india, Obaci	rational otheres, and/or region	Evidence Table 20. Notifalidontized Thata, Observational Otables, and/or Negletiles of Ferrovity intervection 5.0.	
Study	Aim of Study;	Patient	Study Intervention (# patients) /	Endpoint Results	Relevant 2° Endpoint
Acronym;	Study Type;	Population	Study Comparator (# patients)	(Absolute Event Rates, P value; OR or RR; &	(if any);
Author;	Study Size (N)			95% CI)	Study Limitations;
Year Published					Adverse Events

	when compared with nylidrin, GBE and ASA. Cilostazol, PGE1 and flunarizine showed larger improvements in TWD compared with pentoxifylline. • Pentoxifylline appeared to be well tolerated in most				
	<ul> <li>Pentoxitylline.</li> <li>For TWD a larger improvement was shown for Pentoxifylline showed a larger improvement in TWD</li> </ul>				
	Cilostazol (2 studies) and PGE1 (2 study)     showed a larger improvement in PFWD compared with				
	study) and iloprost (1 study).	Comparator: Active agents			
N/A	Pentoxifylline showed a larger improvement in PFWD     when compared with CBE /1 study, buttomedii /1	Study intervention: Pentoxifylline			
	difference, one study snowed a significant improvement in QoL.				
	Operation of the second s				
	PentoXitylline TWD (14 studies): 1%_155 0% with pentovifulline				
	• PFWD (11 studies): -33.8%-73.9% with	Comparator: Placebo			
NA	<ul> <li>Large variability in results. Unable to perform meta- analysis because of variability.</li> </ul>	Study Intervention: Peritoxilyiline	procedures		
		with iloprost and placebo	or percutaneous		
		<ul> <li>1 study compared pentoxifylline</li> </ul>	Indergone surgical		
		with cilostazol and placebo	critical ischemia or		
		<ul> <li>2 studies compared pentoxifylline</li> </ul>	criteria: Pts with		
		with nifedipine	Exclusion	2015)	
		<ul> <li>1 study compared pentoxifylline</li> </ul>		(Current until April	
		<ul> <li>2 studies compared pentoxitylline with PGE1</li> </ul>	<ul> <li>Fontaine stage if due to PAD</li> </ul>	with 3,377 pts	
		with nylidrin hydrochloride		Size: n=94 studies	
		<ul> <li>1 study compared pentoxifylline</li> </ul>	<ul> <li>Symptoms of</li> </ul>	Cochrane review	
<ul> <li>Pentoxitylline was generally well tolerated.</li> </ul>		<ul> <li>1 study compared pentoxitylline with GBE</li> </ul>	intervention	Study type:	
placebo groups.	data.	with aspirin	other	pts with stable IC	
pentoxifylline and	results was generally not possible due to the lack of	<ul> <li>1 study compared pentoxifylline</li> </ul>	placebo or any	walking capacity of	
between the	73.9%Testing for statistical significance of these	with flunarizine	pentoxifylline vs.	improving the	<u>22258961</u>
difference in ABI		<ul> <li>1 study compared pentoxifylline</li> </ul>	RCTs comparing	pentoxifylline in	2015 (178)
statistically significant	pentoxifylline over placebo ranged from 1.2%–155.9%,	<ul> <li>I/ studies compared</li> <li>pentoxifylline with placebo</li> </ul>	Double blind	the efficacy of	al.
There was no	The difference in nercentage improvement in TWD for	<ul> <li>17 etildide commarad</li> </ul>	Inclusion criteria:	<b>Aim:</b> To determine	Salhiwah K et

Study Acronym:	Aim of Study:	Study Acronym: Aim of Study: Patient Population S	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if anv):
Author;	Study Type;		(# patients) /	(Absolute Event Rates,	Study Limitations;
Year Published	Study Size (N)		Study Comparator	P value; OR or RR; &	Adverse Events
			(# patients)	95% CI)	
Villarruz MV, et al.	Aim: To assess the	Inclusion criteria: Pts with PVD,	7 publications	<ul> <li>WMD in ABI: 0.01; 95%</li> </ul>	<ul> <li>Side effects: Faintness: RR: 11.44;</li> </ul>
2008(179)	effects of EDTA	particularly those with IC	representing 5 trials.	Cl: -0.03 – 0.06.	95% CI: 1.51–86.45
<u>12519577</u>	chelation on clinical			<ul> <li>WMD for walking</li> </ul>	<ul> <li>Gastrointestinal symptoms RR: 1.63;</li> </ul>
	outcomes among			distance: -37.93; 95% CI:	95% CI: 0.67–3.99
	people with			-90.32 - 0.06	<ul> <li>Proteinuria RR: 2.60; 95% CI: 0.85–</li> </ul>
	atherosclerotic CV			<ul> <li>WMD for PFWD post-</li> </ul>	7.93
	disease:			treatment: -7.73; 95% CI:	<ul> <li>Hypocalcemia RR: 3.12; 95% CI:</li> </ul>
				-22.59 - 7.13	0.65–14.98
	Study type:				
	Systematic review				
ABI indicates ankle-br	rachial index; EDTA, eth	ABI indicates ankle-brachial index; EDTA, ethylene diamine tetraacetic acid; CI, confidence interval; HR, hazard ratio; IC, intermittent claudication; N/A, not applicable; PFWD, pain free	nce interval; HR, hazard ratio	<ol> <li>IC, intermittent claudication</li> </ol>	N/A not applicable: PFWD, pain free
					i i i i i i i i i i i i i i i i i i i

ABI indicates ankle-brachial index; GBE, ginkgo biloba extract; IC, intermittent claudication; PAD, peripheral artery disease; PFWD, pain free walking distance; PGE1, prostaglandin E1; pt, patient; QoL, quality of life; and TWD, total walking distance.

walking distance; pt, patient; PVD, peripheral vascular disease; RR, relative risk; and WMD, weighted mean difference.

## Evidence Table 28. Nonrandomized Trials, Observational Studies, and/or Registries of Homocysteine Lowering Therapy for Lower Extremity PAD in Patients with Diabetes Mellitus-Section 5.10.1.

	Diabetes Mellin				
Study Acronym;	Study	Patient Population	Primary Endpoint and Results	Summary/Conclusion	
Author;	Type/Design;		(include P value; OR or RR;	Comment(s)	
Year Published	Study Size		& 95% CI)		
Khandanpour N, et al.	Study type:	Inclusion criteria:	1° endpoint: Homocysteine levels in PAD pts vs. controls	<ul> <li>Homocysteine levels are</li> </ul>	
2009 (180)	Meta-analysis of	<ul> <li>Reviewed MEDLINE, EMBASE, and</li> </ul>		elevated among PAD pts as	
19560951	observational	Cochrane databases for studies published	Results:	compared to non-PAD	
	studies and	between 1950—December, 2007	<ul> <li>PAD pts had higher homocysteine levels than non-PAD</li> </ul>	controls	
	clinical trials	<ul> <li>Observational meta-analysis: studies</li> </ul>	controls	<ul> <li>Data lacking to make</li> </ul>	
		with measurement of plasma	<ul> <li>Pooled mean difference vs. controls +4.31 micromol/L</li> </ul>	statement regarding benefit	
	<u>Size:</u>	homocysteine levels in PAD pts and non-	(95% Cl: 1.71–6.31; p<0.0001)	of homocysteine lowering	
	<ul> <li>n=14 studies</li> </ul>	PAD controls	<ul> <li>Mean plasma homocysteine levels higher in PAD pts than</li> </ul>	therapy for clinical benefit in	
	included in meta-	<ul> <li>Clinical trial meta-analysis: Trials for</li> </ul>	in controls in all 14 studies include in meta-analysis, though	PAD	
	analysis (of 214	which PAD pts with treated with single or	magnitude of difference varied across studies		
	retrieved from	combined vitamin therapy (folate, vitamin	<ul> <li>Clinical trial meta-analysis unable to be performed due to</li> </ul>		
	databases)	B6 and/or vitamin B12)	limited study quality and diverse outcomes reported. Among		
		<ul> <li>PAD defined as ABI &lt;0.9, IC, diminished</li> </ul>			
				1	
		reports, homocysteine levels not extractable, non-fasting or post-methionine loading homocysteine levels reported	A	<ul> <li>Studies reported other endpoints including endothelial function testing, inflammatory and other biomarkers</li> </ul>	Jothelial Jers
-----------------------	---------------------------------------------------------	---------------------------------------------------------------------------------------------------------------------------------	----------------------------------	------------------------------------------------------------------------------------------------------------------------------------	--------------------------------------------------------------------------------------------------------
ABI indicates ankle-b	rachial index; CI, confid	ABI indicates ankle-brachial index; CI, confidence interval; IC, intermittent claudication; PAD, peripheral artery disease; and	AD, peripheral artery dis	sease; and pt, patient.	
Evidence Table 2	29. R <u>C</u> Ts Comparin	Evidence Table 29. RCTs Comparing Additional Medical Therapies of Homocysteine Lowering T	^F Homocysteine Lov	wering Therapy for Lov	herapy for Lower Extremity PAD–Section 5.10.1.
Study Acronym;	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Author;	Study Type;		(# patients) /	(Absolute Event Rates,	Study Limitations;
Year Published	study size (N)		study Comparator (# patients)	P value; UK of KK; & 95% CI)	Adverse Events
HOPE-2	Aim: Study effect of	Inclusion criteria:	Intervention: Folic	<u>1° endpoint:</u>	<ul> <li>Homocysteine decreased in interventional</li> </ul>
Lonn E, et al.	vitamin	<ul> <li>Age ≥55 y with documented CAD,</li> </ul>	acid 2.5 mg/vitamin	<ul> <li>No improvement in</li> </ul>	arm and increased in placebo arm (-2.4
2006 (181)	supplementation to	AD, cerebrovascular disease, or DM +	B12 1 mg in a	composite of death from	nicromol/L vs. +0.8 micromol/L)
<u>16531613</u>	levels on risk of	PAD enrollment criteria were prior	combined pill	stroke with intervention	groups (RR: 0.96; 95% CI: 0.81–1.13)
	major CV events	lower extremity revascularization		<ul> <li>Event rates 18.8%</li> </ul>	<ul> <li>No difference in risk of MI between groups</li> </ul>
HUPE-2	among pts with	(bypass or PTA), claudication with ABI	Comparator:	(intervention) vs. 19.8%	(RR: 0.989; 95% CI: 0.85–1.14)
Lonn E, et al.		≥0.% on angiography prior ischemic		(placebo); KK: 0.90; 95% CI: 0 84–1 07:	<ul> <li>Decreased RK stroke among those</li> <li>randomized to intervention /BB: 0.75: 05%</li> </ul>
2006(182)	Study type: Double	limb or foot amputation		p=0.41.	CI: 0.59–0.97).
1045UU17	controlled			<ul> <li>"Average follow-up" 5</li> </ul>	<ul> <li>Increased RR risk of hospitalization with</li> </ul>
	randomized trial	<ul> <li>Use of vitamin supplements with</li> </ul>		У	intervention (RR: 1.24; 95% CI: 1.04–1.49)
	2	significant folic acid content		Safety endpoint: No	<ul> <li>All other secondary outcomes with no</li> </ul>
	<u>oize</u> : ● n=5,522	<ul> <li>Prior adverse reactions to folate/B6/B12</li> </ul>		SAEs related to study treatment.	difference in groups (including VTE, cancer)
	randomized pts with	<ul> <li>Planned cardiac/peripheral vascular</li> </ul>			Summary: Negative study; no overall CV
	PAU	revascularization within 6 mo			benefit of homocysteine lowering therapy in
	<ul> <li>II-133</li> <li>claudication (2.4%)</li> </ul>	<ul> <li>Significant non-</li> <li>atherosclaratio/atherosthrombotic</li> </ul>			Canada Brazil and Europe) which included
	<ul> <li>n=276 with PAD</li> </ul>	cardiovascular disease			a small subset of PAD pts.
	revascularization	<ul> <li>Other non-cardiovascular</li> </ul>			
	(3.0.%)	comorbidities expected to limit			

Exclusion criteria: Lack of non PAD control group, non-English studies, case reports, homocysteine levels not

least major leg artery)

pedal pulses + angiographically demonstrated PAD (obstruction of one at

All 8 studies demonstrated reduction in plasma

homocysteine in folate/vitamin intervention groups

 One study in meta-analysis which reported on ABI and walking distance studied other nutritional supplements not

homocysteine lowering vitamins alone.

8 clinical trials, 3 nonrandomized

Study Acronym;	Aim of Study;	Study Acronym; Aim of Study; Patient Population S	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Author; Year Published	Study Type; Study Size (N)		(# patients) / Study Comparator	(Absolute Event Rates, P value: OR or RR: &	Study Limitations; Adverse Events
	•		(# patients)	95% CI)	
FLUVACS	Aim: To test the effect of	Inclusion criteria: MI pts or	Intervention:	1° endpoint:	Time to first composite triple endpoint of
Gurfinkel EP, et al.	1 yr benefit of influenza	PCI pts	Influenza vaccine	Time to first CVD	CVD, MI, and rehospitalization for severe
2004 (183)	vaccination in pts with		(151)	<ul> <li>At 6 mo: 2% in vaccinated</li> </ul>	recurrent ischemia at 1 yr was significantly
<u>14683739</u>	MI and planned PCI	Exclusion criteria:		intervention group vs. 8% CVD	decreased in the intervention group
		Unstable CAD, prior by-bass	Comparator: No	in unvaccinated controls (RR:	compared to control group (22% in
	Study type: RCT	surgery, angioplasty, or	vaccination ontop of	0.25; 95% CI: 0.07–0.86;	vaccinated intervention group vs. 37% in
		tissue necrosis	standard medication	p=0.01)	unvaccinated control group; RR: 0.59; 95%
	Size: n=301 (200 MI pts		(150)	<ul> <li>At 1 yr: 6% in vaccinated</li> </ul>	Cl: 0.4–0.86; p=0.004)
	and 101 PCI pts)			intervention group vs. 17%	<ul> <li>Reduction in RR of CVD in vaccinated</li> </ul>
				CVD in unvaccinated controls.	group at 1 y.
				(RR: 0.34; 95% CI: 0.17–0.71;	No PAD specific evidence identified
	<b>Δim</b> . Determine effects	Inclusion criteria	Intervention.	1º andnoint:	3º andmint:
Ciszewski A, et al.	of influenza vaccination	• Age 30-80 v	Influenza vaccine	1 vr CVD	<ul> <li>No difference between two arouns for CVD.</li> </ul>
2008 (184)	on coronary events in	<ul> <li>CAD confirmed by</li> </ul>	(325)	• At 1 y: HR: 1.06; 95% CI:	acute MI, or coronary revascularization
<u>18187561</u>	pts with CAD	angiography with ≥50%		0.15–7.56; p=0.95	<ul> <li>At 1 y coronary ischemic events was</li> </ul>
		stenosis of ≥1 large	Comparator:		decreased in intervention group compared to
	<u>Study type</u> : RCT	epicardial coronary artery	Placebo (333)		placebo control group (HR: 0.54; 95% CI: 0.29–0.99: p=0.047)
	Size: n=658 treated	Exclusion criteria:			
	CAD pts (477 men)	Congestive heart failure			Limitations: Small sample size, effect of flu
		<ul> <li>Planned CV surgery within</li> </ul>			selection hise
		6mo			
		<ul> <li>Evolving renal failure</li> </ul>			<ul> <li>No PAD specific evidence identified</li> </ul>
		<ul> <li>Neoplastic disease</li> </ul>			
		<ul> <li>Psycho-organic disorder or</li> </ul>			
		any factor impeding follow-			
		dn			

CAD indicates coronary artery disease; CI, confidence interval; CV, cardiovascular; HOPE, Heart Outcomes Prevention Evaluation; PAD, periphery artery disease; PTA, percutaneous transluminal angioplasty; pt, patient; RCT, randomized controlled trial; RR, relative risk; SAE, serious adverse event; US, United States; and VTE, venous thromboembolism.

compliance or ability to complete study

<u>∞</u>

CAD indicates coronary infarction; NYHA, New Y	artery disease; Cl, co 'ork Heart Associatior	nfidence interval; CVD, cardio ; PCI, percutaneous interventi	CAD indicates coronary artery disease; Cl, confidence interval; CVD, cardiovascular death; CVD, cardiovascular disease; H infarction; NYHA, New York Heart Association; PCI, percutaneous intervention; pt, patient; RCT, randomized controlled trial	CAD indicates coronary artery disease; Cl, confidence interval; CVD, cardiovascular death; CVD, cardiovascular disease; HR, hazard ratio; ITT, ir infarction; NYHA, New York Heart Association; PCI, percutaneous intervention; pt, patient; RCT, randomized controlled trial; and RR, relative risk	IR, hazard ratio; ITT, intention to treat; MI, myocardial ; and RR, relative risk.
Evidence Table 31.	Nonrandomized	Trials for Influenza Vac	Evidence Table 31. Nonrandomized Trials for Influenza Vaccination–Section 5.10.2.		
Study Acronym; Author; Year Published	Study Type/Design; Study Size	sign; Patient Population	tion Primary I (include	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Summary/Conclusion Comment(s)
Davis MM, et al.	Study type: Science		, ,	<ul> <li>COR I LOE B recommendation to immunize with</li> </ul>	Not recommended for persons with CV
2006 (185) 17010820	Advisory Statement	t case control studies and RCTs		inactivated vaccine as part of comprehensive secondary prevention in persons with coronary	conditions to be immunized with live, attenuated vaccine.
	<u>Size</u> : N/A	Exclusion criteria: N/A	-	<ul> <li>1 RCT (FLUVACS) included</li> </ul>	<ul> <li>Immunization coverage levels are below national goals</li> </ul>
			<ul> <li>Summary of obsi control studies der</li> </ul>	<ul> <li>Summary of observational cohort and case control studies demonstrating reduced CV event</li> </ul>	
			rates among pts w	rates among pts with cardiovascular disease who	
)) 	2		received influenza vaccination	vaccination	
COR indicates class of r controlled trial. Evidence Table 32.	ecommendation; CV,	cardiovascular; LOE, level of o	received influenza vidence; N/A, not applicable; F	PCI, percutaneous coronary interv	COR indicates class of recommendation; CV, cardiovascular; LOE, level of evidence; N/A, not applicable; PCI, percutaneous coronary intervention; pt, patient; and RCT, randomized controlled trial.         Evidence Table 32. RCTs for Exercise Therapy–Section 6.
COR indicates class of recommendation; CV, car controlled trial. Evidence Table 32. RCTs for Exercise Study Acronym; Aim of Study; Author; Study Type;	ecommendation; CV, RCTs for Exerci Aim of Study; Study Type;	cardiovascular; LOE, level of o se Therapy–Section 6. Patient Population	received influenza vidence; N/A, not applicable; F Study Intervention (# patients) /	PCI, percutaneous coronary interv CI, percutaneous coronary interv (Absolute Event Rates, P	antion; pt, patient; and RCT, randomized Relevant 2° Endpoint (if any); Study Limitations;
COR indicates class of r controlled trial. Evidence Table 32. Study Acronym; Author; Year Published	ecommendation; CV, RCTs for Exerci Aim of Study; Study Type; Study Size (N)	cardiovascular; LOE, level of o se Therapy–Section 6. Patient Population	received influenza vidence; N/A, not applicable; F Study Intervention (# patients) / Study Comparator (# patients)	² Cl, percutaneous coronary interv <b>Endpoint Results</b> (Absolute Event Rates, P value; OR or RR; & 95% Cl)	antion; pt, patient; and RCT, randomized Relevant 2° Endpoint (if any); Study Limitations; Adverse Events
COR indicates class of r controlled trial. Evidence Table 32. Study Acronym; Author; Year Published CLEVER 18 mo F/U	ecommendation; CV, RCTs for Exerci: Aim of Study; Study Type; Study Size (N) <u>Aim</u> : Report the longer-term (18	cardiovascular; LOE, level of o se Therapy–Section 6. Patient Population Inclusion criteria: • Age >40 v	received influenza vidence; N/A, not applicable; F Study Intervention (# patients) / Study Comparator (# patients) Intervention: OMC, n=22; SE, n=44; ST, n=46. SE	PCI, percutaneous coronary interv CI, percutaneous coronary interv Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI) <u>1° endpoint</u> : PWT improved from baseline to 18 mo for both	<ul> <li>At 18 mo, improvement in disease- specific scales (WIQ, PAQ) was</li> </ul>
COR indicates class of r controlled trial. Evidence Table 32. Study Acronym; Author; Year Published CLEVER 18 mo F/U Murphy TP, et al.	ecommendation; CV, RCTs for Exerci: Aim of Study; Study Type; Study Size (N) Aim: Report the longer-term (18 mo) efficacy of SE	cardiovascular; LOE, level of o se Therapy–Section 6. Patient Population Patient Population Inclusion criteria: • Age >40 y • oderate to severe IC due	received influenza vidence; N/A, not applicable; F Study Intervention (# patients) / Study Comparator (# patients) Intervention: OMC, n=22; SE, n=44; ST, n=46. SE was supervised for 26 wk, 3	PCI, percutaneous coronary interver CI, percutaneous coronary interver Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI) <u>1° endpoint</u> : PWT improved from baseline to 18 mo for both SE (5±5.4 min) and ST (3.2±4.7	antion; pt, patient; and RCT, randomized         Relevant 2° Endpoint (if any);         Study Limitations;         Adverse Events         • At 18 mo, improvement in disease-         specific scales (WIQ, PAQ) was         statistically superior for ST and SE
COR indicates class of r controlled trial. Evidence Table 32. Study Acronym; Author; Year Published CLEVER 18 mo F/U Murphy TP, et al. 2015 (186) 25766947	ecommendation; CV, RCTs for Exerci: Aim of Study; Study Type; Study Size (N) <u>Aim</u> : Report the longer-term (18 mo) efficacy of SE compared with ST and OMC included	cardiovascular; LOE, level of a se Therapy–Section 6. Patient Population Patient Population Inclusion criteria: • Age >40 y • oderate to severe IC due to aortoiliac PAD. IC defined as ability to walk ≥2	received influenza vidence; N/A, not applicable; F Study Intervention (# patients) / Study Comparator (# patients) Intervention: OMC, n=22; SE, n=44; ST, n=46. SE was supervised for 26 wk, 3 times/wk, 1 h for 6 mo followed by a telephone	CI, percutaneous coronary interv         CI, percutaneous coronary interv         Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% CI)         1° endpoint: PWT improved from baseline to 18 mo for both SE (5±5.4 min) and ST (3.2±4 min) more than OMC (0.2±2.1 min); p<0.001 and p<0.04,	<ul> <li>At 18 mo, improvement in disease-specific scales (WIQ, PAQ) was statistically superior for ST and SE compared with OMC, but ST and SE differed significantly from each other</li> </ul>
COR indicates class of r controlled trial. Evidence Table 32. Study Acronym; Author; Year Published F/U Murphy TP, et al. 2015 (186) 25766947	ecommendation; CV, RCTs for Exerci: Aim of Study; Study Type; Study Size (N) <u>Aim</u> : Report the longer-term (18 mo) efficacy of SE compared with ST and OMC included printed advice about exercise and	cardiovascular; LOE, level of a se Therapy–Section 6. Patient Population Patient Population Inclusion criteria: • Age >40 y • oderate to severe IC due to aortoiliac PAD. IC defined as ability to walk ≥2 min on TM at 2 miles/hr at 0% grade but <11 min	received influenza vidence; N/A, not applicable; F Study Intervention (# patients) / Study Comparator (# patients) Intervention: OMC, n=22; SE, n=44; ST, n=46. SE was supervised for 26 wk, 3 times/wk, 1 h for 6 mo followed by a telephone maintenance program through 18 mo during	Vaccination         CI, percutaneous coronary interve         Endpoint Results         (Absolute Event Rates, P         value; OR or RR; &         95% CI)         1° endpoint: PWT improved         from baseline to 18 mo for both         SE (5±5.4 min) and ST (3.2±4.1 min))         more than OMC (0.2±2.1 min); p<0.001 and p<0.04,	<ul> <li>At 18 mo, improvement in disease-specific scales (WIQ, PAQ) was statistically superior for ST and SE differed significantly for PAQ symptoms, PAO treatment satisfaction PAO Cod</li> </ul>

Contraindication to vaccination

GOALSAim: DetermineMcDermott MM, etwhether a home-al.based walking2013 (17)exercise program23821089using a group-mediated coontilve	OMC. Study type: Long- term follow-up of RCT RCT Murphy TP, et al. 2012 (187) 22090168 Size: n=79 of 111 pts initially enrolled assessments at 18 mo. 2012 (187) Walking outcomes and ST on both claudication due to aortoiliac PAD. Size: n=111 pts
è -	
Inclusion criteria: Resting ABI <0.9 or ABI between 0.91–1 with a 20% drop after a heel-rise test or medical evidence of LE revascularization or	<ul> <li>Stenosis of distal aorta or iliac arteries.</li> <li>Exclusion criteria: CLI or 2 comorbid conditions that limited walking ability.</li> <li>Age &gt;40 y</li> <li>Moderate to severe IC due to aortoiliac PAD. IC defined as ability to walk at least 2 min on TM at 2 miles/hr at 0% grade but &lt;11 min (about 5.5 METS maximum).</li> <li>≥50% stenosis of distal aorta or iliac arteries.</li> <li>Exclusion criteria: CLI or 2 comorbid conditions that limited walking ability.</li> </ul>
Intervention: Walking on- ground (not TM) progressing to 50 min 5 times/wk for 6 mo. For pts with IC, walk to pain level 4 of 5, rest, and resume. For	<u>Comparator</u> : N/A <u>Intervention</u> : OMC, n=22; SE, n=44; ST, n=46. SE was supervised for 26 wk, 3 sT/SE group was dropped after 8 pt to enhance enrollment in the other groups. Randomization ratio was 2:2:1 (ST:SE:OMC). <u>Comparator</u> : N/A
1° endpoint: Exercisers increased their 6 min walk distance (357.4–399.8 meters vs. 353.3–342.2 meters for those in the control group; mean difference: 53.5; 95% CI: 33.2–	adverse events occurred in first 6 mo and not in the follow-up. These included an MI in the OMC group; 1 death in SE group; and 1 target limb revascularization in the ST group. 12±2.6 min with OMC alone, 5.8±4.6 min with ST. Compared with OMC alone, SE led to a greater mean improvement in PWT by 4.6 min (95% CI: 2.7– 6.5; p<0.001), whereas ST had a somewhat smaller relative improvement in PWT of 2.5 min (95% CI: 0.6–4.4; p=0.022). A direct comparison of SE and ST showed a greater improvement in PWT of 2.5 min (95% CI: 0.6–4.4; p=0.022). A direct comparison of SE and ST showed a greater improvement in PWT with SE by a mean of 2.1 min (95% CI: 0.0– 4.2; p=0.04) Safety endpoint: 4 SAEs within 30 d of ST. SAEs noted in the 18 mo follow-up report that said they occurred in the first 6 mo were not mentioned.
<ul> <li>Maximal TM walking time (intervention, 7.91–9.44 min vs. control, 7.56–8.09; mean difference: 1.01 min; 95% CI: 0.07–1.95; p=0.04), accelerometer- measured physical activity over 7 ds (intervention, 778.0–866.1 vs. control,</li> </ul>	<ul> <li>the stented pts and changed by 0.00±0.1</li> <li>for OMC, 0.2±0.2 for ST, and 0.00±0.1</li> <li>for SE (p=0.002 for ST vs. OMC and p&lt;0.001 for ST vs. SE)</li> <li>SE had the advantage of improved limb muscle strength, walking efficiency, and performance.</li> <li>ABI improved by 0.29±0.33 in the ST group (p&lt;0.0001) only.</li> <li>The greatest improvements in self-reported QoL were observed in the ST cohort despite greater increases in PWT in the SE group.</li> </ul>

SF-30 IIIG eased by 3.2 (stailuard error.	וש.ט ווופופו ג, ט–ט.טט.		Exclusion criteria:	מטווונץ מווע שטב ווו	
mental nealth WoL subscale score of the	distance average: 39.2; SE:	Exercisers participated in a			
		Expression participated in a	• Type T or 2 UM		
(standard arror: 2.8) unite (n=0.02); the	we mayimal tradmill walking	DM and CVD risk factors		intervention to	
2.2) units and doornoond by 1.0	average: 24 E: CE: 10 E motoro	management behaviors for		walking	
scores increased by 5.7 (standard error:	treadmill walking distance	about PAD and self-	● With PAD or prior surgery	home-based	21873560
respectively, average walking speed		groups received education	● Age ≥40 v	the efficacy of a	2011 (189)
<ul> <li>For the exercise and control groups,</li> </ul>	1° endpoint: The groups did not	Intervention: All pts in both	Inclusion criteria:	Aim: Determine	Collins TC, et al.
			unsupervised exercise	follow-up testing	
			that would preclude	168 completed	
				at 6 mo. At 12 mo.	
		general health topics.	- Other major comercialities	completed testing	
		received calls related to	<ul> <li>Surgery within past 3 mo</li> </ul>	which 178	
		Comparator: Controls	than PAD	enrolled 194 pts, of	
			<ul> <li>Walking impairment other</li> </ul>	Size: Initial study	
		study team.	<ul> <li>Walking aid except cane</li> </ul>		
	events reported	which were sent back to	<ul> <li>Inability to walk 50 ft</li> </ul>	Study type: RCT	
	Safety endpoint: No adverse	exercise and keep logs,	<ul> <li>Wheelchair confinement</li> </ul>		
		and were encouraged to	<ul> <li>LE amputation</li> </ul>	mo intervention	
	14.6–53.5; p<0.001)	from their group facilitator	Exclusion criteria:	completing the 6	
<ul> <li>No adverse events reported</li> </ul>	difference: +34.1 m; 95% CI:	exercisers received call		12 mo, 6 mo atter	
(p=0.139).	in the control group; mean	During the next 6 mo,	evidence of PAD.	a tollow-up study at	
different between groups at 12 mo	intervention vs. 353.1–345.6 m	behavioral techniques.	revascularization or	without IC. This is	
in the WIQ distance score was not	(trom 355.4–381.9 m in the	mediated cognitive	medical evidence of LE	PAD with and	24850615
95% CI: +1.6 - +16.1; p=0.018). Change	baseline to 12 mo tollow-up,	which included group-	atter a neel-rise test or	controis in pts with	2014 (188)
the control group; mean difference: +8.8;	their 6 min walk distance from	weekly group sessions,	0.91–1 with a 20% drop	walking vs.	al.
36.1–46.5 in exercisers vs. 34.9–36.5 in	controls, exercisers increased	phase, exercisers attended	ABI ≤U.9 or ABI between	intervention of	McDermott MM, et
<ul> <li>WIQ speed score increased (from</li> </ul>	1° endpoint: Compared to	Intervention: During 6 mo	Inclusion criteria: Resting	Aim: 6 mo	GOALS
study related.					
CVD with pneumonia, all considered not		topics.	unsupervised exercise	Size: n=194 pts	
and 2 deaths from hypertensive CVD and		discuss general health	that would preclude		
<ul> <li>1 death from cancer among exercisers</li> </ul>		met weekly for 60 min to	other major comorbidities	Study type: RCT	
Cl: 3.4–17.4; p=0.004).		education control group that	surgery within past 3 mo,		
35.3–36.6; mean difference: 10.4; 95%		Comparator: Health	impairment other than PAD,	without IC.	
score (intervention: 36.1–47.7 vs. control:	after recovery.		except cane, walking	with PAD with and	
3.9–18.1; p=0.003), and WIQ speed	CABG and completed study	met once a wk for 90 min.	walk 50 ft, walking aid	control group in pts	
34.4; mean difference: 11.1; 95% CI:	and subsequently required	approach, exercisers also	confinement, inability to	compared with a	
(intervention, 35.3–47.4 vs. control, 33.3–	developed dyspnea on exertion	cognitive behavioral	amputation, wheelchair	performance	
p=0.03), WIQ distance score	Safety endpoint: 1 exerciser	Using a group-mediated	Exclusion criteria: LE	improve functional	
activity units; 95% CI: 12.82–216.5;		14 on Borg RPE scale.		approach, can	
671.6–645.0; mean difference: 114.7	73.8; p<0.001.	pts without IC, walk at 12-	evidence of PAD.	behavioral	

07					
climbing ability were similarly maintained	ability (p=0.011) vs. controls	total time exercise time of		training on	
Improvements in walking speed and stair	(p=0.023) and stair climbing	uptake, 2 d/wk for 24 wk,	<ul> <li>Symptoms 12 mo</li> </ul>	aerobic exercise	
better maintained in upper limb group.		specific maximal oxygen	● ABI ≤0.9	and lower-limb	21215558
perceptions of walking distance were	improvements in the perceived	at 85%–90% of limb-	<ul> <li>PAD with IC by Hx</li> </ul>	effects of upper-	2011 (191)
<ul> <li>At 48 and 72 wk, improvement in</li> </ul>	1° endpoint: After 6 wk,	Intervention: Arm cranking	Inclusion criteria:	Aim: Compare the	Saxton JM, et al.
		Comparator: Non-exercise, usual care control			
		walking.			
		monitors to measure			
		aroups used step activity			
		resume evercise Roth		alze: II-II bis	
		exercise test, to near-		<b>9</b>	
		workload from baseline	liver disease	Study type: RCT	
	effects.	set at 40% of peak	(active cancer, renal, or		
	too few to ascertain group	based program. Intensity	causes, major comorbidities	control.	
	hernia surgery (1). These were	min shorter than home-	exercise limited by other	and usual-care	
	revascularization (2), MI (1), and	a treadmill with durations 5	within 3 mo before study,	exercise program,	
	Events included stroke (2), leg	program was performed on	or pentoxifylline initiated	supervised	
	adverse events in either group.	program. Supervised	asx PAD, use of cliostazol	program, a	
	but though no unanticipated	during final 2 wk of	noncompressible vessels,	based exercise	
	Safety endpoint: Not specified	20 min initially to 45 min	to obtain ABI due to	IC after a home-	
	G	duration progressed from	Exclusion criteria: Inability	activity in PAD with	
	did not change.	selected pace. Walking		daily ambulatory	
	walking time (p<0.01). Controls	maximal pain 3 d/wk at self-	exercise	performance and	
	onset time (p<0.001) and peak	intermittent walking to near-	ABI ≤0.9 or ABI ≤.73 after	exercise	21262997
<ul> <li>Home group only increased daily average cadence (n&lt;0.01)</li> </ul>	<u>1° endpoint:</u> Both exercise	Intervention: 12 wk. Home-based exercise of	Inclusion criteria: Exertional leg pain, resting	Aim: Compare	Gardner AW, et al. 2011 (190)
			program	:	
	group eπects.		unsupervised exercise		
	these were too few to ascertain		preclude participation in		
	complete treadmill testing but	behaviors	Comorbidities that would		
	foot problems, and unable to	to discuss their health	<ul> <li>MI within nast 3 mo</li> </ul>		
	leg bypass surgery, broken hip,	received twice monthly calls	o re revascularization in	Size: n=145 pts	
	included general health issues,	Comparator: Controls	● I E revecularization in		
	either group. Some events			Study type: RCT	
	inanticinated adverse events in	ner wk for 6 mo			
error: 1.5) units (n=0.01)	Safety endnoint: No	4 aroun exercise session	No intention to exercise	and PAD	

		given usual care walking			
		supervised exercise but			
		receive participate in			
		usual care group did not			
		ergometry and walking. A			
		performed both arm			
		combination group			
	with 1 study unrelated injury.	to stop due to IC. A			
	Safety endpoint: Not specified	walk 8 min without having			
		increased when pt could			
	not reach statistical significance.	exercise. Workload			
	p=0.107 vs. control) groups did	4/5 claudication, rest,			
Hg).	control) and combination (+60%;	<u>Comparator</u> : IM walking to	3 mo		
This was maintained at 24 wk (-11 mm	treadmill (+54%; p=0.196 vs.		or LE revascularization past		
in arm group (-17 mm Hg) vs. controls.	control). Change in PFWD in	bouts	health conditions, coronary	Size: n=41 pts	
<ul> <li>Resting SBP was lower after 12 wk on</li> </ul>	group (+82%; p=0.025 vs.	workload and exercise	exercise limited by other	2	
vs. control)	increased in the arm-ergometry	12 wk by increasing	Uncontrolled HBP, CLI,	Study type: RCI	
after a 12 wk follow-up (+123%; p=0.011	pain free walking distance	increase of exercise over	Exclusion criteria:		
maintained in the arm-ergometry group	(p<0.002 vs. control). The 12 wk	min for 60 min. Progressive	I	with IC	
control. PFWD improvement was	combination (+68%) groups	exercise for 2 min, rest for 2	treadmill walking,	usual care in PAD	
combination group declined (p=0.751) vs.	treadmill (+69%), and	baseline test. 3d/wk,	IN ABI OT ≥10% atter	treadmill walking or	6001,0061.
(p=0.019) groups, whereas the	the arm-ergometry (+53%),	below maximal during	claudication, ABI ≤0.9, drop	arm-ergometry vs.	
arm-ergometry (p=0.009) and treadmill	walking distance increased in	ergometry at one work level		effects of aeropic	et al.
<ul> <li>24 wk MWD was maintained in the</li> </ul>	1° endpoint: 12 wk maximal	Intervention: Arm-	Inclusion criteria:	Aim: Compare	Treat-Jacobson D,
	group effects.				
	These were too few to ascertain				
	adverse events in either group.				
	but though no unanticipated				
	Safety endpoint: Not specified				
	9 - F 4F				
	the lower limb aroup (p<0.05).				
	WID domains were improved in				
	controls ( $p \le 0.05$ ), and 3 of the 4		if using long term	Size: n=104 pts	
	the upper limb group vie	oversise control group	<ul> <li>Medications for IC except</li> </ul>	<u>otudy type</u> : RCT	
	(p=0.010). Atter 24 wk, all 4	for any same parameters as	<ul> <li>Severe arthritis</li> </ul>	Chindry tripper DOT	
		comparator: Leg cycling	• SOB		
	observed in the lever limb array		Exercise limiting anglina	ו הומופע עטב ווו טוא	
in both exercise groups vs. control.	The general health domain of the	min rest	past 12 mo	related Ool in hts	
	group, and an improvement in		Revascularization with		
III buti exercise groups vs. curitiois.	aroun and an improvement in	zo IIIII III 40 IIIII session. z	Devecularization with	functional status	
Late and an and a second and a	4mil 2022. 241 4: 62. 2024	a action of a line of a line of	Fuchalas subsula.		

 $\ensuremath{\textcircled{O}}$  American Heart Association, Inc. and American College of Cardiology Foundation

	total number of endovascular and surgical interventions				
	after SET (p=0.001). Yet, the				
(HR: 1.35; 95 % CI: 0.67–2.70; p=0.402)	secondary intervention rate,			Size: n=151 pts	
y was 68% with SET and 74% with ER.	treatments. Except in the			2	
<ul> <li>The cumulative survival probability for 7</li> </ul>	differences between the two	needed		Study type: RCT	
(p<0.001)	comparison showed no	angioplasty and stenting as			
remained higher after ER, 121 vs. 64	both SET and ER. Long-term	revascularization with initial		IC.	
interventions (primary and secondary)	(p≤0.005) had improved after	Comparator: Endovascular		strategy in pts with	
number of endovascular and surgical	distance (p<0.001) and QoL		Exclusion criteria: N/A	first treatment	
73% with ER (p=0.001). Yet, the total	distance and pain free walking	walk at home.		SET-first or an ER-	23842830
significantly lower after SET, 47% vs.	consisting of maximal walking	min, 2 d/wk, and 3 d/wk	femoropopliteal disease.	effectiveness of a	2013 (194)
<ul> <li>The portion of pts not needing secondary intervention rate, was</li> </ul>	<u>1° enapoint</u> : Atter / y, functional performance	supervised TM exercise, 30	IC with iliac and	long-term clinical	Fakhry F, et al.
! : :		pattern as the MT group.			
	related to SET in either group.	walking tollowing the same			
	none were reported as being	rest, and then resumed			
	Among 8 dropouts/withdrawal,	onset of pain, stopped to			
	Safety endpoint: Not specified.	The PFT walked until initial			
		Comparator: Titled PFT.			
	(p<0.01) in the PFT group.				
	the MT group and by 36%	accomplished.	3.2 km/hr.		
	increased by 56% (p<0.001) in	total of 60 min was	disease, unable to walk at	Size: n=60 pts	
	by flow-mediated dilation	by 5 min each 2 wk until a	kidney, liver, or joint		
	Endothelial function assessed	for 35 min and progressed	status due to cardiac, lung,	Study type: RCT	
		session was done initially	angina, impaired function		
	respectively.	until pain abated. The	event in prior 1 y, unstable	IC	
	compared to the PFT,	to moderate pain and rest	Exclusion criteria: CVD	walking in pts with	
	(p<0.001) in the MT group as	intermittent bouts of walking		vs. pain-free	
the program in both groups (p>0.05).	vs. 93%	Walking done with	IC pain.	claudication pain	
and BMI did not change significantly after	and PFWT by 120% (p<0.001)	induced IC within 3–5 min.	not taking medications for	moderate	
SET in either group. The smoking status	(p<0.001) vs. 98% (p<0.001),	km/hr and grade that	medical therapy for 6 mo,	performed to	<u>23117015</u>
hs-CRP and fibrinogen were seen after	MWD was prolonged by 100%	SET, 3 times/wk at 3.2	50–75 y with IC, stable	3 mo of SET	2013 (193)
<ul> <li>No significant changes in the levels of</li> </ul>	1° endpoint: Post-training	Intervention: Titled MT.	Inclusion Criteria: Age	Aim: To compare	Mika P, et al.
		guidelines.			

<b>ERASE</b> Fakhry F, et al. <u>26547465</u>	Mazari FA, et al. 2010 (195) <u>19762206</u>
<u>Aim</u> : To assess the 1 y effectiveness of combination therapy of ER + SET or SET alone in pts with IC <u>Study type</u> : RCT <u>Size</u> : n=212 pts	<u>Aim</u> : To compare the 3 mo effects of PTA, SET, and PTA + SET for the treatment of femoropopliteal disease in pts with IC <u>Study type</u> : RCT <u>Size</u> : n=178 pts
Inclusion criteria: ABI <0.9 or decrease >0.15 with exercise, 1 or more vascular stenosis at the aortoiliac or femropoliteral level or both, and impaired MWD. Exclusion Criteria: Not a candidate for revascularization or prior treatment for the target lesions, limited life expectancy; limited ambulation due to causes other than IC.	Inclusion criteria: Stable IC and suitable for PTA for femoropopliteal lesions after 3 mo of optimal medical therapy for CVD risk factors and DM. Exclusion criteria: CLI, severe systemic disease, inability to tolerate treadmill testing, significant cardiac ischemia; revascularization in prior 6 mo
Intervention: Combination therapy of ER + SET. For ER, a stent was used only if the initial balloon angioplasty was not successful. SET was started 2-4 wk after ER. SET consisted primarily of intermittent bouts of treadmill walking to near- maximum claudication pain. Frequency of 2-3 sessions for 30–45 min for initial 3 mo followed by at least 1 session per wk between mo 3–6 and then 1 session per 4 wk until 1 y.	Intervention: SET, 3 times/wk for 12 wk, consisting of circuit training that included stepping, heel raises, leg press, exercise cycle, knee extension, and elbow flexion. PTA consisting of balloon angioplasty and no stenting. <u>Comparator</u> : Combined PTA + SET.
<ul> <li><u>1° endpoint:</u> After 1 y, MWD increased in both groups (p&lt;0.001) with a greater improvement in the combined therapy group (p&lt;0.001)</li> <li><u>Safety endpoint</u>: See secondary outcomes. No study specific AE's discussed.</li> </ul>	(primary and secondary) remained higher after ER (p<0.001) Safety endpoint: See secondary outcomes demonstrated significant clinical (pt reported walking distance, MWD, PFWD, rest and post- exercise ABI) and QoL improvements (p<0.05). Combined therapy produced greater improvement in clinical outcomes than PTA or SET alone (p<0.05) but not in QoL measures. Safety endpoint: See secondary outcomes. No study specific adverse events reported.
<ul> <li>After 1 y, PFWD increased in both groups (p&lt;0.001) with a greater improvement in the combined therapy groups (p&lt;0.001). Similarly, ABI at rest and after exercise showed significantly greater improvement in the combination therapy group. Also, measures of health-related QoL improved in both groups with greater improvements with combined therapy.</li> <li>A higher proportion of pts without an additional intervention in the combination group (92%) vs. the SET alone (77%), HR: 3.2; 95% CI: 1.1–9.2; p=0.005.</li> </ul>	<ul> <li>21 pts (7%) withdrew, of whom 8 were in the SET group, 3 were in the PTA group, and 10 were in the combined group. 11 pts who had PTA had restenosis but none required revascularization.</li> </ul>

during exercise (p=0.025), large-artery elasticity index (p=0.012), and high- sensitivity C-reactive protein (p=0.041) were also significantly different among the 3 groups. Both walking groups improved time to minimum StO ₂ . Only the NEXT Step home group had	scores for COT (p<0.001), PWT (p<0.001), 6 min walk distance (p=0.028), daily average cadence (p=0.011) were different among the 3 groups, with both walking programs showing changes in these	3 mo of intermittent walking (NEXT STEP) o mild-to- moderate claudication pain 3 d/wk, progressing from 20–45 min/session. Pts used step monitor during each session. Exercise logs	by Hx of ambulatory leg pain or pain confirmed by treadmill exercise or ABI ≤0.90 at rest or ≤0.73 after exercise. Exclusion criteria: ABI	the 12 wk effects of exercise training using a step watch home-exercise program, a supervised exercise program,	2014 (198) 25237048
	Safety endpoint: Not specified. 2 exercisers and 1 control dropped for progression of PAD, 3 exercisers dropped for non- specified medical reasons in first 12 wk.			A im	
	Questionnaire scores in the exercise group were considerably better than those in the control group, 27.94±19.83 vs. 38.54±24.26 (p=0.058), reflecting improved QoL and maintenance of benefits.	resistance or time. <u>Comparator</u> : Usual care, general advice about exercise and smoking cessation, ABI measurement			
	improvement in the exercise group (p=0.066) and a trend towards improved functional capacity (WIQ Stair-climbing p=0.093) in the exercise group, with an increase of 8.55 points in the exercise group and a decrease of 13.42 points in the control group. At 1 v. IC	equipment including treadmill, stepper, elliptical trainer, recumbent cycle, and arm cycle. Intensity of 70%–80% of exercise test maximum HR. On treadmill, walking to leg pain of 3 of 4, rest, and resume walking. Exercise intensity	Hg post-exercise <u>Exclusion criteria:</u> Comorbidities which precluded participation in exercise, MI past 6 mo, acute onset or within one mo of IC, lower limb revascularization past 6 mo	exercise program on functional capacity and QoL for PAD pts <u>Study type</u> : RCT <u>Size</u> : n=44 pts	
N/A	<u>1° endpoint</u> : At 12 wk, there was a trend towards improved QoL in both groups, with a tendency for greater	Intervention: 2 d/wk supervised exercise for 12 wk. 30–40 min of aerobic exercise using a range of	Inclusion criteria: Fontaine Stage II, ABI <0.9 at rest or a decrease of ankle pressure by ≥15 mm	<u>Aim</u> : To assess the 1 y effects of participation in a 12 wk supervised	Guidon M and McGee H 2013 (197) 22804715
		Comparator: SET alone.			

<ul> <li>No exercise-related adverse events.</li> </ul>	Incremental upper- and lower- limb ergometry tests (p<0.001). PFWD and MWD improved in	exercise followed by 2 min	Claudication of >12 mo or revascularization in	training on walking	
SF-36 QoL questionnaire.	power generated during the	groups 2 d/wk for 6 wk.	Evolution onitorio.	limb (arm cranking)	10753273
<ul> <li>Improvements in physical function and role-limitation-physical domains of the</li> </ul>	<u>1° endpoint</u> : Both training groups improved the maximum	Intervention: An upper- limb and lower limb training	Inclusion criteria: Moderate to severe IC	<u>Aim</u> : To compare effects of upper	Walker RD, et al. 2000 (200)
	Safety endpoint: N/A				
	trained group only.				
	Impairment Questionair		limiting exercise		
	(p<0.022) on the Walking		warafin sodium, or	<u>Size</u> : n=52 pts	
	and walking speed scores		current use of vitamin E,	}	
	perceived distance (p<0.001)		revascularization in past y,	Study type: RCT	
	polestriding training program. pt		ulceration, resting ABI <0.4,		
	significantly less after	control	leg pain at rest, ischemic	PAD.	
	claudication pain were	Comparator: Nonexercise	Exclusion criteria: Severe	pain caused by	
	(p<0.001). Perceived	OTICE DEL WIX TOT T WIX.		of persons with IC	
	incremental treadmill tests	and per which own,		exercise tolerance	00112021
postexercise Abi	the constant work-rate and	wk twice ner wk for 8 wk	limiting factor to maximal	exercise increases	12002 (199) 12021703
2° endpoint: No changes in resting or	<u>1° endpoint</u> : Polestriding	Intervention: Polestriding	Inclusion criteria: Pain	Aim: To determine	Langbein WE, et al.
		their visits.			
		monitor to quantify time of			
		pts also wore a step			
		resistance exercise. These			
		walk but performed various			
		exercise aroup that did not			
	walking group.	control. light resistance			
	revascularization in home-based	was also an attention-			
	supervised exercise group: 1 leg	home-based group There		Size: n=180 nte	
		activity monitor tollowing		Study type: RC1	
	Safety endpoint: Not specified.	exercise while wearing step			
		Comparator: Supervised		outcomes.	
	based group (p<0.05).			physiological	
	was greater than the home-	exercise sessions.	comorbidities.	selected	
	in the supervised exercise aroup	aiven to auide subseauent	symptoms, other serious	walking time and	
in I AFL and hs-CRP (n<0.05)	haseline The change for PWT	staff and feedback was	∠ I.40; asx FAD; medications for PAD	control group on	
The second state for the second state in the second state is the s	- House and a second state from the second	the mandator of her shealth			

<ul> <li>Community walking programs may improve with</li> </ul>	Results: Supervised exercise programs were	<ul> <li>English language</li> </ul>	Size: n=10 RC1s	
efficacious.		Cochrane databases	2	24103409
PAD to exercise in the community are not	the treadmill.		Ieview	2013(202)
Unstructured recommendations for pts with sx	1° endpoint: Peak walking performance on	Inclusion criteria:	<b>Study type:</b> Literature	Mays KJ, et al.
- I had a second and a second a stand for the state of	10 and no internet model and the model and t	Inclusion exiterio	Study time literation	Meyo DI of ol
	"tiptoe standing" by /0.5% (group A) and 113 7% (12 mo: p<0.05)			
	by 64.2% (6 mo) and 75.3% (12 mo), and for			
instructed to walk on the weekends on their own.	90.0% (6 mo) and 90.2% (12 mo), for "pulling"			
mimic walking in a community setting. Pts were also	groups showing an increase for "pushing" by			
done on a track in a gym rather than treadmill to	parameters increased significantly in both the			
<ul> <li>Though the program was supervised, walking was</li> </ul>	p<0.001) was seen at 12 vs. 6 mo. All strength		6 mo, n=52 for 12 mo)	
exercise	increase in walking speed (12.1% vs. 5.3%;,	only aorto-illiac stenosis	Size: n=94 pts (n=42 for	
<ul> <li>Strength exercise involved lower extremity</li> </ul>	29.5%, respectively, at 6 and 12 mo a greater	or orthopedic conditions,		
decreased by -9.4 mg/dL in 12 mo group (p=0.0053)	distance increased similarly by 27.5% and	6, exercise limiting CVD	pts with IC.	
sugar in the 6 mo group. Total cholesterol	12 mo training group. The absolute claudication	Rutherford stage 0 or 4–	lasting for 6 or 12 mo in	
<ul> <li>No changes in weight, total cholesterol, or blood</li> </ul>	evaluated, but greater benefit was found in the	Exclusion criteria:	and strength training	
program	Results: Significant increases in all parameters	I	of combined aerobic	
<ul> <li>Greater improvements resulted from the 12 mo</li> </ul>		ABI ≤0.9,	intervention consisting	<u>24825596</u>
MWD, and muscle strength parameters.	walking speed, muscle strength	Rutherford stage 1–3,	Nonrandomized	2014(201)
<ul> <li>Combined exercise increased walking speed,</li> </ul>	1° endpoint: Maximal walking distance,	Inclusion criteria: PAD	Study type:	Pilz M, et al.
	& 95% CI)			Year Published
Comment(s)	(include P value; OR or RR;		Study Size	Author;
Summary/Conclusion	Primary Endpoint and Results	Patient Population	Study Type/Design;	Study Acronym;
Therapy–Section 6.	Evidence Table 33. Nonrandomized Trials, Observational Studies, and/or Registries for Exercise Therap	, Observational Studie	Nonrandomized Trials	Evidence Table 33.
				Q
rment questionnaire.	quanty of line, NC 1, failuonitized controlled that, NN, relative risk, NF E, ravings of bereation: SDE, shortness of breath: StO2, tissue oxygen saturation: ST, stent revascularization: TM, treadmill: and WIQ, walking impairment questionnaire.	erative risk, Kr E, ratings or pe	of breath: StO ₂ , tissue oxvoi	training: SOB. shortness
E supervised exercise: SET supervised exercise	peripriefy aftery disease, i Acc, personal attinuates questioninates of paragraphic avertions. CPD eventies blood processor CPT encoded evention avertion of the second eventies are allocated eventies and the second eventies are allocated eventies and the second eventies are allocated eventies and the second eventies are allocated eventies are allocated eventies are allocated eventies are allocated eventies and the second eventies are allocated eventies are allocated eventies are allocated eventies are all attinued and avertice are allocated eventies are allocated event	estivinitaile, FLT, paintiee un	r AQ, personar attributes qu	peripticity artery disease,
C, optimal medical care; OR, odds ratio; PAD,	LE, lower extremity; METs, metabolic equivalent; MI, myocardial infarction; MWD, maximal walking distance; N/A, not applicable; OMC, optimal medical care; OR, odds ratio; PAD	myocardial infarction; MWD,	s, metabolic equivalent; MI,	LE, lower extremity; MET

ABI indicates ankle-brachial index; ACC, Journal of American College of Cardiology; BMI, body mass index; CABG, coronary artery bypass graft; CI, confidence interval; CLI, critical limb ischemia; COT, claudication onset time; CV, cardiovascular; CVD, cardiovascular disease; ER, endovascular revascularization; HR: hazard ratio; HBP, high blood pressure; HR, hazard

Safety endpoint: N/A

while there was no change in the untrained control group.

distances in pts with claudication.

Study type: RCT

severe arthritis.

group

Comparator: Untrained

of rest; total exercise of 20 min during a 40 min session

Improvements were similar

between the 2 training groups.

both groups (p<0.001).

previous 12 mo; other exercise-limiting comorbidities such as angina, shortness of breath,

Size: n=76 pts

ratio; hs-CRP, high-sensitivity C-reactive Protein; IC, intermittent claudication; JAMA, Journal of American Medical Association; LAEI, large artery elasticity; LDL, low density lipoproteins;

•		-		
of better organized diabetes care"	amputations	amputations from 7/1995–6/2000		al.
Reduction in major amplitations "a result	1º and mint. Incidence of major and minor	Inclusion criteria: All I F	Study type: NR	Canavan R.Let
	remained significant on multivariate analysis			
	limb salvage times. Multidisciplinary care		Size: 85 pts	
CLI	Results: 67 vs. 42% at 1 y; also higher mean	8/2010-6/2012		<u>25073577</u>
amputation-free survival in pts with R5/6		pts" with R5/6 CLI at a single hospital	retrospective cohort	2015(206)
Multidisciplinary care improves	1° endpoint: 1 y amputation-free survival	Inclusion criteria: "All consecutive	Study type: NR,	Chung J, et al.
	interventions		operations	
major amputations	amputations; 44% increase in vascular		Size: n=790 diabetic foot	<u>22431496</u>
from reactive to proactive" and reduces	Results: 37.5% reduction in transtibial			2012(205)
sustained impact in changing surgery type		operations 2006–2008 vs. 2008-2010	retrospective cohort	et al.
Interdisciplinary care as a "rapid and	1° endpoint: Amputation level, case mix	Inclusion criteria: All diabetic foot	Study type: NR,	Armstrong DG,
			prevalence of DM (~4,800)	
	100K; major amputations=16 vs. 3.6 per 100K		population with 2.4%	
diabetic pts"	Results: All amputations=19.1 vs. 9.4 per		Size: n=200,000 pt	
low incidence of major amputations in		between 1982 and 1993		<u>8542736</u>
important role to reduce and maintain a	inhabitant) of major and minor amputation	primary amputations from toe to hip"	retrospective cohort	1995(204)
"Multidisciplinary approach plays an	1° endpoint: Annual incidence (per	Inclusion criteria: "All DM related	Study type: NR,	Larsson J, et al.
			pathway)	
	Results: 23% nonpathway vs. 7% pathway		(55 nonpathway, 60	
infections"			Size: n=115 pts	10028467
treatment of potentially limb-threatening	infection			1999(203)
recognition, evaluation and expedient	amputation among those admitted with	infections 1993 and 1995–1996	retrospective cohort	Werber B
Established pathway allows "earlier	1° endpoint: Prevalence of major (leg)	Inclusion criteria: All diabetic foot	Study type: NR,	Crane M and
				Year Published
	& 95% CI)		,	Author:
Comment(s)	(include P value; OR or RR;		Study Size	Acronym;
Summary/Conclusion	Primary Endpoint and Results	Patient Population	Study Type/Design;	Study

CVD indicates cardiovascular disease; IC, intermittent claudication; MWD, maximum walking distance; PAD, peripheral artery disease; and pt, patient.

exercise groups.

walking time compared to supervised walking

 Used community walking programs to treat PAD pts with IC

more effective than community walking studies at with general recommendations to walk at home. Community trials that incorporated more advice and feedback for PAD pts in general resulted in similar outcomes with no differences in peak

more feedback and monitoring

Exclusion criteria: N/A

<u>18071005</u>	with 1.94% prevalence of DM		Results: Decrease in incidence from 564– 176/100K pts with DM between first and fifth y after change; increase in angioplasty	
	) 		prevalence	····
Williams DT, et al.	Study type: NR, retrospective & prospective	Inclusion criteria: All DM or PAD pts	<u>1° endpoint</u> : Incidence of major and minor amoutation	"Formation of a well-defined Imultidisciplinaryl service has been
2012(208)	cohorts	12/2005 (before service) vs. 1/2006-	<b>)</b> -	associated with further demonstrable
22303433	Size: n=220,000 pts with	12/2009 (after service)	<b><u>Kesults</u>:</b> Fewer major amputations among DM pts (peak of 24.7 to nadir of 1.07 per	foot disease."
	4.2% prevalence of DM (9,328)		10,000); decrease in minor amputations	
Driver VR, et al. 2005(209)	<u>Study type</u> : NR	hetween 1999–2003	1° endpoint: Incidence of LEA (all levels)	Multidisciplinary care improves outcomes, decreases amputation rates
<u>15677774</u>	Size: n=About 350,000		Results: Decreased amputation incidence	
	population (4,940 with DM)		from 9.9–1.6 per 1K (71% of which were minor)	
Wrobel JS, et al.	Study type: Cross-sectional	Inclusion criteria: Surveys of	1° endpoint: Correlation between lower	Improved programming coordination more
2003 (210)	Size: n=10 Veterans Affairs	general, vascular, and orthopedic	extremity amputation rates and	influential than feedback coordination or site rankings on decreasing amoutation
	medical centers	podiatrists; physical therapists;	Results: Significant negative correlation	rates
		pedorthists; orthotists; DM care	between programming coordination and total	
		specialists; DM educators; dermatologists: wound care	and minor amputations	
		specialists: and infectious disease		
		clinicians; and 10 randomly-selected		
		primary care providers		
Vartanian et al.	Study type: NR,	Inclusion criteria: Pts with	<u>1° endpoint</u> : Time to wound healing,	Multidisciplinary care can help effectively
2015 (211) 25596408	Lettosbective Leview	signle institutional amputation	reulceration rate, and ambulatory status.	status in pts with limb threatening
	Size: n=91 limbs from 89 pts	prevention clinic from March 2012-	Results: 67% of wounds were present >6 wks	neuroischemic wounds. Hindfoot or ankle
		July 2013. Pts at highest risk for limb	before referral. A total of 151 podiatric and 86 vascular interventions were prformed with an	wounds can adversely influence the outcome. Healing can be prolonged and a
		(ischemic ulcer or gangrene) or	equal distribution of endovascular and open	substancial proportion of pts can be
		diabetic foot ulcers.	revascularizations. Complete wound healing observed in 59% of wounds. and average time	expected to have a recurrence, therefore surveillance is mandatory. A coordinated
		Exclusion criteria: New pts evaluated for benign conditions (e.g.,	to full healing was 12 wk. Hindfood wounds predictive of failure to heal (OR: 0.21: p <0.01:	amputation prevention program may help to minimize hospital readmissions in the
		arthritis, overuse injuries, simple	95% CI: 0.06–0.68).	high-risk population.
		ulcers minor trauma, radiculopathy)		

2				
However, MRI is costly and may not be	Results:	imaging diagnostic modalities for	Size: n=9 articles from the	
evaluated, MRI was the most accurate.		assess the accuracy of clinical or		2008(215)
Among the imaging tests that we	1° endpoint:	Inclusion criteria: studies that	Study type: Meta-analysis	Dinh MT, et al.
amputation in pts with diabetic foot ulcers.	redness, periwound or pretibial edema, the presence of pus, lymphadenitis/lymphangitis, fever (all p<0.01) and elevated CRP (p=0.01).			
toes are risk factos for lower extremity	Incidence of amputation increased with	<1 y		
but not amputations excluding the lesser	including the hallux	12 mo and those with life expectancy		
independently predicted any amplifation	indenwent amplitations provimal to and	the insilateral foot during the previous		
presence of increased (non)parateric	maior) within 1 v follow-rin: 103 pts (128%)	narticinating centers for an ulcer of	810 010-11:	
periwound or pretibial edema. The	<b>Do: 16.</b> 150 (200/) ato (106 minor and 22	Evaluation anitoria: Dto troated in the	Cize: n=575 nte	
elevated CRP levels, and the presence of	amputation, or death	Eurodiale study.	study	2015(214)
Positive probe-to-bone test, deep ulcer,	1° endpoint: Healing of the foot ulcer, major	Inclusion criteria: Part of the	Study type: Prospective	Pickwell K, et al.
			Size: N/A	
	Results: N/A	Exclusion criteria: N/A	foot infections	<u>22619242</u>
			new guidelines for diabetic	2012(213)
	1º endroint: N/A	Inclusion criteria: N/A	Study type: Summary of	Linekv RA et al
	corracted=0.645; SE=0.0483; 95% CI: 0.559– 0.732			
	Composite predictor c=0.783; coverfitting			
	significant.			
	IDSA combination of signs were not			
	insignificant (c=0.56; n=0.055)			
		<ul> <li>Receiving anticoagulation therapy</li> </ul>		
	Results:	Coagulopathies		
		Patelet count <125,000/mm ³		
	microbial load.	cells/mm ³		
	specific to secondary wounds as compared to	<ul> <li>White blood cell count &lt;1500</li> </ul>		
	predictor computed from the classic and signs	Exclusion criteria:		
	<ul> <li>discriminatory accuracy of a composite</li> </ul>			
	probability of the IDSA confidination of signs	Affiliated beenited		
provides a moderate level of discrimination	Sensitivity, specificity, and concordance	Department of Veterans Attairs		
A composite predictor based on all signs	microbial load (reference standard),	nonarterial diabetic foot ulcers from a	Size: n=64 pts	<u>19147524</u>
signs	probability of each sign as compared to	<ul> <li>Pts with ≥1 full-thickness,</li> </ul>		2009(212)
well nor does the IDSA combination of	<ul> <li>Sensitivity, specificity, and concordance</li> </ul>	<ul> <li>Age ≥18 y of age</li> </ul>	study	<u>a</u>
Individual signs of infection do not perform	1° endpoint:	Inclusion criteria:	Study type: Cross sectional	Gardner SE, et

				. 1
	predictor of nonhealing in PAD pts only.			
	peripheral neuropathy, and PAD. Infection is a			
	without help, ESRD, larger ulcer size,			
lk two separate disease states	age, male sex, HF, inability to stand or walk		Size: n=1,088 pts	
concomitant PAD should be defined as	healed. Predictors of nonhealing are older	Exclusion criteria: N/A		<u>18297261</u>
not diabetic foot ulcers with or without	Results: At 1-y follow up, 23% of pts had not		EURODIALE Study	2008(216)
with and without PAD, suggesting that		EURODIALE Study	cohort study within the	al.
Predictors of healing differ between pts	1° endpoint: Wound healing	Inclusion criteria: Part of the	Study type: Prospective	Prompers L, et
	from 0.28–0.75			
Bu	of osteomyelitis was highly variable, ranging			
osis	Sensitivity of plain radiography for diagnosis			
	49.45	Exclusion criteria: N/A		
	or a positive probe-to-bone test result was			
oone	• The pooled diagnostic OR for exposed bone	as well as the reference test		
	0.14,	to participate in the test being studied		
as	likelihood ratio for a negative test result was	diagnosis of osteomyelitis. All pts had		
radiographs provided limited information	for a positive test result was 9.40, and the	specimens as the reference test for		
atio detection of osteomyelitis. Plain	0.91 (95% CI: 0.89–0.92). The likelihood ratio	and/or microbiologic culture of bone	relevant articles	
/ of had low-to-moderate accuracy for	diagnosis of osteomyelitis and a specificity of	that used histopathologic examination	reference lists of potentially	
scan and indium-labeled leukocyte scans	sensitivity of 0.87 (95% CI: 0.71–0.96) for	diabetes and foot ulcer, and studies	studies identified by perusing	
ad a readily available. Nuclear medicine bone	<ul> <li>A positive probe-to-bone test result in had a</li> </ul>	diagnosis of osteomyelitis in pts with	literature search and 59	

Infectious Disease Society of America; LEA, lower extremity amputation; LPS, Limb Prevention Service; MDC, multidisciplinary care; NR, nonrandomized; OR, odds ratio; pt, patient; and RR, relative risk.

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sociation,
1, Inc. and American
rerican Heart Association, Inc. and American College of Cardiology Foundatic
Foundation

Data Supp	plement 34a	Data Supplement 34a. Functions of a Multidisciplinary Foot Care / Amputation Prevention Team–Section 7.	ıltidisciplinary	/ Foot Care /	Amputation F	Prevention Tea	am–Section 7.		
Study	Patient	<b>Risk Stratification</b> ,	Prophylactic	Protocols,	Wound Care,	Infection	Close	Orthotics	Other
Name	Education	Testing for	Podiatric	Algorithms,	Including	Management	Post-Operative	and	
		PAD	ouigeiy	Pathways	in Clinic		Монноннд	riostietics	
Crane									
1999				<					
<u>10028467</u> (203)				>					
Driver									Research; community
2005	×	×			×	×	×	×	outreach/education
(209)									
Williams									Admission to vascular
2012 22503433	×			×	×				inpatient service for infection; multidisciplinary clinics
(208)									
Pogers									Gait analysis; medical
20804929		×	×		×	×	×	×	
(217)									
Sumpio									
2010		×	×		×	×	×	×	
20488327		>	>		>	>	>	>	
Eitzaorold									
- incherain									
19436764		×			×	×	×		
(219)									
Wrobel									Ease in recruiting staff;
2006	×			×					attendance at diabetic foot
<u>16649651</u>									care education program in
(220)									past 3 yrs
PAD indicate	PAD indicates peripheral artery disease.	tery disease.							

Data Supplement 34a. Functions of a Multidisciplinary Foot Care / Amputation Prevention Team-Section 7.

Evidence Table	35 RCTe Compar	inn Endovæcular Treatme	nt and Endovascula	ar Vereus Noninvasive Ti	Fyidence Table 35 RCTs Comparing Endovascular Treatment and Endovascular Versus Noninvasive Treatment of Claudication-Section 8 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 value; OR or RR; & 95% Cl)	Relevant 2º Endpoint (if any); Study Limitations; Adverse Events
Tetteroo E, et al. 1998(221) 9643685	<u>Aim</u> : Determine superiority of iliac PTAS vs. PTA	<ul> <li>Inclusion criteria:</li> <li>Claudication</li> <li>Iliac artery stenosis &lt;5cm</li> </ul>	Intervention: PTAS Comparator: PTA	<u>1° endpoint:</u> Reduction in symptoms; QoL	<ul> <li>No difference between groups at 2 y</li> <li>Group I=PTAS. Group II=PTA. The mean follow- up was 9.3 mo (range 3–24). Initial hemodynamic success and complication rates were 110 (81%) of</li> </ul>
	<u>Study type</u> : RCT <u>Size</u> : n=279 pts	<ul> <li>Exclusion criteria:</li> <li>Stenosis &gt;10 cm in length</li> <li>Arterial occlusion &gt;5 cm in length, or ≤5 cm not allowing the passage of a guide wire</li> <li>Stenosis involving the distal</li> </ul>			149 limbs and 6 (4%) of 143 limbs (group I) vs. 103 (82%) of 126 limbs and 10 (7%) of 136 limbs (group II), respectively. Clinical success rates at 2 y were 29 (78%) of 37 pts and 26 (77%) of 34 pts in groups I and II, respectively (p=0.6); however, 43% and 35% of the pts, respectively, still had
		(e.g., severe cardiac or cerebrovascular abnormality, malignant disease)			groups during follow-up. 2 y cumulative patency rates were similar at 71% vs. 70% (p=0.2), respectively, as were reintervention rates at 7% vs. 4%, respectively (95% CI: 2%–9%).
Klein WM, et al. 2004(222) <u>15286319</u>	<u>Aim</u> : Determine superiority of iliac PTAS vs. PTA	<ul> <li>Inclusion criteria:</li> <li>Claudication</li> <li>Iliac artery stenosis &lt;5cm</li> </ul>	Intervention: PTAS Comparator: PTA	<u>1° endpoint</u> : Technical success and incidence of reintervention	<ul> <li>No difference between groups</li> <li>Long-term follow-up from above study. The mean follow-up period was 5.6 y±1.3 (±standard deviation). There were no significant differences</li> </ul>
	<u>Study type:</u> <u>Size</u> : n=279 pts	Exclusion criteria: • Stenosis of >10 cm in length • Occlusion of >5 cm in length or of ≤5 cm if it did not			between primary stent placement and primary angioplasty treatment groups in regard to number of reinterventions in the treated iliac arteries (33 [18%] of 187 segments and 33 [20%] of 169
		<ul> <li>enigm, or or so crimination allow the passage of a guidewire; stenosis involving the distal aorta</li> <li>Or severe comorbidity (e.g., severe cardiac or cerebrovascular abnormality, malignant disease)</li> </ul>			segments, respectively) or in the ipsilateral legs (45 [25%] of 181 legs and 50 [30%] of 164 legs, respectively). Sex, presence of critical ischemia, and length of stenosis were predictors of whether a pt would require iliac reintervention.

could be followed up until 2 y after intervention for			<ul> <li>Acute CLI, previous bypass surgery, or stenting of the</li> </ul>	Size: n=104 pts	00070011
superficial femoral artery obstructions, 98 (94%)			Exclusion criteria:	Study type: RCT	2007(225)
<ul> <li>PTAS is superior to PTA for long lesions (lesion length 112 mm PTAS and 93 mm PTA)</li> <li>Of 104 pts with chronic limb ischemia and</li> </ul>	by duplex at 2 y	Comparator: PTA	3–5 and SFA stenosis	PTA	Schillinger M, et
			placement.		
			recanalization before or		
			<ul> <li>Pts with iliac dissection, an</li> <li>associated AAA or iliac</li> </ul>		
			stenoses.		
			as PTA or stenting for iliac	-	
			<ul> <li>Pts undergoing</li> <li>endovascular treatment such</li> </ul>	<u>Size</u> : PTAS n=83 pts vs. ABF n=86 pts	
			Exclusion criteria:		
				Retrospective	
/80% vs 80%) were similar			בט //, ווששמפ וששש, וב //, רובי, 7%)	Study type:	10004040
	secondary patency; survival	Comparator: ABF	(claudication 53% rest pain,	aorto-bifem	2008(224)
ABF than for R/PTAS	success; primary patency;		iliac occlusive disease	disease. PTAS vs.	al.
Primary patency at 3 y was significantly higher for	1° endpoint: Technical	Intervention: PTAS	Inclusion criteria: Sx aorto-	Aim: Iliac occlusive	Kashyap VS, et
long-term failure was reduced by 39% after stent placement compared with PTA.					
placement to treat critical ischemia. The risk of					
stenoses vs. 53% for occlusions after stent					
were 7.7% for sterioses vs. o 1% for occlusions after stent placement to treat claudication and 67% for					
after PTA to treat critical ischemia. These rates					
and were 53% for stenoses vs. 44% for occlusions					
54% for occlusions after PTA to treat claudication					
primary patency rates were 65% for stenoses vs.					
adjusted for lesion type and disease severity. 4 y				Size: n=301 pts	
Analyzed data included technical failures and were				uning one	
significant corrected. Complication and mortality frates were not statistically significantly different			without specified endpoints	analysis	
group (96%), but the difference was not statistically	Safety endpoint: Mortality			Study type: Moto	<u>177CN76</u>
group was 91%; the rate was higher in the stent		Comparator: PTA	<ul> <li>Iliac artery involvement</li> </ul>	PTAS vs. PTA	1997(223)
The immediate technical success rate in the PTA	success; primary patency		<ul> <li>Claudication of CLI</li> </ul>	superiority of iliac	Hunink MG
<ul> <li>No difference between groups</li> </ul>	1° endpoint: Technical	Intervention: PTAS	Inclusion criteria:	Aim: Determine	Bosch JL and

Krankenberg H, et al. 2007 (226) <u>17592075</u>	FAST
PTAS Study type: RCT <u>Size</u> : n= 244 pts	Aim: SFA PTA vs.
<ul> <li>Exclusion criteria:</li> <li>TL that required pretreatment with adjunctive devices, e.g., lasers or debulking catheters</li> <li>A TL that extended into the popliteal artery; previous stent implantation in the targeted SFA</li> <li>Multiple lesions &gt;10 cm in length</li> <li>Acute or subacute (≤4 wk) thrombotic occlusion</li> <li>An untreated ipsilateral iliac artery stenosis</li> <li>Ongoing dialysis treatment</li> <li>Treatment with oral anticoagulants other than antiplatelet agents.</li> </ul>	<ul> <li>SFA</li> <li>Untreated inflow disease of the ipsilateral pelvic arteries (&gt;50% stenosis or occlusions)</li> <li>Inclusion criteria: SFA</li> </ul>
<u>Comparator</u> : PTA	Intervention: PTAS
restenosis	1° endpoint: Technical
difference between PTAS and PTA • Overall, stent fractures were detected in 45 of 121 treated legs (37.2%). In a stent-based analysis, 64 of 261 stents (24.5%) showed fractures, which were classified as minor (single strut fracture) in 31 cases (48.4%), moderate (fracture of >1 strut) in 17 cases (26.6%), and severe (complete separation of stent segments) in 16 cases (25.0%). Fracture rates were 13.2% for stented length ≤8 cm, 42.4% for stented length >8– 16 cm, and 52.0% for stented length >16 cm. In 21 cases (32.8%) there was a restenosis of >50% diameter reduction at the site of stent fracture. In 22 cases (34.4%) with stent fracture there was a total stent reocclusion. According to Kaplan Meier estimates, the primary patency rate at 12 mo was significantly lower for pts with stent fractures (41.1% vs. 84.3%, p<0.0001).	ultrasound and for clinical and hemodynamic outcome by treadmill walking distance and ABI. Restenosis rates at 2 y were 45.7% (21 of 46) vs. 69.2% (36 of 52) in favor of primary stenting compared with balloon angioplasty with optional secondary stenting by an ITT analysis (p=0.031). Consistently, stenting (whether primary or secondary; n=63) was superior to plain balloon angioplasty (n=35) with respect to the occurrence of restenosis (49.2% vs. 74.3%; p=0.028) by a treatment-received analysis. Clinically, pts in the primary stent group showed a trend toward better treadmill walking capacity (average, 302 vs. 196 m; p=0.12) and better ABI values (average, 0.88 vs. 0.78; p=0.09) at 2 y, respectively. Reintervention rates tended to be lower after primary stenting (17 of 46 [37.0%] vs. 28 of 52 [53.8%]; p=0.14) • For short lesions mean length 45 mm no

droups were 2 1.9% vs. 55.0% (p=0.005) at 6 mo by					
			<ul> <li>I Intreated inflow disease of</li> </ul>		
dilation. Restenosis rates in the stent and PTA			stenting of the SFA		
(26%) due to a suboptimal result after balloon			<ul> <li>Previous bypass surgery or</li> </ul>	Size: n=73 pts	
secondary stenting was performed in 10 of 39 pts			Acute CLI	<b>?</b>	
groups (p=0.011), respectively. In the PTA group,			Exclusion criteria:	Study type: RCI	
98±54 mm and 71±43 mm in the stent and PTA		Comparator: PTA	1 - -	2	19859954
<ul> <li>Average length of the treated segments was</li> </ul>	patency		stenosis and claudication	PTA	2009(228)
<ul> <li>PTAS is superior to PTA</li> </ul>	1° endpoint: Primary	Intervention: PTAS	Inclusion criteria: SFA	Aim: SFA SES vs.	Dick P, et al.
			were ineligible		
			immunosuppressive therapy		
			receiving dialysis or		
			vascular bypass or who were		
			inadequate to support		
			poor inflow that was		
			<ul> <li>Angiographic evidence of</li> </ul>		
revascularization.			resolved		
sterit naciales resulted in loss of patericy of TE			treated that could not be		
induction collected in the induction of potencies of T			<ul> <li>Thrombus in the area to be</li> </ul>		
fractures occurred in 3.1% of stants implanted. No			vessel segment to be treated		
/81.3% vs. 36.7%; n<0.0001) Through 12 mo			Aneurysmal disease in the		
natency at 12 mo was better for the stent group					
(p<0.0001). Duplex ultrasound-derived primarv			precided sale illsertion of an		
compared with 45.1% for the angioplasty group			procluded cafe insertion of an		
revascularization was 87.3% for the stent group			Extensive PVD that		
final analysis. At 12 mo, freedom from TL			the target limb		
revascularization and loss of primary patency in the			<ul> <li>Previous bypass surgery of</li> </ul>		
dissection. Bailout stenting was treated as a TL			hepatic insufficiency		
suboptimal angiographic result or flow-limiting			creatinine >2.0 mg/dL) or		
group underwent bailout stenting because of a			<ul> <li>Renal failure (serum</li> </ul>		
83.9%; p<0.01). 29 (40.3%) pts in the angioplasty			medications or materials		
compared with the angioplasty group (95.8% vs.			<ul> <li>Known allergies to study</li> </ul>		
residual stenosis) was superior for the stent group			antihistamines, or both		
angioplasty group. Acute lesion success (<30%			pretreatment with steroids,		
mm for the stent group and 64 mm for the			that was not amenable to		
angioplasty. The mean total lesion length was 71			<ul> <li>Sensitivity to contrast media</li> </ul>		
nitinol stents or percutaneous transluminal			categories 4–6)	<u>Size</u> : n= 206 pts	
artery and IC were randomized to implantation of			<ul> <li>Pts with CLI (Rutherford</li> </ul>		
superficial femoral artery and proximal popliteal			Exclusion criteria:	Study type: RCT	
States and Europe with obstructive lesions of the		Comparator: PTA			20484101
<ul> <li>A total of 206 pts from 24 centers in the United</li> </ul>	derived patency		artery stenosis	PTA	2010(227)
<ul> <li>Mean lesion length 71 mm; PTAS superior</li> </ul>	<u>1° endpoint</u> : 1 y duplex	Intervention: PTAS	Inclusion criteria: Fem/pop	Aim: SFA SES vs.	Laird JR, et al.

<b>DEBATE-SFA</b> Liistro F, et al. 2013(230) <u>24239203</u>		<b>IN.PACT</b> Tepe G, et al. 2015(229) <u>25472980</u>	
<u>Aim</u> : PEB+BMS vs. PTA+BMS <u>Study type</u> : RCT <u>Size</u> : n=104 pts	<u>Size</u> : n= 331 pts	<u>Aim</u> : SFA DCB vs. PTA <u>Study type</u> : RCT	
Inclusion criteria: Claudication and SFA stenosis Exclusion criteria: • Life expectancy <1 y • Contraindication for combined antiplatelet therapy • Known allergy to nickel or paclitaxel • Need for major amputation	<ul> <li>Exclusion criteria:</li> <li>Lesion and/or occlusions located in or extending to the popliteal artery or below the ankle joint space</li> <li>Inflow lesion or occlusion in the ipsilateral iliac, SFA, or popliteal arteries with length ≥15 cm</li> <li>Significant (≥50% DS) inflow lesion or occlusion in the ipsilateral iliac, SFA, or popliteal arteries left untreated</li> <li>Previously implanted stent in the TL(s). Aneurysm in the TL</li> <li>Acute thrombus in the TL</li> </ul>	Inclusion criteria: IC or ischemic rest pain attributable to superficial femoral and popliteal PAD	<ul> <li>the ipsilateral pelvic arteries (&gt;50% stenosis or occlusion)</li> <li>Known intolerance of study medications or contrast agent.</li> </ul>
Intervention: PEB+BMS <u>Comparator</u> : PTA+BMS		Intervention: DCB Comparator: PTA	
<u>1º endpoint</u> : 12 mo binary restenosis		1° endpoint: 12 mo primary patency	
<ul> <li>PEB+BMS is superior to PTA+BMS</li> <li>Mean lesion length was 94±60 vs. 96±69 mm in the PEB+BMS and PTA+BMS groups (p=0.8), respectively. The primary endpoint occurred in 9 (17%) vs. 26 (47.3%) of lesions in the PEB+BMS and PTA+BMS groups (p=0.008), respectively. A near-significant (p=0.07) 1 y freedom from TL revascularization advantage was observed in the PEB+BMS group. No major amputation occurred. No significant difference was observed according to lesion characteristics or technical approach.</li> </ul>	attributable to superficial temoral and popliteal PAD were randomly assigned in a 2:1 ratio to treatment with DCB or PTA. The primary efficacy endpoint was primary patency, defined as freedom from restenosis or clinically driven TL revascularization at 12 mo. Baseline characteristics were similar between the 2 groups. Mean lesion length and the percentage of total occlusions for the DCB and PTA arms were 8.94±4.89 and 8.81±5.12 cm (p=0.82) and 25.8% and 19.5% (p=0.22), respectively. DCB resulted in higher primary patency vs. PTA (82.2% vs. 52.4%; p<0.001). The rate of clinically driven TL revascularization was 2.4% in the DCB arm in comparison with 20.6% in the PTA arm (p<0.001). There was a low rate of vessel thrombosis in both arms (1.4% after DCB and 3.7% after PTA [p=0.10]). There were no device- or procedure-related deaths and no major amputations	<ul> <li>DCB superior to PTA</li> <li>The IN.PACT SFA Trial is a prospective, multicenter, single-blinded, randomized trial in which 331 pts with IC or ischemic rest pain</li> </ul>	CTA, and 2.9% vs. 18.9% (p=0.033), 18.2% vs. 50.0% (p=0.006), and 34.4% vs. 61.1% (p=0.028) at 3, 6, and 12 mo by sonography, respectively. Clinically, pts in the stent group reported a significantly higher maximum walking capacity compared with the PTA group at 6 and 12 mo.

		Werk M, et al. 2012(232) 23102018	Scheinert D, et al. 2014(231) <u>24456716</u>	
	<u>Study type</u> : RCT <u>Size</u> : n=85 pts	<u>Aim</u> : SFA DCB vs. PTA	<u>Aim</u> : SFA DCB vs. <u>Study type</u> : RCT <u>Size</u> : n=101 pts	
	<ul> <li>Exclusion criteria: Key exclusion criteria were:</li> <li>Acute thrombus or aneurysm in the target vessel</li> <li>Failure to cross the TL with a guidewire</li> <li>Inflow lesions that cannot be successfully pretreated</li> <li>Significant disease of all 3</li> </ul>	Inclusion criteria: Sx femoro-popliteal athenoscientic disease	Inclusion criteria: Rutherford class 2–5 femoropopliteal lesions Exclusion criteria: • Life expectancy ≤2 y • Creatinine >2.5 mg/dL or Hx of hemorrhagic stroke ≤3 mo • Previous or planned intervention ≤30 d • Use of adjunctive therapies (including glycoprotein IIb/IIIa inhibitors) • Severe lesion calcification • Sudden symptom onset • Acute or subacute target vessel thrombus or occlusion • Absence of ≥1 patent untreated runoff vessel • Significant inflow disease	at the time of enrollment • Failure to recanalize intended below-the-knee arteries in CLI pts at risk of major amputation
		Intervention: DCB	<u>Intervention</u> : DCB <u>Comparator</u> : PTA	
	blinded angiographic corelab quantitative analyses	<u>1° endpoint:</u> The primary endpoint was late lumen	<u>1° endpoint</u> : The primary endpoint was angiographic late lumen loss at 6 mo. Secondary outcomes included adjudicated major adverse events (death, amputation, TL thrombosis, reintervention), functional outcomes, and pharmacokinetics.	
102	Inserve undergoing per cutaireous it ansignment angioplasty were randomized to paclitaxel-coated IN.PACT Pacific or uncoated Pacific balloons. The primary endpoint was late lumen loss at 6 mo assessed by blinded angiographic corelab quantitative analyses. Secondary endpoints were binary restenosis and Rutherford class change at 6 mo, and TL revascularization + major adverse clinical events (major adverse events=death, target limb amputation, or TL revascularization) at 6 and 12 mo. 85 pts (91 cases=interventional procedures) were randomized in 3 hospitals (44 to DEB and 47	<ul> <li>DEB is superior to PTA</li> <li>Pts with sx femoro-popliteal atherosclerotic</li> </ul>	<ul> <li>DCB superior to PTA</li> <li>Demographic, PVD, and lesion characteristics were matched, with mean lesion length of 8.1 3.8 cm and 42% total occlusions. At 6 mo, late lumen loss was 58% lower for the Lutonix DCB group (0.46 1.13 mm) than for the control group (1.09 1.07 mm; p=0.016). Composite 24 mo major adverse events were 39% for the DCB group, including 15 TL revascularizations, 1 amputation, and 4 deaths vs. 46% for uncoated balloon group, with 20 TL revascularizations, 1 thrombosis, and 5 deaths. Pharmacokinetics showed biexponential decay with peak concentration (Cmax) of 59 ng/mL and total observed exposure (AUC(all)) of 73 ng h/ml. For successful DCB deployment excluding 8 malfunctions, 6 mo late lumen loss was 0.39 mm and the 24 mo TL revascularization rate was 24%.</li> </ul>	

. 104	-	-	-	-	
<ul> <li>DES is superior to PTA±BMS</li> </ul>	1° endpoint: 2 mo rates of	Intervention: DES	Inclusion criteria: Sx	Aim: SFA DES vs.	Zilver PTX
between the 2 treatment groups. Primary patency at 3, 6, 9, and 12 mo of follow-up was 84%, 82%, 75.6%, and 73.5% for the stent graft group and 90%, 81.8%, 79.7%, and 74.2% for the femoral- popliteal surgical group. 13 pts in the stent graft group had 14 reinterventions, and 12 reinterventions occurred in the surgical group. This resulted in secondary patency rates of 83.9% for the stent graft group and 83.7% for the surgical group at the 12 mo follow-up.		Synthetic fem-pop bypass	Exclusion criteria: • No aorto-iliac disease • <1 infrapop artery patent to ankle	<u>Study type</u> : RCT <u>Size</u> : n=86 pts	
<ul> <li>No difference</li> <li>Pts were monitored for a median of 18 mo. No statistical difference was found in the primary patency (p=0.895) or secondary patency (p=0.861)</li> </ul>	1° endpoint: 12 mo duplex primary patency	Intervention: Viabahn Comparator:	Inclusion criteria: Sx femoral-popliteal arterial occlusive disease	<u>Aim</u> : SFA: Viabhan vs. synthetic fem- pop bypass	Kedora J, et al. 2007(236) <u>17126520</u>
ultrasonography (65% vs. 40%, p=0.0003). A patency benefit was seen for lesions at least 3 cm long. At 12 mo, chronic limb ischemia status was 15% further improved for the stent-graft group (p=0.003). There were no significant differences between treatment groups with regard to the occurrence of early or late major adverse events.			popliteal artery of <1 infrapop artery patent to the ankle	<u>Size</u> : n=197 pts	
<ul> <li>Viabahn superior to PTA alone</li> <li>The stent-graft group had a significantly higher technical success rate (95% vs. 66%, p&lt;0.0001)</li> <li>and 1 v primary vessel patency rate at dupley</li> </ul>	1° endpoint: 12 mo duplex primary patency	Intervention: Viabahn Comparator: PTA	Inclusion criteria: Sx SFA PAD Exclusion criteria: Occluded	<u>Aim</u> : SFA: Viabahn vs. PTA Study type: RCT	Saxon RR, et al. 2008(235) 18503895
VIABAHN stent graft (88.8% vs. 69.8%; p=0.04), although secondary patency rates did not differ between bare nitinol stent and stent graft recipients (89.3% vs. 79.5%; p=0.304). There were no instances of procedure-related mortality or amputation. The hemodynamic improvement and quality measures improved equally in both groups.					
differ between pts treated with the VIABAHN stent graft and those who received a bare nitinol stent (24.2% vs. 25.9%; p=0.392). Stent fractures were significantly more common in bare nitinol stents (50.0%) than in the VIABAHN endoprostheses (2.6%). Primary-assisted patency rates were higher in those receiving bare nitinol stents than the			Exclusion criteria: Occluded popliteal artery of <1 infrapop artery patent to the ankle		

-					
restenosis. For both types of stents, improvements			Exclusion criteria: Lesions	Size: n=93 pts	
diseased SEA and in sustaining freedom from			c		
Both the sirolimus-eluting and the bare SMART		Comparator: BIVIS	C CIUSIONS OF STENOSES TASC	Study type: DOT	2006(239)
vs. BMS	trom restenosis		limb ischemia and SFA	BMS	Duda SH, et al.
No meaningful difference between sirolimus DES	1° endpoint: Freedom	Intervention: DES	Inclusion criteria: Chronic	Aim: SFA: DES vs.	SIROCCO
			Frevious target vessel     stenting		
			<ul> <li>Lesion pretreatment with</li> </ul>		
			inflow tract	Size: n=474 pts	
		-	<ul> <li>Utreated &gt;50% DS of the</li> </ul>	•	
clinical durability in comparison with standard endovascular treatments		provisional BMS	exclusion criteria included:	Study type: RCT	
Zilver PTX DES provided sustained safety and	patency	Comparator: DTA w		SIMB	PMC4823823
information previously unavailable.	event-tree survival and	(no polymer)	tem/pop PAD	PIA w provisional	2015(238)
• 5-y results from Zilver PTX study show long-term	1° endpoint: 2 mo rates of	Intervention: DES	Inclusion criteria: Sx	Aim: SFA DES vs.	Dake MD, et al.
0.9% (4/457).					
the stent fracture rate (both DES and BMS) was					
compared with the provisional BMS group, and (3)					
clinical benefit (90.5% and 72.3%; p=0.009)					
patency (89.9% vs. 73.0%; p=0.01) and superior					
provisional DES group exhibited superior primary					
group (88.3% vs. 75.8%; p<0.001), (2) the					
superior clinical benefit compared with the PTA					
evaluations, (1) the primary DES group exhibited					
satisfying the primary hypotheses. In the secondary					
primary patency (83.1% vs. 32.8%; p<0.001),					
event free survival (90.4% vs. 82.6%; p=0.004) and					
the primary DES group exhibited superior 12 mo			c		
and PTA groups. Compared with the PTA group,			stenting		
event free survival and patency in the primary DES			<ul> <li>Previous target vessel</li> </ul>		
(n=59). Primary endpoints were the 12 mo rates of			adjunctive devices		
assignment to provisional DES (n=61) or BMS			<ul> <li>Lesion pretreatment with</li> </ul>		
failure and underwent secondary random			inflow tract	Size: n=474 pts	
approximately 65±40 mm). 120 pts had acute PTA			<ul> <li>Utreated &gt;50% DS of the</li> </ul>		
similar between groups (eg, average lesion length,		provisional BMS	exclusion criteria included:	Study type: RCT	
Demographics and lesion characteristics were		Comparator: PTA w	Exclusion criteria: Major		21953370
implantation (n=236) or PTA (n=238).	patency			BMS	2011(237)
Dte were randomly assigned to primary DEC	event-free survival and	(nn nnlvmer)	fem/non PAD	PTA w nrovisional	Dake MD et al

Tepe G, et al. 2008(240) <u>18272892</u>	
<u>Aim</u> : SFA: PTA vs. PTA with balloon dipped in paclitaxel <u>Study type</u> : RCT <u>Size</u> : n=154 pts	
Inclusion criteria: Pts with Rutherford stages 1–5 sx & stenosis or occlusion of a femoropopliteal artery - Poor inflow; absence of a patent crural artery - Acute onset of symptoms - Pregnancy - Life expectancy of >1 y - Contraindications to required medication	>20 cm
Intervention: Paclitaxel dipped balloon <u>Comparator</u> : PTA	
<u>1º endpoint</u> : Angiographic restenosis at 6 mo and TVR	
• DCB superior • The mean ( $\pm$ SD) age of the pts was 68±8 y, 24% were smokers, and 49% had DM. 27% of the lesions were total occlusions, and 36% were restenotic lesions. The mean lesion length was 7.4±6.5 cm. There were no significant differences in baseline characteristics between the groups. There were no adverse events attributable to the paclitaxel-coated balloons. At 6 mo, the mean late lumen loss was $1.7\pm1.8$ mm in the control group, as compared with $0.4\pm1.2$ mm (p<0.001) in the group treated with paclitaxel-coated balloons and $2.2\pm1.6$ mm (p=0.11) in the group treated with paclitaxel in the contrast medium. The rate of revascularization of TLs at 6 mo was 20 of 54 (37%) in the control group, 2 of 48 (4%) in the group treated with paclitaxel-coated balloons (p<0.001 vs. control), and 15 of 52 (29%) in the group treated with paclitaxel in the contrast medium (p=0.41 vs. control); at 24 mo, the rates increased to 28 of 54 (52%), 7 of 48 (15%), and 21 of 52 (40%)	in ABI and symptoms of claudication were maintained over 24 mo (median 24 mo ABI 0.96 for the sirolimus group vs. 0.87 for the bare stent group, p>0.05). At 24 mo, the restenosis rate in the sirolimus group was 22.9% vs. 21.1% in the bare stent group (p>0.05). The cumulative in-stent restenosis rates according to duplex ultrasound were 4.7%, 9.0%, 15.6%, and 21.9%, respectively, at 6, 9, 18, and 24 mo; the rates did not differ significantly between the treatment groups. The TLR rate for the sirolimus group was 6% and for the bare stent group 13%; the TVR rates were somewhat higher: 13% and 22%, respectively. Mortality rates did not differ significantly between the groups.

mm, mean stent diameter was 6.1±0.4 mm, and					
148±98 mm. mean stented length was 190±116			ţ		
balloons (n=45 lesions). Mean lesion length was			and >60 mm in length.		
using cryoplasty (n=45 lesions) or conventional			of stents >5 mm in diameter	Size: n=/4 pts	
lesions, were randomly assigned to post-dilation			lesions requiring implantation	2	00010677
<ul> <li>74 pts, with 90 stented superficial temoral artery</li> </ul>		Comparator: PTA	<ul> <li>Superficial femoral artery</li> </ul>	Study type: RCI	2012(242)
angioplasty			• Sx PAD		al.
binary restenosis compared to conventional balloon	restenosis	Cryoplasty PTA	• DM	PTAS with Cryo PTA	Banerjee S, et
<ul> <li>Post-dilation with cryoplasty balloon reduced</li> </ul>	1° endpoint: 12 mo binary	Intervention:	Inclusion criteria:	Aim: SFA: PTAS vs.	COBRA
			proximal to the target stent.		
			affecting target stent or		
			<ul> <li>Grade 4/5 stent fracture</li> </ul>		
			completion of 30 d follow-up		
			interventions prior to		
			cardiac surgery or		
			<ul> <li>Planned or predicted</li> </ul>		
			the TL		
			<ul> <li>DES or covered stents in</li> </ul>		
			Aneurysm within TL		
			dependent		
			mg/dL unless dialysis-		
			<ul> <li>Serum creatinine ≥2.5</li> </ul>		
			to study procedure		
			target vessel within 3 mo prior		
TI R /HR· 0.48· 95% CI· 0.31_0.74)			<ul> <li>Previous treatment to the</li> </ul>		
FI A+PTA was associated with a 52% reduction in					
were 5.8% vs. 20.5% (p<0.001), respectively.			- Systemic infaction in TI		
(p<0.005), and 30 d major adverse event rates			hypercoanulability		
mo freedom from TLR was 73.5% vs. 51.8%			Uncontrolled		
procedural complications. ELA+PTA and PTA pt 6-			procedure		
vs. 82.7%; p=0.01) with significantly fewer			allergies that could affect the		
demonstrated superior procedural success (93.5%			<ul> <li>Contraindications or</li> </ul>		
exhibited total occlusion. ELA+PTA pts			or MI 60 d prior to procedure		
cm vs. 19.3±11.9 cm, and 30.5% vs. 36.8% of pts			<ul> <li>Cerebrovascular accidents</li> </ul>		
were enrolled. Mean lesion length was 19.6±12.0			<ul> <li>Life expectancy &lt;12 mo</li> </ul>		
PTA pts (61.7% male; mean age 67.8±10.3 y)			• ALI	Size: n=250 pts	
pts (62.7% male; mean age 68.5±9.8 y) and 81			<ul> <li>Pregnancy</li> </ul>		
specified interim analysis. A total of 169 ELA+PTA	MACE	Comparator: PTA	Exclusion criteria:	Study type: RCT	25499305
early efficacy demonstrated at a prospectively-	Safety endpoint: 30 d				2015(241)
<ul> <li>Study enrollment was stopped at 250 pts due to</li> </ul>		ELA+PTA	Class 1–4 SFA ISR	ELA+PTA vs. PTA	Dippel EJ, et al.
<ul> <li>ELA+PTA superior to PTA alone for SFA ISR</li> </ul>	1° endpoint: 6 mo TLR	Intervention:	Inclusion criteria: Rutherford	Aim: SFA ISR:	EXCITE ISR

oin <u>t</u> : Max treadmill nset of claudication low-up <b>ndpoint:</b> Non-	tion: dical therapy herapy ASA+advise	Inclusion criteria: • Unilateral IC • Short stenoses Exclusion criteria: • Previous angioplasty or arterial surgery to the sx leg	<u>Size</u> : n=62 pts (30 PTA+Meds, 32 Med	
It: Non-	tion: dical therapy herapy	Inclusion criteria:     Unilateral IC     Short stenoses     Exclusion criteria:	Size: n=62 nts (30	
lax treadmill claudication	tion: dical therapy	<ul> <li>Inclusion criteria:</li> <li>Unilateral IC</li> <li>Short stenoses</li> </ul>		
lax treadmill claudication	dical therapy	Inclusion criteria:     Unilateral IC	<u>Study type</u> : RCT	1997(244) <u>9357454</u>
			Aim: 2 y tollow-up ot above study	Whyman MR, et al.
mo in PTA group (p≤0.05)		treadmill examination <ul> <li>Any psychiatric illness or other reason making follow-up difficult</li> </ul>	<b>Size:</b> n=62 pts (30 PTA+Meds, 32 Med Tx ) 47 femoral; 15 iliac	
mo in PTA group (p≤0.05)		<ul><li>Inability to manage the</li></ul>	Study type: RCT	
mo in PTA group (p≤0.05)		<ul> <li>Duration of symptoms &lt;1</li> </ul>	disease.	
mo in PTA group (p≤0.05)	exercise	Pts taking oral	Duplex measured extent of occlusive	
-	therapy=ASA+advise on smoking and	<ul> <li>MI within 6 mo</li> </ul>	reported MWD, ABI, QoL (NHP) and	
<ul> <li>ABI higher in PTA group at 6 mo (p≤0.05)</li> <li>Lower Nottingham Health Score pain scores at 6</li> </ul>	Medical therapy (Medical	<ul> <li>Exclusion criteria:</li> <li>Previous angioplasty or</li> </ul>	claudication, treadmill MWD, pt	
			until onset of	8760978
time to onset or claudication   (p≤∪.∪1) at 6 mo follow-up p<0.01	at 6 m	Short stenoses	treadmill distance	ai. 1996(243)
		Inclusion criteria:	Aim: Compare PTA	Whyman MR, et
		dimensions.		
		Imaging tield as a reference for determining vessel		
		had radio-opaque tape in the		
		infrapopliteal run-off. All pts		
		<ul> <li>Absence ≥1 vessel</li> </ul>		
in the cryoplasty group (29.3% vs. 55.8%; p=0.01; OR: 0.36: 95% CI: 0.15–0.89)		diameter stenosis) iliofemoral		
At 12 mo, binary restenosis was significantly lower		<ul> <li>Had obstructive (≥50%)</li> </ul>		
balloon angioplasty groups, respectively (p=0.02).		contrast		
5.51±0.72 mm in the cryoplasty and conventional		<ul> <li>Allergic to ASA, cioploogrei, or iodine-based radiographic</li> </ul>		
50% of the lesions were total occlusions. Post-		Exclusion criteria:		

			<ul> <li>Life incapacitating cardiac</li> <li>disease (≥NYHA class III)</li> </ul>	training in pts with IC	
	PFD, and MWD SF-36 QoL	training	• AAA	supervised exercise	
	in terms of ABI, maximum	supervised exercise	Exclusion criteria:	revascularization or	
exercise group—still no dimerence at 12 mo	Functional capacity defined	Hospital based	• ABI <0.9	mo after	
2010 correction of statistical methods—better for	Cafaty and point:	Comparator	• Max PFWD <350 m	functional capacity,	<u>19188327</u>
<ul> <li>By 12 mo no difference</li> </ul>	in one Rutherford category	with provisional stent	•	clinical success,	2009(246)
<ul> <li>At 1 wk endo superior</li> </ul>	1° endpoint: Improvement	Intervention: PTA	Inclusion criteria:	Aim: To compare	Spronk S, et al.
		a regular basis)			
		exercises at home on			
		also encouraged to			
		improved. Pts were			
		exercise tolerance			
		increasing as the pt's			
		intensity of exercise			
		performed, with the			
		exercises were			
		min. Dynamic leg			
		Each class lasted 30			
		the pt's progress.			
		basis according to			
		required on a regular		Size: n=56 pts	
		attendance was	specified in article		
		first 6 mo. After this,	Exclusion criteria: Not	Study type: RCT	
		classes 2x/wk for the		Binner actiona	
	v follow-in	(Supervised exercise	surgeons and radiologists	exercise training	
2 DTA cally (as starts ar mod Ty)	to claudication or MMD at 6	Exercise training	andonlasty as agreed by	treated with PTA vs	21855020
• No difference in endpoints at 6 y follow-up (only	difference in ADI distence	Comparator	ciadulcation lesion(s) on	dictoron in DAD ato	2011/2/E/
• Small study	<u>1° endpoint:</u> Better ABI in	Intervention: PTA	Inclusion criteria: Unilateral	Aim: Compare ABI	Perkins, JM, et
			difficult		
			Ally psychiatric inness of     other reason making follow-rin		
			<ul> <li>Inability to manage the</li> </ul>		
			mo		
			<ul> <li>Duration of symptoms &lt;1</li> </ul>		
			anticoaculants		

				<u>18771879</u>	Spronk S, et al. 2008/247)	
	Size: n=76 endo; n=75 hospital based supervised exercise	Study type: RCT	study	analysis of above	<u>Aim</u> : Cost- effectiveness	<u>Size</u> : n=76 endo; n=75 hospital based supervised exercise
same-side stenoses at both the iliac and femoral levels, requiring multiple revascularization procedures) • Isolated tibial artery disease • Lesions deemed unsuitable for revascularization (iliac or femoropopliteal TASC type D	<ul> <li>AAA</li> <li>Life incapacitating cardiac disease (≥NYHA class III)</li> <li>Multilevel disease (i.e.,</li> </ul>	Exclusion criteria:	• ABI <0.9	• Max PFWD <350 m	Inclusion criteria: ● IC	<ul> <li>same-side stenoses at both the iliac and femoral levels, requiring multiple revascularization procedures)</li> <li>Isolated tibial artery disease</li> <li>Lesions deemed unsuitable for revascularization (iliac or femoropopliteal TASC type D and some TASC type B and/or C lesions, such as a unilateral external iliac occlusion that involved the origins of the internal iliac and/or common femoral artery or single or multiple femoral artery or single or multiple femoral artery distal bypass procedure)</li> <li>Prior treatment for the lesion (including exercise table)</li> </ul>
	e U	supervised exercise	<u>Comparator</u> : Hospital based		Intervention: PTA with provisional stent	
	Safety endpoint: Not reported	costs, and incremental	capacity over a 12 mo period, cumulative 12 mo	related QoL and functional	1° endpoint: Mean improvement of health-	
				difference between QoL at 12 mo	<ul> <li>Endo costs more than exercise program when adjusted for OALV however this study had no</li> </ul>	

Taft C, et al. 2001(249) <u>11472043</u>	Gelin J, et al. 2001(248) <u>11472042</u>	
<u>Aim</u> : QoL analysis of above study <u>Study type</u> : : RCT single center <u>Size</u> : Invasive (n=87 pts; 17 were endo) vs. Meds (n=89) vs. Control (n=89)	<u>Aim</u> : Invasive vs. supervised exercise vs. control <u>Study type</u> : RCT single center <u>Size</u> : Invasive (n=87 pts; 17 were endo) vs. meds (n=89) vs. control (n=89)	
Inclusion criteria: IC with ABI <0.6 Exclusion criteria: Pts with a medical Hx contraindicating surgery and/or with other disorders severely limiting walking evaluation on a treadmill	Inclusion criteria: IC with ABI <0.6 Exclusion criteria: Pts with a medical Hx contraindicating surgery and/or with other disorders severely limiting walking evaluation on a treadmill	and some TASC type B and/or C lesions, such as a unilateral external iliac occlusion that involved the origins of the internal iliac and/or common femoral artery or single or multiple femoral popliteal lesions in the absence of continuous tibial vessels to improve inflow for a distal bypass procedure) • Prior treatment for the lesion (including exercise training)
Intervention: Surgery or endo <u>Comparator</u> : Supervised exercise (3 30 min sessions for 6 mo and then 2 sessions per wk) <u>Control</u> : Advise on risk factor management and walking	Intervention: Surgery or endo <u>Comparator</u> : Supervised exercise (3 30 min sessions for 6 mo and then 2 sessions per wk) <u>Control</u> : Advise on risk factor management and walking	
<u>1° endpoint</u> : Invasive therapy improved disease specific symptoms (waling pain) but no difference in other aspect of QoL other aspect of QoL	<u>1° endpoint:</u> ABI (p<0.01) and max treadmill time (p<0.01) improved only in invasive group <u>Safety endpoint</u> : No difference in 1 y mortality	
	• Only 59% of exercise pts competed training	

<u>17055756</u>	2007(251) single center		Nylaende M, et	OBACT <u>Aim</u> : Endo vs. OMT		stent (n=46)	vs. SE (n=42) vs.		al. 2015/186) Study type: RCT	Murphy TP, et meds	F/U exercise vs. stent vs	CLEVER 18 mo Aim: Supervised		stent (N=46)	<u>Size</u> : Meds (n=22) vs	2	22090168 Study type: RCT	al. meds	P, et	CLEVER <u>Aim</u> : Supervised	therapy (n=7)	Best medical		revascularization+be	Size: Endovascular		2006(250) Study type: RCT	
<ul> <li>Both Aortoiliac and</li> </ul>	0	RCT	<ul> <li>PAD with disabling IC</li> </ul>		comorbid conditions that limited walking ability		2) vs.		RCT Gardner protocol) and	but <11 min on a graded	•	vised Inclusion criteria: Severe IC	Exclusion criteria: CLI or comorbid conditions that limited walking ability		0	Objective evidence of a	RCT Gardner protocol)	to walk ≥2 but <11 min on a	•	vised Inclusion criteria:	7)	<u> </u>	тегару	ation+be	ascular	Exclusion criteria: N/A		
and	Medical therapy		ing IC	<u>a:</u> Intervention: PTA	ins that		stenosis	ficant	g tne <u>Comparator:</u> and <u>Stenting vs Medical</u>	ä	to walk ≥2 Supervised exercise		<u>a</u> : CLI or ins that ility		significant	nce of a therapy alone	Stenting vs. medical		ΪŤ	i: Intervention:				therapy	Optimal medical	a: N/A Comparator:		
PTA group compared to			<ul> <li>PFWD, MWD at 3, 12</li> </ul>	PTA <u>1° endpoint</u> :			SE vs. ST p=0.16	meds vs. ST p=0.04	edical 5 0+5 4 min ST 3 7+4 7	compared to baseline	peak walking time at	1° endpoint: Change in			SE vs. SI p=0.022	vs. SE p<0.001	(meas 1.2±2.0 mms, 5.8±4.6, ST 3.7±4.9)	compared to baseline	peak walking time a	1° endpoint: Change in						(p=0.008)	(p=0.013) and MWD	
				<ul> <li>On QoL questionnaires pain was less in PTA</li> </ul>						1	18 mo	in N/A					) meds	<b>1</b>	0	in    Both SE and ST experienced improvement in								_

			<ul> <li>serious cardiopulinionary comorbidity (NYHA III–IV)</li> </ul>	(n=35) Consisted of	
			in a SET program	Size. Endovaccular	
	endo alone (p=0.011)		<ul> <li>Hx of or current participation</li> </ul>	single center	
	<ul> <li>Endo+SE superior to</li> </ul>	Comparator: Endo	Exclusion criteria:	Study type: RCT	2011(203) 21571547
N/A	<ul> <li>1° endpoint:</li> <li>6 mo absolute walking</li> </ul>	Intervention: Endo+SE	Inclusion criteria: PAD pts with Rutherford 1–4	<u>Aim</u> : Endo vs. Endo+SE	Kruidenier LM, et al.
			which prohibits exercise.		
			musculoskeletal or cardiac)		
			Concomitant disease (e.g.,		
			Doppler pressure >50 mm hg)		
			ulcers or gangrene with a	mo	
			< 50 mm hg or presence of	times per wk tor b	
	longer with PTA(p=0.05)		CLI (absolute Doppler BP	walking training 3	
	(p=0.05) and ICD was		mandatory	(n=88) I readmill	
	was 78% greater with PTA		severe that intervention was		
	<ul> <li>Aorto-iliac disease AWD</li> </ul>		consider angioplasty or so	or procedures vs.	
	longer with PTA (p=0.004)		<ul> <li>Symptoms too mild to</li> </ul>	(n=8/) multiple types	
	(p=0.04) and ICD was		Exclusion criteria:	revascularization	
	was 38% greater with PTA			Size: Endovascular	
	<ul> <li>Fem-pop disease AWD</li> </ul>	once a wk for 6 mo	disease		<u>19022184</u>
	distance	Comparator: SE	disease, 34 pts with aortoiliac	single center	2008(252)
	time and initial claudication		<ul> <li>93 pts with femoropopliteal</li> </ul>	Study type: RCT	RM, et al.
	<ul> <li>24 mo average walking</li> </ul>	PTA±stent	<ul> <li>PAD pts with IC (ABI &lt;0.9)</li> </ul>		Greenhalgh
N/A	1° endpoint:	Intervention:	Inclusion criteria:	Aim: Endo vs. SE	MIMIC
			<ul> <li>Renal Insufficiency</li> </ul>		
			<ul> <li>Use of warfarin</li> </ul>		
			abrogating organized exercise		
			<ul> <li>Other physical disability</li> </ul>		
			DM ulcer		
			endovascular surgery		
			<ul> <li>Previous vascular or</li> </ul>		
			• CLI	therapy (n=28)	
			<ul> <li>Subjective PFWD &gt;400 m</li> </ul>	Optimal medical	
	ABI p=0.0013		Exclusion criteria:	therapy (n=28)	
	p=0.0001, MWD p=0.0009,			timal medical	
	<ul> <li>94 mon values PEWD</li> </ul>		nonulation was included	revascularization+on	

			2012(254) 22021102	Mazari FA, et al														
	(n=60), SE (n=60) Endovascular revascularization+su pervised exercise (n=58)	single center Size: Endovascular revascularization	Study type: RCT	Aim: Endo vs. SE	exercise program 2x/wk for 6 mo	Nonspecified	pervised exercise (n=35)	revascularization+su	Vs. Endovascular	and femoral	primary stent	recanalization with	stenoses or	primary stent	angioplasty with	stenoses,	placement for iliac	iliac angioplasty
	<ul> <li>Inability to tolerate treadmill testing</li> <li>Ischemic changes on ECG during treadmill testing</li> <li>Ipsilateral surgery/PTA in previous 6 mo</li> </ul>	Exclusion criteria: • Critical ischemia • Incanacitating systemic	suitable for angioplasty and femoropopliteal lesions	Inclusion criteria: PAD with sx unilateral claudication										<ul> <li>Major amputation or tissue loss</li> </ul>	<ul> <li>No insurance for SET</li> </ul>	Dutch language	<ul> <li>Insufficient knowledge of the</li> </ul>	<ul> <li>Other serious comorbidity preventing physical activity</li> </ul>
Concomitant therapy: All pts were prescribed antiplatelet therapy	Percutaneous transluminal angioplasty Supervised exercise therapy: Circuit of exercises 3x/ wk for 12 wk	alone vs. SE alone Endovascular therapy:	Comparator: Endo	Intervention: Endo+SE														
			procedural complications	<u>1° endpoint:</u> ICD, MWD, reneat revascular peri-														
				<ul> <li>No significant difference at 12 mo in ICD and MWD or Col</li> </ul>														

	Mazari FA, et al. 2010(195) <u>19762206</u>	
<u>Size</u> : n=178 pts	<u>Aim</u> : 3 mo data for above trial <u>Study type</u> : RCT	
<ul> <li>Critical ischemia</li> <li>Incapacitating systemic disease</li> <li>Inability to tolerate treadmill testing</li> <li>Ischemic changes on ECG during treadmill testing</li> <li>Ipsilateral surgery/PTA in previous 6 mo</li> </ul>	Inclusion criteria: PAD with sx unilateral claudication suitable for angioplasty Exclusion criteria:	
<ul> <li>alone</li> <li>Endovascular therapy: Percutaneous transluminal angioplasty Supervised exercise therapy: Circuit of exercises 3 times per wk for 12 wk</li> <li>Concomitant therapy: All pts were prescribed antiplatelet therapy (ASA and/or clopidogrel), received smoking cessation advice and support (including nicotine replacement therapy and NHS smoking cessation program),</li> </ul>	Intervention: Endo+SE <u>Comparator:</u> • Endo alone vs. SE	(ASA and/or clopidogrel), received smoking cessation advice and support (including nicotine replacement therapy and NHS smoking cessation program), and risk factor
reported	<u>1° endpoint</u> : ICD, MWD, repeat revascular, peri- procedural complications Safety endpoint: None	
	At 3 mo PTA + SEP provided greater improvement in claudication than SEP or PTA alone. See above for 12 mo results	
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		Nordanstig J, et al. 2011(255) <u>21397530</u>
Size: Inv (n=100) vs. OMT(n=101)	Study type: RCT multicenter	<u>Aim</u> : Invasive+OMT vs. optimal medical tx
<ul> <li>Pts with ≥2 previously occluded vascular reconstructions.</li> </ul>	<ul> <li>Incorrect Dx</li> <li>Other disorders limiting</li> </ul>	Inclusion criteria: IC >6 mo Exclusion criteria: ● Age ≥85 y
	ization:	Intervention: Invasive+OMT Comparator:
INV vs. the NON group. 2 SF-36 physical subscales, Bodily Pain (p<0.01) and Role Physical (p<0.05) improved significantly more in the INV vs. the NON group. There were 7% crossovers against the study protocol in the INV group. group.	MWP was not significantly (p=0.104) improved in the	<u>1° endpoint</u> : 2 y Mean Walking Performance and QoL
		N/A

	2015(257) 25721067	Maloor RD of al				<u>25095886</u>	al. 2014(256)	IRONIC Nordanstig J, et									
Study type: Meta- analysis of RCTs	surgical vs. SE vs. Meds	Aim. Epoto ve		(n=79) vs. OMT (n=79)	Size: Invasive	Study type: RCT (single center)		<u>Aim</u> : Invasive+OMT vs. optimal medical									
exclusively enrolling pts with CLI, defined as rest pain or tissue loss	IC pts Exclusion criteria: Trials	<ul> <li>⇒2 previously failed</li> <li>⇒2 previously failed</li> <li>ipsilateral vascular</li> <li>interventions</li> </ul>	<ul> <li>occasionari est pari,</li> <li>infrarenal aortic thrombosis)</li> <li>Weight &gt;120 kg (maximum nossible load on treadmill)</li> </ul>	IC, subcritical ischemia with	considered mandatory (main criteria according to protocol:	<ul> <li>Symptoms so severe that invasive treatment was</li> </ul>	Very mild symptoms	Inclusion criteria: IC >6 mo									
	vs. surgical vs. SE vs. Meds	Intervention: Con-	• 2 pts hybrid	<ul><li>16 pts open</li></ul>	<ul><li>• 52 pts</li></ul>	invasive Rx 70 received	79 allocated to	Intervention: Endo except for TASC D	home several times per d.	perform an additional	distance as often as possible and to	maximal claudication	least 1 H/d and to	Instructed to walk at	and a written training	Verbal training advice	were managed according to national
superior to medical management in terms of walking distance and	Open surgery, endovascular therapy, and exercise therapy were	10 00 00 00 00 00 00 00 00 00 00 00 00 0					VascularuQoL (p<0.01) at 12 mo better with Inv	<u>1° endpoint:</u> SF 36 (p<0.001) and									
<ul> <li>Evidence is sparse supporting superiority of one of three approaches</li> <li>Isolated iliac or femorpopliteal disease pts. may</li> </ul>	<ul> <li>Efficacy of surgery, endovascular and exercise therapy seemed to be superior to medical mgmt for walking distance, pain and claudication</li> </ul>	Minimal data on port offortivopor	population independent of whether surgical or endovascular approach was required	<ul> <li>Both aortoiliac and femoropoliteal disease pts were enrolled. Pragmatic design to include large IC</li> </ul>	<ul> <li>Outcomes not stratified by surgical vs.</li> <li>endovsacular procedures.</li> </ul>	<ul> <li>Invasive therapy group included 18 pts treated with surgical and hybrid approach to invasive Rx</li> </ul>	<ul> <li>group (p=0.003)</li> <li>No difference Inv vs. Meds for MWD change</li> </ul>	• Distance to onset of claudication better with Inv. Invasive (+124 m) vs. the noninvasive (+50 m)									

																																enrolling 1,548 pts	reviews and 12 trials	Cita: n=0 contamatio
Safety endpoint: Not reported	distance but not MWD	overall QoL score and IC	parameters, better SF 36,	increased leg BP and flow	<ul> <li>Invasive revasc generally</li> </ul>	mgmt or exercise:	<ul> <li>Revasc with medical</li> </ul>	better in surgical arm.	patency was generally	info about 30-d mortality but	<ul> <li>Some SRs had conflicting</li> </ul>	and a high 30-d mortality.	stay, high complications	significantly longer hospital	open bypass had	<ul> <li>Studies generally showed</li> </ul>	surgery:	<ul> <li>Endovascular vs. open</li> </ul>	be a better approach	Combination of both may	mgmt. or exercise:	approaches with medical	Endovascular	alone but not exercise	<ul> <li>ABI improved vs. surgery</li> </ul>	alone or exercise alone	vs. Medical management	walking distance improved	<ul> <li>Max. and symptom free</li> </ul>	physical training):	RCTs for Surgery (with	Results:	ciaudicalion	alaridization
																																	limited data.	da hattar than amhinad diegoed geografing to the

pressure; CI, confidence interval; CLI, critical limb ischemia; CTA, computed tomography angiography; DCB, drug coated balloon; DEB, drug eluting balloon; DES, drug eluting stent; DS, 119

et al	currainal via avaraina		ve eurninal ve	improvided MNN/D p=0.01	
2015(258)	vs. Meds	Exclusion criteria: N/A	exercise vs. Meds	SF-36 improved in all	inconclusive to determine superiority for walking distance or QoL for claudication
25963038				groups compared to meds	
	Study type: Meta- analysis of RCTs		Comparator: Medication alone	(usual care)	
				Safety endpoint: Not	
	Size: n=35 studies			reported	
	of 7,475 pts				
McPhail IR, et al.	<u>Aim</u> : Compare the standard LE	<ul> <li>Inclusion criteria:</li> <li>Known or suspected IC</li> </ul>	Intervention: Active pedal plantarflexion	<u>1° endpoint:</u> Active pedal plantarflexion compared	N/A
2001(259)	vascular laboratory	<ul> <li>Referred for LE treadmill</li> </ul>		favorably with treadmill	
11300450	treatmill exercise	exercise testing	Comparator: LE	exercise for the noninvasive	
	with the office-based		treadmill exercise	objective assessment of	
	active pedal	Exclusion criteria:	testing	PAOD	
	plantarflexion	<ul> <li>Ankle SBP &gt;300 mmHg or</li> </ul>			
		brachial systolic BP		reported	
	Study type:	<ul> <li>CLI and inability to walk on</li> </ul>			
	Prospective,	a treatmill or perform active			
	crossover study	peual piantamexion			
	<b>Size:</b> n=50 pts (100				
	LE) · ·				
Schulte KL, et al.	<u>Aim</u> : Compare primary placement	<ul> <li>Pts undergoing treatment</li> </ul>	Intervention: Primary placement of	<u>1° endpoint</u> : Sustainable clinical improvement after	<ul> <li>Sustained improvement at 1 y in 74.3% of the pts treated with primary stenting and in 68.6% of the</li> </ul>
2015(260)	of a self-expanding	for infrapopliteal stenosis in	a self-expanding	12 mo, defined as ≥1	pts treated with PTA and bailout stenting (p>0.05).
26245919	nitinol stent to PTA	11 European centers	nitinol stent vs. PTA	category increase for	• Freedom from TLR (76.6% and 77.6%), mortality
	with bailout stenting		with bailout stenting	Rutherford category 3 pts, a	(7.4% vs 2.1%), and amputation [8.9% (major
	in infrapopliteal	Exclusion criteria:		≥2 category increase for	6.7%) vs 13.2% (major 8.7%)] at 1 y were not
	arteries of pts with	• N/A		CLI pts compared with	significantly different.
	claudication or CLI			paseilne.	<ul> <li>Primary self-expanding nitinol stenting did not show statistically different clinical outcomes</li> </ul>
	) - - -			Safety endpoint: TLR,	compared to PTA with bailout stenting
	<u>Study type</u> : RC1			mortality, and amputation assessed after 12 mo.	
	2				

stent revascularization; TASC, transatlantic inter-society consensus; TL, target lesion; TLR, total lesion revascularization; TPP, treatment per-protocol; TVR, target vessel distance; MWP, mean walking performance; N/A, not applicable; NEJM, New England Journal of Medicine; NHP, Nottingham Health Score; NYHA, New York Heart Association; OR, odds group; ISR, in stent restenosis; ITT, intention to treat; JACC, Journal of American College of Cardiology; LE, lower extremity; MACE, major adverse cardiac event; MWD, maximal walking revascularization; and VIA, viabahn treatment. percutaneous transluminal angioplasty, and stenting; RR, relative risk; SE, supervised exercise; SEP, supervised exercise; SES, self-expanding stents; SFA, superficial femoral artery, ST percutaneous angioplasty stent; PVD, peripheral vascular disease; QALY, quality adjusted life year; QoL, quality of life; RCT, randomized controlled trail; R/PTAS, recanalization, ratio; OMT, osteopathic manipulative treatment; PAD, periphery artery disease; PEB, paclitaxel eluting balloon; PFWD, pain free walking distance; PTA, percutaneous angioplasty; PTAS, diameter stenosis; ECG, electrocardiogram; ELA, excimer laser antherectomy; HR, hazard ratio; IC, intermittent claudication; ICD, International Classification of Disease; Inv, intervention

Treatment of Cla	Treatment of Claudication–Section 8.1.			
Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Summary/Conclusion Comment(s)
Scheinert D, et al. 2005 (261) <u>15653033</u>	Study type: Prospective series assessing SES fracture incidence	Inclusion criteria: PTAS for claudication or chronic ischemia Exclusion criteria: None	<ul> <li><u>1° endpoint:</u></li> <li>Stent fracture incidence</li> <li>Restenosis incidence</li> </ul>	<ul> <li>Stent fractures predict restenosis</li> <li>Overall, stent fractures were detected in 45 of 121 treated legs (37.2%). In a stent-based analysis, 64 of 261 stents (24.5%) showed fractures, which were</li> </ul>
	<u>Size</u> : n=93 pts	reported	<b><u>Results:</u></b> The primary patency rate at 12 mo was significantly lower for pts with stent fractures (41.1% vs. 84.3%, p<0.0001).	classified as minor (single strut fracture) in 31 cases (48.4%), moderate (fracture of >1 strut) in 17 cases (26.6%), and severe (complete separation of stent segments) in 16 cases (25.0%). Fracture rates were 13.2% for stented length $\leq 8$ cm, 42.4% for stented length >8–16 cm, and 52.0% for stented length >16 cm. In 21 cases (32.8%) there was a restenosis of >50% diameter reduction at the site of stent fracture. In 22 cases (34.4%) with stent fracture there was a total stent reocclusion. According to Kaplan Meier estimates, the primary patency rate at 12 mo was significantly lower for pts with stent fractures (41.1% vs. 84.3%; p<0.0001).
Sakamoto Y, et al. 2013(262)	Study type: Case series evaluating PTAS patency for SFA CTO	Inclusion criteria: SFA CTO undergoing PTAS	<u>1° endpoint</u> : 5 y primary and secondary patency rates and the rates of freedom from bypass surgery, major	<ul> <li>Stent diameter predicts restenosis</li> <li>Mean age was 72±9 y and 31% were female pts. In total, 58% of the pts had DM and 25% were pts with</li> </ul>
<u>23536429</u>	<u>Size</u> : n=352 pts	Exclusion criteria: None reported. Lack of CTO	or minor amputation, and all-cause death <u>Results:</u> Female gender (OR: 1.95; p=0.0051) and mean stent diameter	CLI. Occluded length was 194±89 mm, mean total stent length was 198±7 mm, and mean stent diameter was 7.1±0.9 mm. 5 y primary and secondary patency rates were 51.8% and 79.5%, respectively, and the rates of freedom from bypass surgery, major or minor

Evidence Table 36. Nonrandomized Trials, Observational Studies, and/or Registries of Endovascular and Endovascular Versus Noninvasive

Shammas NW, et al. 2009(269) <u>19966364</u>	Sachs T, et al. 2011(268) <u>21880457</u>	Kalbaugh CA, et al 2006(267) <u>16814976</u>	Pell JP and Lee AJ 1997(266) <u>9507581</u>
<u>Aim</u> : Determine predictors of distal embolization in pts undergoing LE arterial peripheral endovascular	<u>Aim</u> : Determine national estimates for the costs, utilization, and outcomes of angioplasty and bypass graft for the treatment of claudication <u>Study type</u> : Retrospective analysis <u>Size</u> : n=563,143 pts	<u>Study type</u> : Case series <u>Size</u> : IC n=54 CLI n=30	revascularization (n=os) Percutaneous transluminal angioplasty or surgery (n=31) Comparator Conservative treatment (N=64) No description provided <u>Study type:</u> Observational multicenter <u>Size</u> : Endovascular revascularization (n=19) Percutaneous transluminal angioplasty or surgery (n=19) Comparator Conservative treatment (n=157) No description provided
Inclusion criteria: Pts undergoing peripheral intervention enrolled in a single center registry	Inclusion criteria: Pts who underwent endo or surgery for PAD based on ICD-9 codes Exclusion: Atherosclerosis unspecified ICD-I code	Inclusion criteria: Endo treatment of IC or ALI Exclusion criteria: None reported	<u>Inclusion criteria</u> : IC <u>Exclusion criteria</u> : N/A
<u>1° endpoint</u> : Predictors of distal embolization <u>Results</u> : Prior Hx of amputation;	<u>1° endpoint</u> : Costs and clinical outcomes <u>Results</u> : Unclear cost analysis as more PTA procedures were performed compared to surgery; lower mortality with PTA	<u>1° endpoint</u> : QoL at 1 y <u>Results</u> : Improved QoL in both IC and ALI compared to baseline	<u>1° endpoint</u> : 6 mo QOL <u>Results</u> : PTA or surgery provided improved QOL at 6 mo compared to conservative Tx
Limitation is that this is a single center registry analysis	Study limited by methodology; ICD-9 code analysis	<ul> <li>No comparative arm</li> </ul>	<ul> <li>Study did not report pts' baseline characteristics</li> <li>Study did not report pts' comorbid conditions</li> <li>Comparator(s) not well described</li> </ul>

3I indicates ankle-brachial index; A	Size: n=410 procedures in 295 pts		analysis	Study type: Retrospective		for claudication or CLI	undergoing endo revasc	in a cohort of pts	1998(270) predictors of complications	Manninen HI complications and	Matsi PJ and Aim: To report	Size: n=577 pts	analysis; case-control study	2	revasc
LI, acute limb is	procedures in			Retrospective		in or CLI	ndo revasc	pts	complications	and	7	ots	-control	-	
schemia; CAD, coronary artery disea							Exclusion: None reported		intervention at a single center	undergoing peripheral	Inclusion criteria: Pts			Exclusion: None reported	
ease; CI, confidence interval; CLI, critical limt		CAD and cerebrovascular disease	claudication; mortality was driven by	had higher mortality compared to	complications in women; pts with CLI	stenosed arteries; more bleeding	with occluded arteries compared to	Results: More complications in pts		predictors of complications	1° endpoint: Complications and			lesions predicted distal embolization	presence of thrombus, and TASC-D
ABI indicates ankle-brachial index; ALI, acute limb ischemia; CAD, coronary artery disease; CI, confidence interval; CLI, critical limb ischemia; CTO, chronic total occlusion; HF, heart										analysis	Limitation is that this is a single center retrospective				

failure; HR, hazard ratio; IC, intermittent claudication; ICD, International Classification of Diseases; JACC, Journal of American College of Cardiology; LE, lower extremity; MACE, major adverse cardiac event; OR, odds ratio; PAD, periphery artery disease; PTA, percutaneous angioplasty; PTAS, percutaneous angioplasty stent; pt, patient; QoL, quality of life; RR, relative risk; SES, self-expanding stents; SFA, superficial femoral artery; and TASC, Trans-Atlantic Inter-Society Consensus.

## Evidence Table 37. RCTs Evaluating Surgical Treatment for Claudication–Section 8.1.2.

Study	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint (if any);
Acronym;	Study Type;		(# patients) /	(Absolute Event Rates, P value;	Study Limitations;
Author;	Study Size (N)		Study Comparator	OR or RR; &	Adverse Events
Year Published			(# patients)	95% CI)	
IRONIC	Aim: Compare invasive vs.	Inclusion criteria: Stable (>6 mo)	Intervention:	1° endpoint: HRQL assessed by	<ul> <li>Exclusion criteria somewhat</li> </ul>
Nordanstig, et	noninvasive treatment	IC symptoms	<ul> <li>Invasive treatment</li> </ul>	SF-36, VascuQol. Greater	arbitrary
al.	strategies for IC		(Open surgical repair	improvement in VascuQol	<ul> <li>Only 18/158 pts had surgical</li> </ul>
2014(256)		Exclusion criteria: Mild or severe	reserved for TASC D	improved significantly more in	or hybrid procedures (Total
25095886	Study type: RCT (single	symptoms	lesions)	invasive group (p<0.01) including	procedures: 1 aortobifemoral
	center, open label)		<ul> <li>79 allocated to</li> </ul>	3/5 domain scores; claudication	bypass, 3 femoral-femoral
			invasive Rx	distance improved more in invasive	bypass, 8 ccommon femoral
	Size: n=158 pts with stable		<ul> <li>70 received</li> </ul>	group (+124m vs. +50m); change	endarterectomy/profundaplasty,
	IC (79 allocated to invasive		intervention:	in MWD not different between	5 femoral-popliteal artery
	Rx 79 to noninvasive Rx)		52 pts	groups	bypass, 1 distal to popliteal

124					
(الم_م: الحم), assisted fritting	and $0 > 10 $ ( $p - 0.700$ ), and secondary	bypass		Study type: RCT	1 000000
00 vs. 47 % IUL KOLAE	and 60% (n=0.406) and secondary	Supragenicular	<ul> <li>Previous surgery or PTA with</li> </ul>		21035693
FR the AZY for DEFAF	00% 101 bypass (p=0.107),	Comparator			0010/072\
prostructure grants (n-ou) at o y	allel o y was 47 % IUL ROFAE allu		A IC AID IO SIDISAL	TASC C and D losions of	קוטעפו גר טט, פו
For venous (n=25) and     ind     ind	<u>1° endpoint</u> : 3 y primary patency	Intervention:	Inclusion criteria: IASC C and D	AIM: Compare KSFAE or	Cichortz CC of
-	-		Popliteal arteries (P2-3 segments)		
		stenting (N=44)			
		angioplasty and	<ul> <li>Common femoral</li> </ul>	Size: n=95 pts	
		Subintimal	<ul> <li>Occlusion of iliac</li> </ul>		
	at 36 mo	Comparator:	creatinine 1.5 mg/dL)	Study type: RC1	
	1.89; 95% CI: 0.94–3.78; p=0.07)		<ul> <li>Chronic renal insufficiency (serum</li> </ul>		
	4/./% (21 OT 40) IN ENDU (HK:	(IC-NI) BUILLAIS	Intervention or bypass)	predictive or resteriosis.	
	47 70/ 101 of 40) in ENIDO (UD)	stopting (NI-51)	intervention or burner)	prodictive of restances	
	was 62 7% (32 of 46) in RF and	angioplasty and	<ul> <li>Previous treatment (endovascular)</li> </ul>	and identified factors	
claudication)	Cl: 0.99–4.2; p=0.05) at 24 mo and	distal endpoint	Exclusion criteria:	D femoropopliteal lesions	
ischemia (<50% of pts had	(25 of 44) in ENDO (HR: 2.6; 95%	endarterectomy with		interventions on (TASC)-II	23044257
group had Rutherford 4–5	76.5% (39 of 51) in RE and 56.8%	Kemote	lesions (not claudication-specific)	OT RE VS. ENDO	2012(272)
<ul> <li>61% of RE and 52% of endo</li> </ul>	1° endpoint: Primary patency was	Intervention:	Inclusion criteria: TASC-II D	Aim: Evaluated outcomes	Gabrielli R, et al.
			Pregnancy	Size: n=80 pts	
			<ul> <li>Renal insufficiency</li> </ul>	2	
			• Trauma	center, open label)	
			Redo CFE	and the second surface	
			bypass grafting	Study type: DCT (single	
<ul> <li>Limb salvage (p=0.51)</li> </ul>			<ul> <li>Simultaneous aneurysm repair or</li> </ul>	group).	
CFE vs. 84% BASI; p=0.01)			Urgent CLI		
• 1 y secondary patency (100%			Exclusion criteria:	artery enderterectomy (CEE	
				or by common femoral	
CEE vs 80% BASI: n=0 007)				implantation (BASI group)	
<ul> <li>1 v primarv patency (100%)</li> </ul>		implantation	<ul> <li>Atherosclerosis</li> </ul>	by bioabsorbable stent	
CFE vs. 92.5% BASI; p=0.038)		Comparator: BASI	<ul> <li>CFA stenosis or occlusion</li> </ul>	CFA atherosclerotic lesions	
<ul> <li>30d primary patency (100%)</li> </ul>			duration	pts undergoing treatment of	<u>25101576</u>
CFE vs. 97.5% BASI)	(7 for CFE vs. 0 for BASI, p=0.002)	randomization	<ul> <li>Claudication or CLI &gt;2 wk in</li> </ul>	hemodynamic outcome in	2014(271)
<ul> <li>Technical success (100%)</li> </ul>	1° endpoint: Surgical site infection	Intervention: 1:1	Inclusion criteria:	Aim: Compare clinical and	Linni K, et al.
		allocated)			
		treatment (N=79 pts			
		Noninvasive			
		Comparator:			
procedures		2 pts. hybrid			
surgical vs. endovsacular		surgery.			
<ul> <li>Outcomes not stratified by</li> </ul>		16 pts open			
hynass)		Endovascular			

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Heart
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Inc.
and
Inc. and American
n College of
Cardiology
American Heart Association, Inc. and American College of Cardiology Foundation

REVASAim: Compare R: supragenicular by graftingal. 2009(275)grafting Study type: RCT18990592Study type: RCT18990592had IC)	van Det RJ, et al. 19231253 19231253 femoro-popliteal b grafts. Study type: RCT (multicenter) Size: n=228 bypa: (176 [77%] for IC)	<u>Size</u> : n=1 had IC)
<u>Aim</u> : Compare RSFAE vs. supragenicular bypass grafting <u>Study type</u> : RCT <u>Size</u> : n=116 pts (77 [66%] had IC)	<u>Aim</u> : To compare ePTFE prosthesis and collagen- impregnated knitted polyester (Dacron) for AK femoro-popliteal bypass grafts. <u>Study type</u> : RCT (multicenter) <u>Size</u> : n=228 bypass grafts (176 [77%] for IC)	<u>Size</u> : n=116 pts (77 [66%] had IC)
Inclusion criteria: TASC C and D lesions of the SFA Exclusion criteria: • Previous treatment (endovascular intervention or bypass) • Chronic renal insufficiency (serum creatinine 1.5 mg/dL) • Occlusion of iliac, common femoral, and popliteal arteries (P2-3 segments)	<ul> <li>Inclusion criteria:</li> <li>Disabling claudication</li> <li>Rest pain</li> <li>Tissue loss for whom suprageniculate femoral-popliteal bypass was feasible</li> <li>Exclusion criteria:</li> <li>Previous ipsilateral femoro- popliteal procedures</li> <li>Contraindication to long-term anticoagulant therapy</li> <li>Life expectancy &gt;1 y and current treatment with chemotherapy or radiotherapy.</li> </ul>	additional stent placement of the target SFA • An SFA diameter <4 mm. SFA occlusion had to start <4 cm from the proximal SFA
Intervention: RSFAE <u>Comparator:</u> Supragenicular bypass	Intervention: AK femoro-popliteal bypass grafts were randomly allocated to either an ePTFE (n Z 114) or a Dacron (n Z 114) vascular graft <u>Comparator</u> : N/A	
<u>1° endpoint</u> : Primary patency after 1 y follow-up was 61% for RSFAE and 73% for bypass (p=0.094). Secondary patency was 79% for both groups. Subdividing between venous (n=25) and prosthetic grafts (n=30) shows a primary patency of 89% and 63% respectively at 1 y follow-up (p=0.086).	<b>1° endpoint:</b> After 5 y, the primary, primary assisted and secondary patency rates were 36% (95% CI: 26%–46%), 46% (CI: 36%–56%) and 51% (95% CI: 41%–61%) for ePTFE and 52% (95% CI: 42%–62%; p=0.01) and 70% (95% CI: 66%–76%; p=0.01) for Dacron, respectively. After 10 y these rates were respectively 28% (95% CI: 18%–38%), 31% (95% CI: 19%–43%) and 35% (95% CI: 23%–47%) for ePTFE and 28% (95% CI: 18%–38%), 49% (95% CI: 37%–61%) and 49% (95% CI: 37%–61%) for Dacron.	patency was 69 and 73% (p=0.541), respectively
NA	NA	<ul> <li>patency was 84% and 56 vs.</li> <li>63% for RSFAE (p=0.052), and secondary patency was 89% and 59 vs. 69% for RSFAE (p=0.046).</li> <li>Pts were randomized to RSFAE or bypass with the ipsilateral saphenous vein. When the saphenous vein was not available or not suitable, 23 pts received a PTFE bypass</li> </ul>

Jensen LP, et al. 2007(277) <u>17400486</u>	Ricco JB and Probst H 2008(276) <u>17997269</u>
<u>Aim</u> : Compare PTFE and polyester grafts for femoral to above-knee popliteal artery bypass <u>Study type</u> : RCT (multi- center), Scandinavia <u>Size</u> : n=427 pts (270 [65%] had IC)	<u>Aim</u> : Compare crossover vs. direct bypass for unilateral iliac occlusive disease in claudicants <u>Study type</u> : RCT (multicenter) <u>Size</u> : n=143 pts
<ul> <li>Inclusion criteria:         <ul> <li>Consecutive pts with chronic lower limb ischemia</li> <li>Considered suitable for surgical revascularization using a supragenicular prosthetic bypass graft</li> <li>Provided the pts consented to take part</li> </ul> </li> <li>Exclusion criteria:         <ul> <li>Age &lt;18 y</li> <li>Pregnant</li> <li>Previously enrolled in the study</li> <li>Considered impossible to follow</li> <li>Informed consent could not be obtained.</li> </ul> </li> </ul>	Inclusion criteria: Unilateral iliac artery occlusive disease and disabling claudication <u>Exclusion criteria</u> : N/A
Intervention: 6 mm Dacron conduit <u>Comparator</u> : 6 mm PTFE conduit	Intervention: Crossover bypass (N=74) <u>Comparator</u> : Direct bypass (N=69)
<b>1° endpoint:</b> 2 y primary patency rates for Dacron and PTFE were 70% and 57% (p=0.02), whereas the secondary patency rates were 76% and 65% (p=0.04), respectively. Primary patency at 2 y was significantly influenced by the number of patent crural vessels (2 or 3 67%, 1 50%, p=0.01). At 2 y, pts treated for CLI had a major amputation more often than pts operated on for IC, 10 and 3 respectively (p=0.003), and had higher mortality rates, 20% and 8% respectively (p=0.001).	<u>1° endpoint</u> : Primary patency and assisted primary patency Primary patency at 5 y was higher in the direct bypass group than in the crossover bypass group (92.7 vs. 73.2, p=0.001). Assisted primary patency and secondary patency at 5 y were also higher after direct bypass than crossover bypass (92.7 vs. 84.3, p=0.04 and 97.0 vs. 89.8, p=0.03, respectively). Patency at 5 y after crossover bypass was significantly higher in pts presenting no or low-grade SFA stenosis than in pts presenting high-grade (>50%) stenosis or occlusion of the SFA (74.0% vs. 62.5%, p=0.04). In both treatment groups, patency was comparable using PTFE and polyester grafts. Overall survival was 59.5±12% at 10 y.
<ul> <li>Medical therapy was not standardized</li> <li>Amputations at 2 y, (major in 4% and minor in 3%), 30 d mortality and complications (wound infections: 3% and other wound complications: 13%) occurred equally frequent in both groups.</li> </ul>	NA

	by patency rate (73%) than HUV	(n=261), or SV (n =	<ul> <li>Prior ipsilateral prosthetic fem-pop AK or below-knee bypass graft</li> </ul>	Study type: RC1	
	a significantly (p≤0.01) better	externally supported	• ABI >0.9	2 JOT	
ار م	above-knee SV bypass grafts had	Comparator:	Noncompressible vessels	knee bypass grafts.	
ຫຼຸ. ~	HUV: 70%: PTFE: 69%). After 5 v.	gran.	Exclusion criteria:	femoral-popliteal above-	7601001
÷	were similar among the different	popliteal bypass	grafting at 20 VA Medical Centers	with different bypass graft	1999(280)
	assisted primary patency rates	knee femoral-	for femoral-AK popliteal bypass	improved patency exists	and Lee KK
	1º endnoint: The cumulative	Intervention: above-	Inclusion criteria: Pts scheduled	<b>Aim:</b> To identify whether	Johnson WC
			femoral bypass grafts were eligible,		
			aortofemoral iliofemoral or femoral-		
			at the time of the temoropopliteal		
			inflow procedures were not allowed	claudication)	
			Exclusion criteria: Adjunctive	Size: n=240 pts (59% had	
		tex vs. Hemasnieid orafts	<ul> <li>Not undergone any earlier</li> <li>infrainguinal vascular procedures</li> </ul>	Study type: RCT	
		Comparator: Gore-	ment above the knee	grafts	
			with reconstitution of a popliteal seg-	femoropopliteal bypass	
	at 5 yrs between the 2 grafts.	popliteal bypass	superficial femoral artery occlusion	prosthetic above-knee	10709052
	primary or secondary patency rates	knee femoral-	<ul> <li>An angiographically demonstrated</li> </ul>	affecting patency of	2000(279)
	1° endpoint: No difference in	Intervention: Above-	Inclusion criteria:	Aim: Identify factors	Green RM, et al.
		control			
		served as their own			
		same pts; each pt			
		Contralateral leg in			
		Comparator			
		PTFE-SV)			
		(either SV-PTFE or		Size: n=43 pts (86 legs)	
		staged interventions	EXCIDSION CITEMA. NOTE mentioned		
	שפופ ווושוופו וטו טעט (ט-ט-טט).	terms of the order of	Exclusion oritoria: Nono	randomized	
	assisted primary patency rates	grait in the other,	knee reconstitution.	Study type: Drocpective	
2		and saphenous vein	<ul> <li>Long SFA occlusion with above-</li> </ul>	bypass	<u>10520903</u>
	primary and secondary patency	PTFE graft in 1 leg	<ul> <li>Failed medical therapy</li> </ul>	grafts for above-knee	1999(278)
	significant differences between	received above-knee	Bilateral disabling claudication	PTFE vs. saphenous vein	et al.

OR, odds ratio; PTA, percutaneous tra	indicates confidence interval; CFA, co	ABF indicates aortobifemoral bypass;		
OR, odds ratio; PTA, percutaneous transluminal angioplasty; PTAS, percutaneous transluminal angioplasty stent; PTFE, polytetrafluoroethylene; pt, patient; RCT, randomized controlled	indicates confidence interval; CFA, common femoral artery; CFE, common femoral artery endarectomy; CLI, critical limb ischemia; EIA-external illac artery; ENUO, endovascular interventions: ePTFE expanded notivetrafilioroethylene. HR hazard ratio: HI V human imbilical vein: IC intermittent claudication: MM/D maximum walking distance. N/A not applicable	ABF indicates aortobifemoral bypass; ABI, ankle-brachial index; AK, above knee; BASI, bioabsorbable stent; CFA, common femoral artery; CFE, common femoral endarterectomy; CI	Inadequate vein	Composite grafts
tent; PTFE, polytetrafluoroethylene; pt, patient; RCT, randomized con	ermittent claudication: MMD_maximum walking distance: N/A_not an	CFA, common femoral artery, CFE, common femoral endarterectomy		

1986(282)

PTFE vs. saphenous vein Aim: Compare patency of

control ischemia caused by popliteal or an infrapopliteal artery to

Inclusion criteria: Bypass to the

Intervention: PTFE

1° endpoint:

atherosclerosis

reconstructions for infra-inguinal arterial Veith FJ, et al.

Study type: RCT

bypass graft procedure in the same

graft conduit Venous vs. PTFE

PTFE bypasses (p=0 .036).

79.7% for vein and 57.2% for Secondary patency rates were for PTFE grafts (p=0.035).

times.

prostheses were used 76

polytetrafluoroethylene

Comparator:

leg or with the greater saphenous

Exclusion criteria: Earlier arterial

Size: n=151 bypasses (120

vein removed earlier.

for claudication)

above the knee

with the distal anastomosis femoropopliteal bypasses polytetrafluoroethylene for

knee

to the popliteal artery above the

bypass

Femoral-AK popliteal

rates after 5 yrs were 75.6% for

1° endpoint: Primary patency

venous bypass grafts and 51.9%

stretched

bypass grafts, and 6 mm

Reversed vein was used in 75

Intervention:

bypass with the distal anastomosis

nclusion criteria: Femoropoplitea

3510323

**Study type:** prospective, randomized, multicenter

claudication. <20% of pts had Size: n=845 bypasses

> approach or through deep femora Ability to treat with endovascular

Bypass for non-PAD diagnosis

Exclusion criteria:

saphenous vein graft

above-knee grafts. No difference in

rates of limb salvage for CLI.

No difference in 4 y patency for

gratt to popliteal artery.

on diameter <3.0 mm for graft to tibial artery or <4.0mm for

Inadequate vein defined based

Patency and limb salvage by

distal anastomotic site.

4 y primary patency for

Autogenus

<u>Comparator</u>

revascularizastion without bypass

p<0.001).

inferior for PTFE (49% vs. 12%) intrapopliteal bypasses were

Sequential bypasses

2003(281)

12514593

Klinkert P, et al.

Aim: To compare vein with

were not excluded, but

randomization was limited to either

HUV or PTFE

 Prior ipsilateral SV bypass graft Platelet count >106/mm² higher than 7.5 × 106/mm³) Polycythemia (red blood cell count Serum creatinine >2.0 mg/dL Size: n=752 pts

emergency surgery

226)

significantly (p≤0.01) better

patency rate than PTFE bypass

grafts (39%).

• <1 y life expectancy</p>

Oral anticoagulation,

Popliteal aneurysmal disease

Aihara H, et al.	<u>24142958</u>	Siracuse JJ, et al. 2014(285)			24080134 (784)	Lo RC, et al.	<u>25702917</u>	Nguyen BN, et al. 2015(283)	Evidence Lable So. Study Acronym; Author; Year Published
Study type: NR,	<u>Size</u> : n=1,513 pts from the ACS- NSQIP dataset (no stratification by IC/CLI/other)	<u>Study type</u> : NR		700	<u>Size</u> : n=1,797,885 nts	<u>Study type</u> : NR	<u>Size</u> : 1,843 procedures	Study type: NR	Type/Design; Study Study Study Size
Inclusion criteria:	<u>Exclusion criteria</u> : N/A	Inclusion criteria: Elective CFE		Exclusion criteria: N/A	dataset based on ICD-9 primary	Inclusion criteria: Pts admitted	NSQIP database <u>Exclusion criteria</u> : Other major procedures, hybrid procedures	Inclusion criteria: Common femoral endarterectomies in	Patient Population
1° endpoint: Primary patency	<b>Results:</b> Partial- and total-dependent functional status (OR: 9.0; 95% CI: 2.8–28.4 and OR: 21.3; 95% CI: 3.3–139.4) and dyspnea at rest (OR: 8.2; 95% CI: 1.2–58.8) predicted mortality	<u>1° endpoint</u> : 30 d mortality	<ul> <li>Women had higher mortality rates than men for all procedures (open: 1.0% vs7%; OR: 1.37; 95% CI; 1.25–1.49; p&lt;0.01; endovascular: 0.5% vs. 0.2%; OR; 1.99; 95% CI: 1.72–2.30; p&lt;0.01; open+endo: 1.8% vs8%; OR: 2.13; 95% CI: 1.76–2.58; p&lt;0.01).</li> </ul>	<ul> <li>Involution yourses an invergence young endovascular procedures and highest among those undergoing open+endo procedures.</li> </ul>	Results:	1° endpoint: In-hospital mortality stratified by gender	<b><u>Results</u>:</b> 3.4% mortality; mortality predictors included age, nonindependent functional status, preoperative dialysis, sepsis, emergency status, and ASA class 4 or 5	1° endpoint: Operative mortality	Evidence Fable So. Normalidonitzed Trials, Observational Studies, and/or Registries of Surgical Treatment to Study Acronym;       Study         Study Acronym;       Study       Patient Population       Primary Endpoint and Results         Author;       Type/Design;       (include P value; OR or RR;         Year Published       Study Size       & 95% CI)
Overall complication rate was 14.4% in the	<ul> <li>predictors of morbidity include steroid use (OR: 2.4; 95% CI: 1.4–4.1), DM (OR: 1.8; 95% CI: 1.3–2.4), and obesity (OR: 1.6; 95% CI: 1.1–2.4).</li> <li>Postoperative morbidities included cardiac (1.0%), pulmonary (1.9%), renal (0.4%), urinary tract infection (1.7%), thromboembolic (0.5%), neurologic (0.4%), sepsis (2.7%), superficial (6.3%), and deep surgical site complications (2.0%).</li> <li>At least 1 complication, including major and minor, was seen in 7.9% of the pts.</li> </ul>	<ul> <li>No claudication-specific results or ABI data</li> <li>Major morbidity (aggregate): Independent</li> </ul>	<ul> <li>In-hospital mortality lowest among pts undergoing endovascular procedures</li> </ul>	<ul> <li>Inypolicies and induces based on genues</li> <li>In-hospital mortality highest among pts who had hybrid (open+endo) procedures</li> </ul>	mortality rates stratified by procedure type - Hypothesis and models based on conder	Claudication pts were a subgroup analysis,		<ul> <li>Not claudication-specific</li> </ul>	ts Summary/Conclusion

trail; RE, remote endarterectomy; R/PTAS, percutaneous transluminal angioplasty, and stenting; RR-relative risk; RSFAE, remote superficial artery endarterectomy; SA-RIEA, stent assisted remote iliac endarterectomy; SFA, superficial femoral artery; SIA, subintimal angioplasty; SV, saphenous vein; TASC, transatlantic inter-society consensus; and TL, target lesion.

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			cohorts of CLI and	
	strategies are unknown.		evaluated mixed	
	treatment strategies and the timing of these combined		specific, while 12	
	usual care. The potential additive effects of combined	Exclusion criteria: N/A	claudication	
	intervention, and surgical intervention compared with		evidence; 35 were	
	cilostazol, exercise training, endovascular	since January 1995	contributed	
	effective. QoL showed significant improvement from	language studies published	Size: n=83 studies	
	definitively conclude which treatment is most	Reviews for relevant English		
treatments.	Results: For claudication, data were too sparse to	Database of Systematic	(AHRQ)	23844447
comparative evidence to support it over other		Embase, and the Cochrane	Systematic review	2013(289)
Surgery is effective for claudication, but limited	1° endpoint: N/A	Inclusion criteria: PubMed,	Study type:	Jones WS, et al.
patency compared with ABF."				
secondary natency but inferior primary				
hospital length of stay, and equivalent	in the PCIS aroup (76% vs. 68%; p=0.013).			
<ul> <li>"Iliac stenting has lower morbidity, shorter</li> </ul>	the ABF bypass group was significantly greater than		claudicants)	
24%; p=0.001).	in both groups (98% ABF vs. 85% PCIS). Survival in		stent group were	
superficial femoral artery disease (45% vs.	p=0.010). Secondary patency rates were equivalent		percutaneous iliac	
70%; p=0.02), and a greater incidence of	bypass was greater than for PCIS (91% vs. 73%;	Exclusion criteria: N/A	71% of	
(97% vs. 86%; p=0.002), COPD (85% vs.	Results: At 72 mo, the primary patency for ABF		(66% of ABF and	
more commonly had a Hx of nicotine abuse		aortofemoral bypass	Size: n=229 pts	
(age 60±0.9 y vs. age 65±1.2 y; p=0.002) and	Survival	undergoing illac stenting or		2317/535
<ul> <li>Pts in the ABF grafting group were younger</li> </ul>	Patency	artery occlusive disease	retrospective	2013(288)
<ul> <li>Includes pts with CLI</li> </ul>	<u>1° endpoint:</u>	Inclusion criteria: Sx iliac	Study type: NR	Sachwani GR, et al.
		Exclusion criteria: N/A	SIA/stent)	
	90.1%; and 88.2% vs. 90.1%.		(82 bypass, 58	
	was, respectively, 66.6% vs. 70.1%; 76.5% vs.	stenting	Size: n=150 limbs	
	secondary patency for bypass vs. SIA+stent groups	femoropopliteal bypass or SIA +		
	Results: 24 mo, primary, primary-assisted, and	treated with above-knee	(France)	23835109
difference between treatments		with femoropopliteal disease	retrospective	2013(287)
<ul> <li>No statistical test provided for patency</li> </ul>	1° endpoint: Patency	Inclusion criteria: Claudicants	Study type: NR	Boufi M, et al.
		TASC A/B	(111 limbs)	
		• CLI	bypass: 86 pts	
		<ul> <li>Acute ischemia</li> </ul>	pts (202 limbs);	
		<ul> <li>Hybrid procedures</li> </ul>	endovascular: 177	
		Exclusion criteria:	(313 limbs);	
	test)		Size: n=263 pts	
	the endovascular treatment group (p<0.01, log-rank	disease	(onlow)	
	69.4% in the bypass group and 67.8% and 45.2% in	TASC C/D femoropopliteal	(Japan)	<u>24232123</u>
bypass surgery group and 3.5% in the EVI	<b>Docuto:</b> 1 and 5 warmon, notonow rates 80.10/ and	Endovascular therapy or bypass	pooled data	2014(286)
				17

131				
assisted 75%, secondary 77%)		or IC	England)	
secondary 89%. CI I: primary 66% primary-	major amplitution at 1 v than ofs with CI I (2% vs	tissue loss or ischemic rest pain)	Group of New	
(IC: primary 79% primary-assisted 87%)	Results. Dts with IC experienced a lower rate of	indication of CLL (defined as	Wascular Study	22608039
<ul> <li>Graft patency was also significantly better in</li> <li>the IC grain when compared to the CI I grain</li> </ul>	1° endpoint: Amputation-free survival	Inclusion criteria: Elective and	Study type: NR	Simons JP, et al.
	groups.			2 i
	There was no difference in survival between the 2			
	<ul> <li>Overall survival was 93% at 1 y and 77% at 5 y.</li> </ul>	grafting		
	(p=0.01; p=0.02).	<ul> <li>Common femoral interposition</li> </ul>		
	Limb salvage was also lower in pts with RC 5 and 6	<ul> <li>Cross-femoral bypass</li> </ul>		
	<ul> <li>Secondary patency was 100% at both time points.</li> </ul>	bypass		
group B pts were claudicants	group B were 96% and 92%, respectively.	aorto-,axillo-, or iliotemoral	procedures	
• 113 (67%) of group A and 37/85 (40%) of	<ul> <li>Cumulative 5 y primary patencies for groups A and</li> </ul>	revascularization, including	pts/262	
(p=0.003).	Results:	<ul> <li>Hx of infrainguinal</li> </ul>	Size: n=230	
anticoagulation		Exclusion criteria:	2	
(p<0.0001), DM (p=0.04), and being on	(Group A) vs. CFE+distal revascularization (Group B)		single center	22944568
RC 5 or 6 (p<0.001), TASC D lesions	limb salvage; analysis stratified by use of CFE alone	CHE	retrospective,	2012(291)
<ul> <li>Predictors for distal revascularization were</li> </ul>	1° endpoint: Mortality, patency, reintervention, and	Inclusion criteria: Consecutive	Study type: NR	Malgor RD, et al.
	respectively).			
	Cl: 1.07–1.61 and OR: 1.29; 95% Cl: 1.04–1.61,			
	were found in the bypass group at 4 y (OR: 1.31; 95%			
	<ul> <li>Higher amputation free and overall survival rates</li> </ul>			
	0.55; 95% CI: 0.39–0.77) y of intervention.			
	second (OR: 0.60; 95% CI: 0.42–0.86) and third (OR;		CLI.	
	commonly in the endovascular group at the end of the		claudication or	
	<ul> <li>Progression to amputation was found to occur more</li> </ul>		pts with either	
	Cl: 1.12–1.97) y of intervention.		while 4 included	
	(OR: 2.03; 95% Cl: 1.20–3.45), and 3 (OR: 1.48; 95%		claudication only,	
	arm was found at 1 (OR: 2.42; 95% CI: 1.37–4.28), 2		study was	
	<ul> <li>Higher primary patency in the surgical treatment</li> </ul>		endovascular). 1	
	CI: 0.55–1.51).		1430	
	between the 2 groups were identified (OR: 0.92; 95%		139=87 open,	
offered as a first line interventional treatment.	surgery, whereas no differences in 30 d mortality	Exclusion criteria: N/A	studies (2,817 pts;	
term perspective a bypass procedure may be	failure (OR: 0.10; 95% CI: 0.05–0.22) than bypass		observational	
comorbidity, whereas for fit pts with a longer	(OR: 2.93; 95% CI: 1.34–6.41) and higher technical	femoropopliteal arterial disease	and 6	
be advisable in pts with significant	<ul> <li>Endovascular treatment had lower 30 d morbidity</li> </ul>	methods for the treatment of	Size: n=4 RCT	
lacking. An endovascular first approach may	Results:	percutaneous transluminal		<u>23159476</u>
superiority of one method over the other is		comparing open surgical and	analysis	2013(290)
Light lovel evidence demonstrating the	10 0000 01/A	Inclusion pritoria: Ottodion		Antonial CA at al
			claudication	

			21704539	2011(295)	Simó G, et al.			<u>21865062</u>	2011(294)	Kakkos SK, et al.														<u>22301210</u>	2012(293)	Siracuse JJ, et al.					
	procedures (79	<b>2</b>	retrospective)	(single center	Study type: NR	<u>Size</u> : n=269 pts (86 [32%] for IC)	2	retrospective)	(single center	Study type: NR											PTAS)	(113 bypass, 105	Size: n=218 nts	retrospective)	(single center	Study type: NR			(י טי [בטיט] וומט וט)	(797 [28%] had IC)	ota 200 C-a .c.: S
	<ul> <li>stenotic aorta and/or</li> </ul>	Long chronic CIA occlusion	Exclusion criteria: N/A		Inclusion criteria: SA-RIEA			Exclusion criteria: N/A		Inclusion criteria: AFB										-	Secondary procedures	Limb salvage procedure	Exclusion criteria:	ciaudication	procedures at single center for	Inclusion criteria: All LEB	<ul> <li>No specified indication</li> </ul>	disease	Rypage for applingemal		
	rates were 80.2%, /4./% and 69.3%; 84.8%,82.4% and 78.2%; and 86.8%, 84.2% and 79.6%,	primary, primary-assisted and secondary patency	Results: The 1, 3, and 5 y		1° endpoint: Patency		Dx other than IC; p=0.013)	Results: 60% survival at 10 y (vs. 42% for pts with		1° endpoint: Long-term survival, complications	<ul> <li>Recurrence of symptoms was similarly predicted by PTA/S (HR: 3.0; 95% CI: 1.8–5) and TASC D lesions (HR: 3.1; 95% CI: 1.4–7).</li> </ul>	lesions.	1.4–4.4) and TASC D (HR: 3.7; 95% CI: 3.5–9)	0.40).	last follow-up (83% vs. 49%; HR: 0.18; 95% CI: 0.08-	95% CI: 0.2–0.56), and freedom from symptoms at	(/3% vs. 42% at 3 y; HR: 0.4; 95% CI: 0.23–0./1),	Bypass showed improved freedom from restenosis	Results:		• Major amputation	Keinterventions	Symptom recurrence	Restenosis	Complications,	<u>1° endpoint:</u>					
132				iliofemoral reconstructive procedure	<ul> <li>10 pts required conversion to a conventional</li> </ul>		<ul> <li>No other results were stratified by Dx</li> </ul>	significant in multivariable model	survival vs. CLI or aneurysm Dx, but not	<ul> <li>IC associated with improved long-term</li> </ul>						vs. 0%; NS) or 3 y mortality (9% vs. 8%; NS).	93 % CI. U.30-U.30). ● No differences in perioperative mortality (2%)	freedom from recurrent symptoms (HR: 0.6;	patency (HR: 0.6; 95% CI: 0.35–0.97) and	Statin use postoperatively was predictive of	reintervention (77% vs 66% at 3 v. NS)	<ul> <li>There was no difference in freedom from</li> </ul>	(13%) in 2%; in 20 01 Indiana	• A (17% vs. 40%; p<0.01) and more for	<ul> <li>Bypass grafts were used less for TASC</li> </ul>	<ul> <li>Claudication-specific retrospective study</li> </ul>					

		Piazza M, et al. 2011(297) <u>21531527</u>			Sachs T, et al. 2011 (268) <u>21880457</u>		Eugster T, et al. 2011(296) <u>21850598</u>	
60% of hybrid repair pts were claudicants	Size: n=162 pts (248 limbs) 74% of	Study type: NR (single center retrospective)		<u>Size</u> : n=264,231 pts (claudication subaroup)	<u>Study type</u> : NR (NIS database 1997–2009)		Study type: NR (single center retrospective) Size: n=124 pts	[51%] had IC as indication)
<ul> <li>Exclusion criteria:</li> <li>Aortic thrombosis</li> <li>Abdominal aortic or iliac aneurysms</li> </ul>	and temoral reconstruction in pts with extensive iliac and common femoral occlusive disease	Inclusion criteria: Hybrid repair (combining iliac stenting and open CFE) or open aortoiliac	Exclusion criteria: N/A	peripheral bypass) or aortofemoral bypass	Inclusion criteria: Pts with ICD- 9 defined Dx atherosclerotic disease who underwent intervention of angioplasty stent,	Exclusion criteria: N/A	Inclusion criteria: Pts operated on for severe IC (walking distance\200 m) ≥y ago after failing nonoperative management	<ul><li>aneurysmal degeneration</li><li>Heavily calcified EIAs or</li><li>bilateral lesions</li></ul>
<b><u>Results:</u></b> • 30 d morbidity (3% vs. 5%, p=0.55) and mortality (1.1% vs. 1.4%, p=0.85) were equivalent between hybrid and open repair.	<ul> <li>Long-term patency</li> <li>Procedurally related limb salvage</li> <li>Overall survival</li> </ul>	<ul> <li><u>1° endpoint:</u></li> <li>30 d mortality and morbidity</li> <li>ABI increase</li> </ul>	<ul> <li>Average cost per procedure of PTA (w p 0.007)</li> <li>Average cost per procedure of PTA was higher than BPG for claudication (\$13,903 vs. \$12,681; p=0.02).</li> <li>Number of pts per y undergoing PTA for IC increased threefold (15,903 to 46,138)</li> </ul>	<ul> <li>Results:</li> <li>In-hospital mortality was similar for PTA and BPG proups for claudication (0.1%, vs. 0.2%; n=0.04)</li> </ul>	<u>1° endpoint</u> : Demographics, costs, and comorbidities, as well as multivariable adjusted in-hospital mortality and major amputation.	<ul> <li>Results:</li> <li>In-hospital and 30 d mortality of 0.8%; survival rate was 50.3% (SE±5.42%)</li> <li>Primary patency rate at 10 y was 63.5% (SE±7.50%)</li> <li>Assisted-primary patency rate was 87.3% (SE±5.19%)</li> <li>Patency rates of spliced and nonspliced vein bypasses were not different</li> </ul>	<ul> <li><u>1° endpoint:</u></li> <li>Survival</li> <li>Primary patency rate</li> <li>Assisted primary patency rate</li> </ul>	respectively
	<ul> <li>Multiple selective sub-group tests without</li> <li>Multiple stratified comparisons by dichotomized TASC classification</li> </ul>	<ul> <li>"Procedurally related" limb salvage is likely biased endpoint</li> <li>Reported 100% limb salvage rate is atypical</li> </ul>			N/A		<ul> <li>In-hospital and 30 d mortality of 0.8%</li> </ul>	

	Bailotta E, et al. 2010(300) <u>19828166</u>	2010(299) <u>20101647</u>	Derksen WJ, et al. 2010(298) <u>20167515</u> Konsister A of al	
	single center cohort)(Italy) Size: n=117 pts (121 procedures [60% of procedures were for claudication])	(retrospective case-control) <u>Size</u> : n=48 pts (24 matched pairs)	Study type: NR (prospective cohort) Size: n=90 pts (72 [80%] had IC)	
	<ul> <li>• CFA occlusive disease (isolated or with additional infrainguinal lesions in the ipsilateral limb)</li> <li>• Amenable to endarterectomy of the CFA (isolated or combined with a profundoplasty or with the endarterectomy of the superficial or deep femoral artery first tract, not &gt;1 cm long)</li> </ul>	peripheral arterial occlusive disease undergoing femoropopliteal supragenicular bypass or profundaplasty <u>Exclusion criteria</u> : None mentioned	Inclusion criteria: RSFAE performed TASC C/D SFA obstruction with or without an additional open CFE Exclusion criteria: N/A	<ul> <li>Concomitant visceral artery revascularization</li> <li>ALI</li> <li>Pts &lt;40 y with traumatic etiology for their disease from high performance sport (competitive cyclists).</li> </ul>
	<ul> <li><u>Presults:</u></li> <li>7 y PP, APP, and LS rates were 96%, 100%, and 100%, respectively</li> <li>The 7 y rates of freedom from further revascularization and survival were 79% and 80%, respectively.</li> </ul>	<ul> <li>Bypass occlusion</li> <li>Surgical revision</li> <li>Amputation</li> <li>Death</li> <li>Death</li> <li>Results: No significant outcome differences between supragenicular bypass surgery orprofundaplasty in pts who had surgery for IC</li> </ul>	<ul> <li><u>1° endpoint</u>: Restenosis following RSFAE</li> <li><u>Results</u>:</li> <li>57 pts (63%), a restenotic lesion was diagnosed within 12 mo.</li> <li>In multivariate analysis, age, duration of ischemic walking complaints, and lumen diameter before RSFAE were associated with increased restenosis</li> </ul>	<ul> <li>Primary patency of hybrid vs. open repair at 3 y was similar (91% vs. 97%; p=0.29) and was maintained after stratification by TASC A/B (89% vs. 100%; p=0.38) and TASC C/D (95% vs. 97%; p=0.54).</li> <li>Multivariate analysis for patency indicated that major tissue loss (Rutherford class 6) at presentation in the hybrid group was predictive of decreased long-term patency (p=0.02).</li> <li>Limb salvage at 3 y was 100% in both groups.</li> <li>Overall survival was 74% for OR vs. 40% for HR (p=0.007).</li> </ul>
134	• No comparison group		Complicated inclusion/exclusion criteria make generalization challenging	

Twine CP and McLain AD 2010(302) <u>20464717</u>	Burke CR, et al. 2010(301) <u>20122461</u>	
Study type: Cochrane systematic review Size: n=13 RCT with 2,313 pts (1955 above knee, 358 below knee bypasses)	Study type: NR (retrospective single center) <u>Size</u> : n=118 AFB and 174 aortoiliac angioplasty and AS procedures	
<u>Inclusion criteria</u> : Randomized trials comparing femoro-popliteal grafts. <u>Exclusion criteria</u> : N/A	Inclusion criteria: All pts undergoing treatment AIOD at the University of Michigan Hospitals between 1997–2007 Exclusion criteria: None mentioned	Exclusion criteria: Major tissue loss for which a contemporary infrainguinal revascularization was performed
<u>1° endpoint:</u> N/A <u>Results:</u> 7 graft types were compared (reversed and in situ autologous vein, PTFE with and without vein cuff, HUV, Dacron and HBD. Above the knee, there was a benefit in primary patency for autologous vein over PTFE (p=0.0001) and HUV (p=0.0003) by 60 mo. Dacron showed primary patency benefit over PTFE by 24 mo (p=0.02), continuing to 60 mo (p=0.02). HUV also showed benefit over PTFE by 24 mo (p=0.003) in 1 trial. Below the knee, in the 1 trial there was a significant benefit in primary patency for PTFE with a vein cuff when compared to PTFE alone at all time intervals to 24 mo (p=0.03). Limited data were available for limb survival. Antiplatelet and anticoagulant protocols varied extensively between	<ul> <li>1° endpoint:</li> <li>Mortality</li> <li>Mortality</li> <li>Adverse events</li> <li>Results:</li> <li>Long-term mortality, freedom from amputation, and freedom from revision procedure of any type (endovascular or open) were not different between groups.</li> <li>AFB was associated with increased surgical complication rates including the need for emergency surgery (6.8% and 1.7%; p=0.029), infection/sepsis (16.1% and 2.3%; p&lt;0.001), transfusion (16.1% and 5.7%; p=0.004), and lymph leak (8.5% and 0.6%; p=0.001).</li> <li>No difference between AFB and AS groups with respect to 30 d mortality (0.8% and 1.1%; p=0.64), MI (1.7% and 1.1%; p=0.35), or renal failure requiring hemodialysis (3.4% and 1.2%; p=0.19).</li> </ul>	
There was a clear primary patency benefit for autologous vein when compared to synthetic materials for above knee bypasses. In the long term (5 y) Dacron confers a small primary patency benefit over PTFE for above knee bypass. PTFE with a vein cuff improved primary patency when compared to PTFE alone for below knee bypasses. Further randomized data is needed to ascertain whether this information translates into improvement in limb survival.	<ul> <li>Large number of statistical comparisons without adjustment of significance level</li> <li>Not claudication specific (60 % of PTA and 41% of AFB pts had IC)</li> </ul>	

Goodney I 2009(305) <u>19497502</u>	Al-Khoury 2009(304) <u>19628359</u>	Chiesa R, et al 2009(303) <u>19540713</u>
Goodney PP, et al. 2009(305) 19497 <u>502</u>	Al-Khoury G, et al. 2009(304) <u>19628359</u>	<u>∪</u> 33), et al.
Study type: NR (prospective registry) (Vascular	<u>Study type</u> : NR (retrospective single center cohort) <u>Size</u> : n=95 pts (105 limbs); 65% of procedures done for IC	Study type: NR (retrospective single center cohort) Size: n=822 pts (777 [94%] had claudication as indication)
Inclusion criteria: LEB for arterial occlusive disease	Inclusion criteria: Pts who underwent an isolated femoral endarterectomy Exclusion criteria: N/A	Inclusion criteria: Consecutive pts undergoing aortoiliac or aortofemoral reconstruction employing a bifurcated ePTFE stretch graft Exclusion criteria:
1° endpoint: Predictors of ambulation status 1 y postoperatively	<ul> <li>1° endpoint:</li> <li>Change in ABI (based on cut-point of 15)</li> <li>Change in Rutherford class</li> <li>Repeat intervention</li> <li>Patency</li> <li>Results:</li> <li>83.8% of pts with marked initial clinical improvement remained symptom free at 2 y, whereas only 28.6% in the group with mild and moderate initial response maintained their clinical status.</li> <li>2 y freedom from repeat intervention was 61.8%.</li> <li>Multivariate analysis revealed that TASC C / D lesions (OR: 9.3, 95% CI: 1.01–13.15; p=0.048) were predictive of recurrent symptoms while extensive endarterectomy and ≥2 vessel tibial runoff decreased the need for repeat intervention.</li> <li>Patency was 100% with a mean follow-up of 11 mo (1–72).</li> <li>Complete resolution of symptoms was noted in 91% of limbs.</li> <li>ABI increase achieved in 85.1% with a mean ABI increase of 0.27±0.20, and this correlated with ≥2 runoff vessels (OR: 0.20; 95% CI: 0.04–0.96; p=0.04).</li> </ul>	trials, and in some cases within trials.          1° endpoint:         • Survival         • Graft-patency survival         • Amputation-free survival         • Amputation-free survival         • At the secondary graft-patency rate 90.6%         • The secondary rate patency rate was 97.9%
	WA	<ul> <li>Amputation-free survival only evaluated in subset of pts with CLI as indication</li> <li>Primary patency reported was for total 11 y duration of study period but mean follow-up of only 72 mo</li> <li>No survival analysis; descriptive analysis without models accounting time considerations</li> </ul>

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	<ul> <li>Clinical improvement was seen in 92% of pts.</li> </ul>			
	7%; p<0.01).			
	significantly higher primary patency (87% 5% vs. 53%			
	compared with bare stents was associated with			
	<ul> <li>By logistic regression analysis, use of stent grafts</li> </ul>			
	pts; inflow surgical procedures were required in 10%.			
			indication)	
	<ul> <li>Endovaccular reintervention was required in 1.1% of</li> </ul>	Exclusion criteria: N/A	claudication as	
	patencies were 60%. 97%. and 98% respectively.		(46% had	
	<ul> <li>5 y primary, primary-assisted, and secondary</li> </ul>	illotemoral occlusive disease	193 procedures	
	<ul> <li>30 d mortality was 2.3% and 5 y survival was 60%.</li> </ul>		$\underline{SIZe}$ : n=1/1 pts,	
		and ondovecular monodure for		
		single combined hybrid open	cohort)	
	adverse events, length of stay	stenting or stent grafting in a	retrospective	18572359
	(based on AHA classification), ABI change, patency,	patch angioplasty and primary	(single center	2008(306)
N/A	1° endpoint: Technical success, clinical success	Inclusion criteria: CFE with	Study type: NR	Chang RW, et al.
	being nonambulatory at 1 y.			
	were independently associated with a nigher risk of			
	here independently appropriated with a bight wind of			
	nrenneratively (HR: 3.5: 95% CI: 1.5-7 8: n<0.003)			
	<ul> <li>Graft thrombosis during follow-up (HR: 1.6; 95% CI:</li> </ul>			
	Cl: 1 G 1 1: m/0 001) and by major amputation (UD:			
	1.2–3.4: p<0.007). by postoperative MI (HR: 2.5: 95%			
	95% Cl: 1.5–3.7; p<0.0001), by CLI (HR: 2.0; 95% Cl:			
	1.8; 95% CI: 1.2–2.6; p<0.007) and 80-89 y (HR: 2.3;			
	1.6; p<0.0001), by increasing age of 70–79 y (HK:			
	$1 \text{ G}_{1} \approx 1000001$ km increasing and $1 \text{ G}_{1} \approx 10000000000000000000000000000000000$			
	nonambulatory preoperatively (HR-1.5-95% CI-1.3–			
	postoperatively was increased in pts who were			
	<ul> <li>The risk of dying or being nonambulatory 1 y</li> </ul>			
	ambulatory.			
	95% who survived 1 y postoperatively remained			
	<ul> <li>All claudicant pts walked before surgery, and the</li> </ul>			
	claudicant pts (p<0.001).			
	<ul> <li>Amputation rates were 12% for CLI pts and 1% for</li> </ul>		(%CZ	
	p<0.001) than CLI pts.		was indication for	
	and ambulatory I y postoperatively (90% vs. 81%;			
	gratt patency rates and were more likely to be alive		4F64 bimpion (IC	
	p<0.001) and secondary (87% vs. 81%; p<0.001)		2	
	<ul> <li>Claudicant pts had higher primary (79% vs. 73%;</li> </ul>		New England)	
	Results:	EXCIUSION CRITERIA: IN/A	Study Group of	
			> - >	

Jaquinandi V, et al. 2007(307) <u>17264010</u>	KoivunenK and Lukkarinen H <u>18221916</u>
<u>Study type</u> : NR, prospective <u>Size</u> : n=105 pts	<u>Study type:</u> NR, prospective <u>Size:</u> n=180 pts (64 conservative, 85 endovascular, 31 surgery)
Inclusion criteria:         Age ≥18 y         Had a patent AFB for ≥4 mobefore his or her visit         Able to walk on treadmill         Exclusion criteria:         Acute CLI         Uncontrolled hypertension         New York Heart Association (NYHA) cardiac insufficiency function class of III or IV         MI ≤3 mo         Arterial aneurysm         Major respiratory limitation (resting dyspnea)         Stroke or major neurologic disorders         Lived too far from the	Inclusion criteria: IC (Fontaine II), surgery clinic pt at university hospital in Finland Exclusion criteria: Nonatherosclerotic disease, lack of angiographic verification of Dx, previous surgery/endovascular treatment <5 y, CLI
<u>1° endpoint</u> : Symptoms based on modified San Diego Claudication questionnaire, change in TcPO ₂ before and after treadmill ambulation <u>Results</u> : 30 pts reported proximal exercise-related pain consistent with vascular criteria by Hx before exercise. However, 59 pts (56%) reported symptoms compatible with proximal claudication, and TcPO ₂ values were abnormal on one or both sides in 52. The persistence of at least one (prograde or retrograde) pathway to the hypogastric circulation did not decrease proportion of pts reporting proximal claudication (33% by Hx; p=0.51 compared with at least one prograde hypogastric pathway and 61% based on treadmill test, p=0.65 compared with at least one prograde hypogastric pathway).	<ul> <li>Mean ABI increased from 0.38 0.32 to 0.72 0.24.</li> <li>Median length of stay was 2 d (range, 1–51 d).</li> <li><u>1° endpoint</u>: HRQoL (Nottingham Health Profile)</li> <li>Conservative group's clinical outcomes (ABI, asx walking distance) remained stable, while these measures improved significantly in the surgery group</li> <li>Conservative group had improved quality of sleep and emotional reactions</li> <li>Endo group had significant improvement in emotional reactions and energy + reduction in social isolation. No significant changes in pain or mobility</li> <li>Surgery group had improvements in sleep, pain, emotional reactions, social isolation, and physical mobility</li> <li>Large effect size for surgery vs. small for conservative, endo</li> </ul>
	<ul> <li>Pts treated with conservative approach exercised more often at baseline</li> <li>Surgery group had more baseline hypertension</li> <li>Smoking increased significantly in conservative management group</li> </ul>

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	table analysis was 60.6±4.8% (SE) at 33 mo, (mean 17.1 mo: range 133 mo)	lesion	Size: n=210 pts (158 [75%] were	
<ul> <li>12 pts (5.7%) had wound complications</li> </ul>	<ul> <li>Results:</li> <li>Primary cumulative patency rate by means of life-</li> </ul>	aSpire stenting for TASC D SFA	multicenter cohort)	<u>16953157</u>
<ul> <li>Did not stratify results by diagnostic indication</li> </ul>	<u>1° endpoint</u> : Primary cumulative patency	Inclusion criteria: Remote superficial femoral	Study type: NR (retrospective	Rosenthal D, et al. 2006(310)
		<ul> <li>Repeat inclusion of bypasses</li> <li>Unreliable or unattainable reconstruction of life tables from graphs or texts.</li> </ul>		
		<ul> <li>Predominance of composite bypass grafts</li> <li>Predominance of bypasses to the infrapopliteal arteries</li> </ul>		
	secondary graft patency was 73.2%, 80.1%, and 79.7%, respectively (p>0.05).	<ul> <li>Exclusion criteria:</li> <li>Clinical symptoms not described</li> <li>Predominance of blind</li> </ul>		
	3, 4, and 5 y (p<0.05). The corresponding pooled	publication after 1986	specific subgroup	
possible.	pooled primary graft patency was 57.4% for above- knee polytetrafluoroethylene, 77.2% for above-knee	minimum of 30 bypasses in at least 1 series when article	Size: n=73 articles included; analysis	
The great saphenous vein performs better than polytetrafluoroethylene in femoropopliteal bypass graffing and should be used whenever	<u>1° endpoint</u> : Pooled primary graft patency Results: For claudication-specific meta-analysis	Inclusion criteria: graft patency included as outcome, follow up of 1 v for at least some grafts.	<u>Study type</u> : Meta- analysis	Periera CE, et al. 2006(309) 16950427
	primary patency was significantly higher in the bypass group after 12 mo (OR: 1.6; 95% CI: 1.0–2.6) but not after 4 y (p=0.14). Blood flow restoration was significantly greater in bypass than in thromboendarterectomy pts (Peto OR: 9.2; 95% CI: 1.7–50.6); mortality and amputation rates did not differ. Bypass surgery outcomes did not differ differ. Bypass surgery outcomes did not differ	<u>Exclusion criteria</u> : N/A	SIZE: n=19 trials (2 claudication only, 4 with claudication and CLI) and CLI)	
of bypass surgery compared with other treatments; no studies compared bypass to no treatment. Further large trials are required.	Results: Mortality and amputation rates did not differ significantly between bypass surgery and PTA;	bypass surgery for chronic lower limb ischemia vs. any other treatment	Systematic review (Cochrane)	GC 2008(308) <u>18425879</u>
There is limited evidence for the effectiveness	1º androint: N/A	laboratory. Inclusion criteria: RCTs of	Study type:	Fowkes F and I end

		<u>9507581</u>	Pell JP and Lee AJ 1997(266)						10642712	Feinglass J, et al 2000(263)						<u>11821823</u>	Mori E, et al. 2002(312)					16476609	Martin JD, et al. 2006(311)	
	<u>Size</u> : n=201 pts	observational)	Study type: NR (prospective,	angioplasties)	bypasses and 44	revascularization,	<u>Size</u> : n=526 pts (104 had		observational)	Study type: NR (prospective,		conservative=168]	legs)	Size: n=427 pts	2	observational)	Study type: NR		(57% had IC)	<b>Size:</b> n=133 nts	cohort)	single center	<u>Study type</u> : NR (retrospective	claudicants)
		Exclusion criteria: N/A	Inclusion criteria: newly referred pts with IC	• Gangrene	Rest pain     Ulcers	<ul> <li>Prior revascularization</li> </ul>	Exclusion criteria:	symptoms	revascularization or CLI	ABI without prior LE					Exclusion criteria: N/A		Inclusion criteria: Admitted to the hospital for IC	Exclusion criteria: N/A	arteries.	total occlusions of the external lian and/or superficial femoral	reconstruction of >10 cm length	incision for vascular	Inclusion criteria: Remote endarterectomv from an inquinal	Exclusion criteria: N/A
in pain and physical function after adjustment for case	Conservative treatment	<ul> <li>All aspects of QoL deteriorated following</li> </ul>	<u>1° endpoint</u> : QoL (SF-36)	<ul> <li>Mean ABI improved significantly for bypass, modestly for angioplasty</li> </ul>	<ul> <li>Conditions of unmatched medical management pts</li> <li>declined on all outcome measures</li> </ul>	symptom improvement	significant improvements in mean physical function and walking distance scores, and reported greater leg	Bypass and angioplasty groups maintained highly	Results:	<u>1° endpoint</u> : SF-36 physical functioning score			<ul> <li>Intrainguinal and conservative were not significantly different</li> </ul>	improvement than conservative	Surgery group had significantly better QOL	Results:	<u>1° endpoint:</u>			salvage was 94%.	patency of 70% at 30 mo by life-table analysis. Limb	Results: Mean follow-up was 19 mo, with a primary	<u>1° endpoint</u> : Primary patency	<ul> <li>During follow-up percutaneous transluminal balloon and/or stent angioplasty was necessary in 50 pts for a primary assisted patency of 70.2±4.8% at 33 mo.</li> <li>Mean ABI rose from 0.58–0.95</li> </ul>
defined beyond lack of procedural intervention 140	<ul> <li>76% managed conservatively</li> <li>"Conservative management" was not</li> </ul>	∘10% had PTA ∘10% had reconstruction	<ul> <li>F/U data available on 81% of 195 pts alive at final timepoint.</li> </ul>			<ul> <li>No mention of exercise therapy</li> </ul>	<ul> <li>Does not include adverse event rates</li> <li>No standardized medical management</li> </ul>	admission or separate hospitalizations)	if procedures were staged within a single	<ul> <li>Pts who underwent angioplasty and surgery were classified as surgical bypass (regardless</li> </ul>	rates	Does not report adverse events, amputation	No defined pharmacounerapy     No exercise comparator	results	revascularization may be overinterpretation of	<ul> <li>Recommendation for surgical</li> </ul>	<ul> <li>Inferior 3 and 5 y patency observed for</li> <li>below knee bypass</li> </ul>						<ul> <li>12% technical failure rate (bypass performed in these pts)</li> </ul>	

5,209 pts at the initial examination; of these 4,030	1° endpoint: Incidence of claudication by age and sex	Inclusion criteria: General	Study type: NR	Kannel WB et al.
	of 15.5 per 1,000 person-years)	Exclusion criteria: N/A	<u>Size</u> : n=1,592 pts	
surgery or amputation, and 1.4% developed leg ulcers.	Results:116 new cases of claudication identified (incidence	registers of 10 general practices in Edinburgh, Scotland	study (Edinburgh Artery Study)	9027521
Among those with baseline claudication, 28.8% still had pain after 5 yr, 8.2% underwent vascular	<u>1° endpoint</u> : Incidence and natural hx of claudication; incidence of CV events in sx and asx PAD.	Inclusion criteria: Age 55–74 y selected randomly from the age-sex	Study type: Prospective cohort	Leng GC, et al. 1996(317)
			grafts in 578 legs; 85% were for CLI.	
	Initial bypass indication. Early graft failure (<30 d) nad worse prognosis.		infrainguinal bypass	
tissue loss and those with early graft failure."	50% at 2 yr. Limb salvage was 100% among pts with IC as			<u>15111843</u>
"The overall prognosis for limb salvage in pts with failed infrainguinal bypass grafts is poor,	<u>1º endpoint</u> : Limb salvage	Inclusion criteria: N/A	<u>Study type:</u> Retrospective single	Baldwin ZK, et al. 2004(316)
• 25 limbs had acute ischemia			(none had IC)	
All pts had CLI		Exclusion criteria: N/A	Size: n=211 grafts in	
<ul> <li>Primary bypass procedures had superior</li> </ul>	Results: 5 yr cumulative limb salvage was 51%	limb salvage	center)	<u>8230575</u>
<ul> <li>5 yr primary/secondary patency 23% / 25%</li> </ul>		(tibial) PTFE grafts implanted for	(retrospective single	1993(315)
• 7 vr primary/secondary patenny 37% / 15%	1° endpoint: 5 vr cumulative limb salvane	Inclusion criteria: See above	Study type: NR	Schweiner H et al
		Exclusion criteria: Cap above		
		not duplicate other published material.		
	for claudication.	the distribution of covariates, and	analysis of patency	
	primary patencies after surgery varied from 33%-80% with	errors, define patency as	studies were included in life table	
	49% for bypass surgery using PTFE graft. Adjusted 5 yr	the number at risk or standard	popliteal bypass	
analysis was stratified based on indication.	Results: Unadjusted pooled 5 yr patency was 45% tor and and oblights and a vein graft, and	table or Kaplan-Meier analysis with	Size: n=17 femoral-	<u>8152359</u>
CLI/limb salvage as well as claudication, but		language articles had to report	(meta-analysis)	1994(314)
Pooled data included bypasses performed for	1° endpoint: Patency	Inclusion criteria: English	Study type: NR	Hunink MG, et al.
<ul> <li>Overall PTFE failure rate was 3–4 times higher than that of GSV.</li> </ul>		Exclusion criteria: N/A	bypasses in 285 pts (39% had IC as indication)	
<ul> <li>Above-knee GSV primary patency &gt;below-knee GSV &gt;above-knee PTFE.</li> </ul>	Results: GSV patency superior to PTFE at 3 and 5 yr; P<0.01.	reversed GSV when available and PTFE when not.	center)	7811585
<ul> <li>Patency for GSV vs. PTFE was 87% vs. 54% at 3 yr and 81% vs. 48% at 5 ys.</li> </ul>	<u>1° endpoint</u> : Patency	Inclusion criteria: Femoropopliteal bypass using ipsilateral autologous	<u>Study type</u> : NR (retrospective, single	Archie JP Jr 1994(313)
<ul> <li>No defined pharmacotherapy</li> <li>No exercise therapy comparison group</li> </ul>	mix			

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	symptoms after presentation, and 1.5–5% had major amputation.			
	Results: Reported prevalence of claudication in general population ranges from 0.4%–6.9% in men and 0.2%–3% in women. 25% of pts with claudication had worsening of	Exclusion criteria: Publications based on small numbers of pts or inconclusive data	Size: n=52 studies published between 1958–1986	<u>2041/101</u>
N/A	<u>1° endpoint</u> : Fate of pts presenting with chronic leg ischemia	Inclusion criteria: English language published data	<u>Study type</u> : NR (Review)	Dormandy J, et al. 1989 (323)
	Results: N/A	Exclusion criteria: N/A	Size: N/A	<u>19310276</u>
N/A	<u>1° endpoint</u> : N/A	Inclusion criteria: N/A	Study type: Topic overview	Bloor K 1961(322)
<ul> <li>clinical physiology at a single hospital over 1 y.</li> <li>At a mean follow up of 6.5 ± 0.5 yts, 44% of pts had died.</li> </ul>	Kesults: /.5% rate or progression in the worst affected leg during first yr after referral; 2.2% per yr thereafter.	Exclusion criteria: Rest pain, ulcers, or foot gangrene.		
<ul> <li>Unclear whether design was prospective or retrospective.</li> <li>Recruitment occurred from the department of</li> </ul>	<u>1° endpoint:</u> Rate of clinical progression (to rest pain or gangrene).	Inclusion criteria: Pts referred consecutively for the first time for claudication during a 1 y period.	Study type: NR Size: n=257 pts	Jelnes R, et al. 1986(321) <u>3094806</u>
<ul> <li>Authors concluded that "the course of the disease in the lower limbs does not affect life expectancy to any considerable extent."</li> </ul>		peripheral arterial insufficiency that appeared in the final stage of a severe disease (e.g., heart failure or cancer).	2 - - -	
<ul> <li>Crassified pts with DW separate from those with atherosclerosis.</li> <li>Included pts with CLI but did not stratify results in a similar fashion.</li> </ul>	attributed to arteriosclerosis had an amputation during the observation period. Amputation rate among this subgroup was 2.24/1000 mo for men and 1.23/1000 mo for women.	causing a suspicion of arterial insufficiency	<u>Size</u> : n=466 pts	<u>143   7320</u>
Study included pts suspected to have Beurger's disease.	<u>1° endpoint</u> : Survival, amputation, adverse CV events.	Inclusion criteria: Pts treated at hospitals in Stockholm for complaints in the lower limbs	<u>Study type</u> : NR (retrospective)	Tillgren C 1965(320)
<ul> <li>cardiovascular impairments, and terminates tatally".</li> <li>Significant overlap with Kannel 1970 (making it challenging to identify distinct findings within this report).</li> </ul>	attributatie to impaired leg circulation. A total or o amputations occurred. Among those followed for ≥4 y from onset of claudication symptoms, 45% had their symptoms disappear for at least 4 y	Exclusion criteria: None stated	אסליכי-וו :	<u>5119838</u>
<ul> <li>Purpose of study was "to examine in a general population the manner in which IC arises, evolves, and becomes complicated by more serious</li> </ul>	<u>1° endpoint</u> : Adverse cardiovascular events, mortality <u>Results</u> : No death in the study group was directly	Inclusion criteria: General population of adult men and women (Framingham; 16 y follow up)	Study type: NR (prospective cohort)	Kannel WB and Shurtleff D 1971(319)
analysis.	<b>Results:</b> 79 men and 46 women developed claudication. Overall annual incidence per 10,000 was 26 for men and 12 for women. No death was attributable to impaired limb circulation, and no amputation related to circulatory diseased occurred over 14 yr study period.	(Framingham; 14 y follow up) (Framingham; 14 y follow up) Exclusion criteria: None stated	Size: n=5,209 pts	5444530
returned for the 8 examination covered in this		population of adult men and women	(prospective cohort)	

National Surgical Quality Improvement Program, NS, not significant; NYHA, New York Heart Association; OR, odds ratio; PAD, peripheral artery disease; PCIS, percutaneous iliac stent; PP, primary patency; PTAS, percutaneous angioplasty/stent; PTFE, polytetrafluoroethylene; pt, patient; QoL, quality of life; RC, routine care; RCT, randomized controlled trial; RR, relative risk; RSFAE, remote superficial artery endarterectomy; SA RIEA, Stent-assisted remote iliac endarterectomy; SE, supervised exercise; SFA, superficial femoral artery; SIA, subintimal angioplasty; TASC, transatlantic inter-society consensus; and TcPO₂, transcutaneous oxygen pressure. greater saphenous vein; HBD, heparain bonded Dacron; HR, hazard ratio; HRQoL, heath-related quality of life; HUV, human umbilical vein; ICD, International Classification of Disease; ischemia; COPD, chronic obstructive pulmonary disease; CV, cardiovascular; EIA, external iliac artery; ePTFE, expanded polytetrafluoroethylene; EVT, endovascular treatment; GSV, Society of Anesthesiologist; BPG, bypass graft; CFA, common femoral artery; CFE, common femoral endarterectomy; CIA, common iliac artery; CI, confidence interval; CLI, critical limb aortobifemoral bypass; AHRQ, Agency for Healthcare Research and Quality; AIOD, aortoiliac occlusive disease; APP, assisted primary patency; AS, aortoiliac stenting; ASA, American ABF indicates aortobifemoral; ABI, ankle-brachial index; ALI, acute limb ischemia; ACS NSQIP, American College of Surgeons National Surgical Quality Improvement Program; AFB, IC, intermittent claudication; LEB, lower extremity bypass; LE, lower extremity; LS, limb salvage; N/A, not applicable; NIS, National Impatient Sample; NR, nonrandomized; NSQIP,

Evidence Table	39. RCTs Comp	Evidence Table 39. RCTs Comparing Endovascular Revascularization for Chronic CLI–Section 8.2	ation for Chro	nic CLI–Section 8.2.	
Study	Aim of Study;	Patient Population	Study	Endpoint Results	Relevant 2° Endpoint (if any);
Acronym;	Study Type;		Intervention	(Absolute Event	Study Limitations;
Author; Year Published	Study Size (N)		(# patients) /	Rates, P value; OR or	Adverse Events
			Comparator	95% CI)	
			(# patients)		
Werk M, et al.	Aim: SFA DCB	Inclusion criteria: Sx femoro-popliteal	Intervention:	1° endpoint: The	DEB is superior to PTA
2012(232)	vs. PTA	atherosclerotic disease	DCB	primary endpoint was	<ul> <li>Pts with sx femoro-popliteal atherosclerotic disease</li> </ul>
<u>23192918</u>				late lumen loss at 6	undergoing percutaneous transluminal angioplasty were
	Study type:	Exclusion criteria:	Comparator:	mo assessed by	randomized to paclitaxel-coated IN.PACT Pacific or
	RCT	<ul> <li>Acute thrombus or aneurysm in the</li> </ul>	PTA	blinded angiographic	uncoated Pacific balloons. The primary endpoint was late
		target vessel		corelab quantitative	lumen loss at 6 mo assessed by blinded angiographic
	Size: n=85 pts	<ul> <li>Failure to cross the target lesion with</li> </ul>		analyses	corelab quantitative analyses. Secondary endpoints were
		a guidewire			binary restenosis and Rutherford class change at 6 mo,
		<ul> <li>Inflow lesions that cannot be</li> </ul>			and target lesion revascularization + major adverse
		successfully pretreated			clinical events (major adverse events=death, target limb
		<ul> <li>Significant disease of all 3</li> </ul>			amputation, or target lesion revascularization) at 6 and 12
		infrapopliteal vessels			mo. 85 pts (91 cases=interventional procedures) were
		<ul> <li>Renal failure (serum creatinine &gt;2.0</li> </ul>			randomized in 3 hospitals (44 to DEB and 47 to uncoated
		mg/dL)			balloons). Average lesion length was 7.0±5.3 and
		<ul> <li>Known intolerance or allergy to study</li> </ul>			6.6±5.5 cm for DEB and control arm, respectively.
		medication			Procedural success was obtained in all cases. 6 mo
		<ul> <li>Life expectancy &lt;2 v</li> </ul>			quantitative angiography showed that DEB were
					associated with significantly lower late lumen loss (-0.01
					mm; 95% Cl: -0.29–0.26 vs. 0.65 mm; 95% Cl: 0.37–
					0.93; p=0.001) and fewer binary restenoses (3 [8.6%] vs.
					11 [32.4%]; p=0.01). This translated into a clinically

Size		ABSOLUTE Aim: SF	IN.PACT       Aim: SF         Tepe G, et al.       vs. PTA         2015(229)       Study ty         25472980       RCT         RCT       RCT         Size: n=       1
Size: n=104 pts	<u>/pe:</u>	A PTAS	A DCB 331 pts
<ul> <li>Previous bypass surgery, or stenting of the SFA</li> <li>Untreated inflow disease of the ipsilateral pelvic arteries (&gt;50% stenosis or occlusions)</li> </ul>	•ALI	Inclusion criteria: Rutherford 3–5 and SFA stenosis	Inclusion criteria:       IC or ischemic rest         pain attributable to superficial femoral         and popliteal PAD         Exclusion criteria:         • Lesion and/or occlusions located in or         extending to the popliteal artery or         below the ankle joint space         • Inflow lesion or occlusion in the         ipsilateral iliac, SFA, or popliteal arteries         with length ≥ 15 cm         • Significant (≥50% DS) inflow lesion or         occlusion in the ipsilateral iliac, SFA, or         popliteal arteries left untreated         • Previously implanted stent in the TL(s)         • Aneurysm in the target vessel. Acute         thrombus in the TL
	Comparator: PTA	Intervention: PTAS	Intervention: DCB <u>Comparator:</u> PTA
	at 2 y	<u>1º endpoint:</u> Restenceis hy dunley	<u>1° endpoint</u> : 12 mo primary patency
duplex ultrasound and for clinical and hemodynamic outcome by treadmill walking distance and ABI. Restenosis rates at 2 y were 45.7% (21 of 46) vs. 69.2% (36 of 52) in favor of primary stenting compared with balloon angioplasty with optional secondary stenting by an ITT analysis (p=0.031). Consistently, stenting (whether primary or secondary; n=63) was superior to plain balloon angioplasty (n=35) with respect to the occurrence of restenosis (49.2% vs. 74.3%; p=0.028) by a treatment-received analysis. Clinically, pts in the primary stent group showed a trend toward better	<ul> <li>Of 104 pts with chronic limb ischemia and SFA obstructions, 98 (94%) could be followed up until 2 y after intervention for occurrence of restenosis (&gt;50%) by</li> </ul>	PTAS is superior to PTA for long lesions (lesion length     112 mm PTAS and 93 mm PTA)	<ul> <li>relevant benefit with significantly fewer major adverse events for DEB vs. uncoated balloons up to 12 mo (3 [7.1%] vs. 15 [34.9%]; p&lt;0.01) as well as target lesion revascularizations (3 [7.1%] vs. 12 [27.9%]; p=0.02).</li> <li>OCB superior to PTA</li> <li>The IN.PACT SFA Trial is a prospective, multicenter, single-blinded, randomized trial in which 331 pts with IC or ischemic rest pain attributable to superficial femoral and popliteal PAD were randomly assigned in a 2:1 ratio to treatment with DCB or PTA. The primary efficacy endpoint was primary patency, defined as freedom from revascularization at 12 mo. Baseline characteristics were similar between the 2 groups. Mean lesion length and the percentage of total occlusions for the DCB and PTA arms were 8.94±4.89 and 8.81±5.12 cm (p=0.82) and 25.8% and 19.5% (p=0.22), respectively. DCB resulted in higher primary patency vs. PTA (82.2% vs. 52.4%; p&lt;0.001). The rate of clinically driven target lesion revascularization was 2.4% in the DCB arm in comparison with 20.6% in the PTA arm (p&lt;0.001). There was a low rate of vessel thrombosis in both arms (1.4% after DCB and 3.7% after PTA [p=0.10]). There were no device- or procedure-related deaths and no major amputations</li> </ul>

	2013(230) 24239203	<b>DEBATE-SFA</b> Liistro F, et al.			Gandini K, et al. 2013(324) 24325697											17592075	et al.	<b>FASI</b> Krankenberg H,		
RCI	Study type:	<u>Aim</u> : PEB+BMS vs. PTA+BMS	RCT <u>Size</u> : n=448 pts	Study type:	AIM: CLI & SFA ISR: DCB vs. laser+DCB									eid ++7-11 - <u>3710</u>	Cize: n=011 nte	RCT	Study type:	AIM: SFA PTA vs. PTAS		
<ul> <li>Life expectancy &lt;1 y</li> </ul>	Exclusion criteria:	Inclusion criteria: Claudication or CLI and SFA stenosis		Exclusion criteria: Denovo stenosis without ISR	in-stent occlusion	<ul> <li>Ungoing dialysis treatment</li> <li>Treatment with oral anticoagulants other than antiplatelet agents.</li> </ul>	stenosis	<ul> <li>Untreated iosilateral iliac artery</li> </ul>	<ul> <li>Acute or subacute (≤4 wk) thrombotic</li> </ul>	<ul> <li>Multiple lesions exceeding a total length of 10 cm</li> </ul>	targeted SFA	<ul> <li>Previous stent implantation in the</li> </ul>	• A TL that extended into the popliteal	debulking catheters	A TL that required pretreatment with	criteria were:	Exclusion orthonia: Major avaluation	Inclusion criteria: SFA stenosis & claudication or CLI		
PTA+BMS	Comparator:	Intervention: PEB+BMS		<u>Comparator</u> : DCB	Intervention: Laser+DCB											PTA	Comparator	Intervention: PTAS		
		<u>1° endpoint</u> : 12 mo binary restenosis			1° endpoint: 12 mo primary patency												duplex restenosis	<u>1° endpoint:</u> Technical success, 1 y		
of lesions in the PEB+BMS and PTA+BMS groups 145	PEB+BMS and PTA+BMS groups (p=0.8), respectively. The primary endpoint occurred in 9 (17%) vs. 26 (47.3%)	<ul> <li>PEB+BMS is superior to PTA+BMS</li> <li>Mean lesion length was 94±60 vs. 96±69 mm in the</li> </ul>	Laser+DEB group and 50% in the DEB only group (p=0.01). 2 (8%) pts needed major amputations in the Laser+DEB group vs. 11 (46%) in the DEB only group at 12 mo (p=0.003).	higher (p=0.01) than in the DEB only pts (58.3% and 37.5%, respectively). TLR at 12 mo was 16.7% in the	<ul> <li>Laser+UEB superior to UEB alone</li> <li>In the Laser+DEB group, the patency rates at 6 and 12 mo (91.7% and 66.7%, respectively) were significantly</li> </ul>			mo was significantly lower for pts with stent tractures (41.1% vs. 84.3%, p<0.0001).	Kaplan-Meier estimates, the primary patency rate at 12	site of stent fracture. In 22 cases (34.4%) with stent fracture there was a total stent reocclusion. According to	there was a restenosis of >50% diameter reduction at the	lengtn ≤6 cm, 42.4% for stented lengtn >6–16 cm, and 52.0% for stented length >16 cm. In 21 cases (32.8%)	cases (25.0%). Fracture rates were 13.2% for stented	Ridderate (fracture of > 1 Strut) in 17 cases (20.0%), and severe (complete separation of stent segments) in 16	as minor (single strut fracture) in 31 cases (48.4%),	stents (24.5%) showed fractures, which were classified	Overall, stent fractures were detected in 45 of 121	<ul> <li>For short lesions mean length 45mm, no difference between PTAS and PTA</li> </ul>	p=0.09) at 2 y, respectively. Reintervention rates tended to be lower after primary stenting (17 of 46 [37.0%] vs. 28 of 52 [53.8%]; p=0.14)	treadmill walking capacity (average, 302 vs. 196 m; p=0.12) and better ABI values (average, 0.88 vs. 0.78;

rates
revascularization, index-limb amputation
patency death, repeat
angiographic restenosis vessel
<u>1° endpoint</u> : 1 y
TLR.
amplifation and CD-
was a composite of all-
primary safety
Safety endpoint: The
loss (LLL).
TLR) and late lumen
driven target lesion
1° endpoint: Clinically

2016(327) 26777329 BASIL Katsanos K, et ACHILLES 2005 (328) Adam DJ, et al. 16325694 Aim: Infrapop: DES vs. PTA Study type: RCT Study type: RCT PTA for CLI Size: n=452 pts Aim: Bypass vs. Size: n=200 pts vascular surgeon and interventional infrainguinal PAD of the knee joint or in an artery subject Stent placement across or within 1 cm Known intolerance to antiplatelet Life expectancy <12 mo</li> Hx of thrombophlebitis, deep venous Prior stenting within the target to external compression Infrapopliteal trifurcation lesions popliteal artery Common or superficial femoral and the common or external iliac Untreated lesions (>75% stenosis) in bypass or angioplasty in the opinion of a Exclusion criteria: trial above thrombosis, or impaired renal function popliteal artery vessel(s) or aneurysm in the SFA or requiring 2- or 3-branch treatment radiologist treated equally well with infrainguinal Exclusion criteria: Pt who could not be Refer to ACHILLES trial above (Cr >2.5 mg/dl) nclusion criteria: CLI due to medication. nclusion criteria: Refer to ACHILLES <u>Comparator</u>: PTA Intervention: DES Bypass Comparator: ΡΤΑ Intervention: angiographic patency death, repeat survival Amputation free 1° endpoint: rates index-limb amputation revascularization, restenosis vessel <u>1° endpoint</u>: 1 y angioplasty underwent an attempt at their allocated bypass surgery and 216 (96%) of 224 to balloon • The trial ran for 5.5 y, and follow-up finished when pts p=0.08) in the whole study and 0.17 (95% CI: -0.03-0.35) compared with PTA up to 1 y after randomization. amputation, and 130 (29%) dead without amputation. 38 (8%) alive with amputation, 36 (8%) dead after 248 (55%) pts were alive without amputation (of trial leg) after randomization, respectively. At the end of follow-up, intervention at a median (IQR) of 6 (3–16) and 6 (2–20) d first y of follow-up. 195 (86%) of 228 pts assigned to these, 3 were lost (1 angioplasty, 2 surgery) during the randomization (3 assigned angioplasty, 2 surgery); of ankle or death). 7 individuals were lost to follow-up after reached an endpoint (amputation of trial leg above the Equal outcomes p=0.09) in the wound subgroups comparison. Relative QALY gain was 0.10 (95% CI: -0.01-0.21; There was a trend of more QALYs gained with SES superior to PTA for CLI Infrapop SES axcellerates wound healing and is ES After 6 mo, the 2 strategies did not differ significantly in

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	24456716	2014 (231)	LEVANT 1 Schienert D, et al				2014 (330)	W, et			20307380	al. 2010 (329)	BASIL Bradbury AW, et		
<u>Size</u> : DEB=49 pts; Standard PTA=52 pts	<u>Study type:</u> RCT	bailout stenting	<u>Aim</u> : Assess efficacy of DEB vs_PTA with	Size: n=452 pts	RCT	analysis of a	Study type: ITT	<u>Aim</u> : Bypass vs. angiography for CLI		Size: n=452 pts	RCT	Study type:	<u>Aim</u> : Bypass vs. PTA for CLI		
	<ul> <li>Listed in methods</li> <li>Notably highly calcified lesions</li> </ul>	Exclusion criteria:	Inclusion Criteria: Rutherford 2–5 symptoms		vascular surgeon and interventional radiologist	bypass or angioplasty in the opinion of a	Exclusion criteria: Pt who could not be	Inclusion criteria: CLI due to infrainguinal PAD	radiologist	bypass or angioplasty in the opinion of a vascular surgeon and interventional	treated equally well with infrainguinal	Exclusion criteria: Pt who could not be	Inclusion criteria: CLI due to infrainguinal PAD		
	Standard PTA with bailout stenting	Comparator:	Intervention: DEB			n Abass	Comparator:	Intervention: PTA			Bypass	Comparator:	Intervention: PTA		
	lower lumen loss than standard PTA (p<0.016)	At 6 mo DEB had	<ul> <li><u>1° endpoint:</u></li> <li>Angiography lumen</li> </ul>					<u>1° endpoint</u> : AFS and OS					<u>1° endpoint</u> : AFS		
			Small study					Bypass was associated with improvements in OS and AFS of about 7 and 6 mo, but long term no significant difference between the treatments					N/A	1.07; 95% CI: 0.72–1.6; adjusted HR: 0.73; 95% CI: 0.49–1.07). We saw no difference in health-related quality of life between the 2 strategies, but for the first y the hospital costs associated with a surgery-first strategy were about 1/3 higher than those with an angioplasty-first strategy.	amputation-free survival (48 vs. 60 pts; unadjusted HR:

	classification after 1 y		≤44 mm in length	Prospective,	
	Rutherford-Becker		that was 2.5–3.5 mm in diameter, and	Sinny type:	
	<ul> <li>Changes in</li> </ul>		lesion in a native infrapopliteal artery	Ctudu tuno.	
	rate	metal stent	<ul> <li>Presence of a single primary target</li> </ul>	artery	
	<ul> <li>Secondary patency</li> </ul>	coated bare-	lesions to improve runoff status	Intrapopilteal	
	patency rate	Placebo-	intervention of TASC A femoropopliteal		
	<ul> <li>6-mo primary</li> </ul>	<u>comparator</u> :	Rutherford-Becker classs 2 if successful	therapy of local	
	2° endpoints:		<ul> <li>lifestyle-limiting claudication</li> </ul>	Interventional	
		eluting stent	<u>с</u> л	rates atter	
	rate	sirolimus-	<ul> <li>PAD with Rutherford-Becker class 3–</li> </ul>	primary patency	21622669
	<ul> <li>1-y primary patency</li> </ul>	Polymer-tree	• Age ≥21 y	It SES Improves	2011 (334)
SES improved mid-term patency rates compared to BMS	<u>1° endpoint:</u>	Intervention:	Inclusion criteria:	Aim: Determine	Rastan A, et al.
				Size: n=140 pts	
	:			RCT	
	BMS (p=0.001)			Study type:	
	<ul> <li>DES was superior to</li> </ul>	BMS	outflow to the foot		
	mo	Comparator:	Exclusion criteria: Lack of ≥1 vessel	vs. BMS for CLI	22169682
	the target lesion at 12			PTAS with DES	2012 (333)
compared with bare metal stents	<ul> <li>Binary restenosis of</li> </ul>	DES	stenosis	infrapopliteal	Bosiers M, et al.
Reduced restenosis and the need for reintervention	1° endpoint:	Intervention:	Inclusion criteria: CLI and infrapop	Aim: Assess	DESTINY
				Size: n=476 pts	
	endnoints				
	DEB noninferior with		Detailed in NEJM	BCT	
	(p<0.02)	Standard PTA	• Lesion length ≥15 cm	<b>2</b>	26106946
	<ul> <li>DEB superior</li> </ul>	Comparator:	Exclusion criteria:	bailout stenting	2015 (332)
	target lesion at 12 mo	•		vs. PTA with	<u>a</u>
	<ul> <li>Primary patency of</li> </ul>	DEB	symptoms	efficiacy of DEB	Rosenfield K, et
N/A	1° endpoint:	Intervention:	Inclusion criteria: Fontaine 2-4	Aim: Assess	LEVANT-2
				PTA=25 pts	
				pts; Standard	
				Size DER=95	
	group (p<0.01)		residual stences >50%		
	lower in the DEB	Standard PTA	provisional stenting after angioplasty	Study type:	
	<ul> <li>Late lumen loss was</li> </ul>	Comparator:	Exclusion criteria: Pts requiring		23046320
	loss at 6 mo			vs. PTÁ	2012 (331)
	Angiography lumen	DEB	symptoms	efficacy of DEB	Fanelli F, et al.
Small study	1º endnoint:	Intervention:	Inclusion criteria: Fontaine 2h-4	Aim: Assess	DFREITUM

				Tepe G, et al. 2015 (336) <u>25616822</u>		Siablis D, et al. 2014 (335) <u>25234679</u>		
	<u>Size</u> : n=154 pts	Study type: multicenter RCT	restenosis rate after peripheral arterial interventions.	<u>Aim</u> : Evaluate 5-y follow-up of PCB on the	<u>Study type:</u> Prospective PCT <u>Size</u> : n=50 pts	<u>Aim</u> : To compare PCB vs. DES in long infrapopliteal lesions		randomized, multi-centre, double-blind trial <u>Size</u> : n=161 pts
			Exclusion criteria: N/A	<ul> <li>Inclusion criteria:</li> <li>Included in the THUNDER study</li> </ul>	Exclusion criteria: N/A	<ul> <li>Inclusion criteria:</li> <li>Rutherford classes 3–6</li> <li>Angiographically documented infrapopliteal disease ≥70 mm</li> </ul>	<ul> <li>Known systemic coagulopathy</li> <li>Buerger's disease</li> <li>ALI</li> <li>Life expentency &lt;1 y</li> <li>Intolerance of aspirin, clopidogrel, and heparin</li> </ul>	<ul> <li>Diameter stenosis of ≥70%</li> <li>Exclusion criteria:</li> <li>Pregnant pts</li> <li>Visible thrombus within target lesion</li> </ul>
	and paclicaxel added to standard nionic contrast medium (paclitaxel-in- CM Group)	<ul> <li>Plain old</li> <li>balloon</li> <li>angioplasty</li> </ul>	nonionic contrast medium (PCB aroun)	<ul> <li>Intervention:</li> <li>PCB and standard</li> </ul>	<u>Comparator</u> : Placebo- coated bare- metal stent	Intervention: Polymer-free sirolimus- eluting stent		
	<ul> <li><u>2° endpoints:</u></li> <li>freedom from TL revascularization, binary restenosis rate, and amputation</li> </ul>	evaluated by quantitative angiography)	the postprocedural and 6-mo follow up minimal lumen diameter	<u>1° endpoint:</u> ● Angiographic LLL (difference between	<ul> <li><u>2° endpoints:</u></li> <li>Immediate post- procedure stenosis</li> <li>Target lesion revascularization</li> </ul>	<ul> <li><u>1° endpoint:</u></li> <li>Target lesion restenosis &gt;50% at 6 mo</li> </ul>		
160				<ul> <li>5-y follow up period resulted in maintained reduced TL revascularizationrate following PCB treatment. No signs of drug-related local vessel abnormalities were detected.</li> </ul>		<ul> <li>Significant lower residual immediate post-procedure stenosis in DES compared with PCB in long infrapopliteal lesion</li> <li>At 6 mo, significantly reduced vessel restenosis in DES compared with PCB</li> </ul>		

confidence interval; CLI, critical limb ischemia; DCB, drug coated balloon; DEB, drug eluting balloon; DES, drug eluting stent; DM, diabetes mellitus; HR, hazz	ABI indicates ankle-brachial index; AFS, amputation-free surivival; ALI, acute limb ischemia; BMS indicates bare metal stent; CD-TLR, clinically driven target lesion revascularization; CI,	(Control group)	nonionic CM	and standard	angioplastic	balloon	Plain old Plain old	Comparator:
; DM, diabetes mellitus; HR, hazard ratio; IA-DEB, apmhirion-	ally driven target lesion revascularization; CI,							

drug eluting balloon; IC, intermittent claudication; ISR, in stent restenosis; IQR, interquartile range; JACC, Journal of American College of Cardiology; LLL, late lumen loss; N/A, not applicable; OR, odds ratio; OS, overall survival; PAD, periphery artery disease; PCB, paclitaxel-coated blaoon; PEB, paclitaxel eluting balloon; PTA, percutaneous angioplasty, PTAS, percutaneous angioplasty stent; pt, patient; RCT, randomized controlled trial; RR, relative risk; SES, self-expanding stents; and SFA, superficial femoral artery; and TL, target lesion.

## 8.2.1. Evidence Table 40. Nonrandomized Trials, Observational Studies, and/or Registries of Endovascular Revascularization for Chromic CLI–Section
<u>16730466</u>	Faglia E, et al. 2006 (339)	Ferraresi R, et al. 2009 (337) <u>19112033</u> 2013 (338) <u>23975668</u>	
<u>Size</u> : n=564 total pts: 420 PTA, 117 bypass, 27 both	<u>Study type:</u> Case series	<u>Study type</u> : Case series: infrapop PTA for CLI <u>Size</u> : n=101 pts <u>Study type:</u> Case series <u>Size:</u> n=64 pts	
Exclusion criteria: • Pts without DM • No stenosis >50%	Inclusion criteria: Pts with DM with CLI	Inclusion criteria:       Pts with DM         with CLI due to infrapop PAD         Exclusion criteria:       Above the         knee >70% stenosis         Inclusion criteria:       CLI due to         CTO in below the knee artery         Exclusion criteria:         CTO in below the knee artery         Exclusion criteria:         Pts with         concomitant above-knee arterial         steno-occlusive lesions including         the aortoiliac and femoropopliteal         arterial lesions, clinical or         imaging signs of embolic         disease, or who had undergone         thrombolysis prior to         endovascular or surgical         procedures.	
<b>Results:</b> Major amputation was associated with absence of revascularization (OR: 35.9; p<0.001; 95% CI: 12.9–99.7), occlusion of each of the 3 crural arteries (OR: 8.20; p=0.022; 95% CI: 1.35–49.6), wound infection (OR: 2.1; p=0.004; 95% CI: 1.3– 3.6), dialysis (OR: 4.7; p=0.001; 95% CI: 1.9–11.7) increase in TcPO ₂ after revascularization (OR: 0.80; p<0.001; 95% CI:	<u>1° endpoint:</u> Limb salvage	1° endpoint: Limb salvage         Results: 93% limb salvage rate;         no comparator         1° endpoint: Limb salvage         and 59.1% primary patency rate         at 1 y. No comparator group.	
<ul> <li>23 above-the-ankle amputations (4.1%) were performed at 30 d: 6 in PTA pts, 3 in BPG pts, 14 in nonrevascularized pts. In the follow-up of 558 pts (98.9%), 62 repeated PTAs and 9 new BPGs, 32 new major amputations (16 in PTA pts, 14 in BPG pts and 2 in nonrevascularized pts) were performed. Major amputation was associated with absence of revascularization (OR: 35.9; p&lt;0.001; 95% CI: 1.29–99.7), occlusion of each of the 3 crural arteries (OR: 8.20; p=0.022; 95% CI: 1.35–49.6), wound infection (OR: 2.1; p=0.004; 95% CI: 1.3–3.6), dialysis (OR: 4.7; p=0.001; 95% CI: 1.9–11.7) increase in TcPO₂ after revascularization (OR: 0.80; p&lt;0.001; 95% CI: 0.74–0.87). 173 pts died during follow-up and this</li> </ul>	<ul> <li>PTA was carried out in 420 (74.5%), BPG in 117 (20.7%)</li> <li>pts. In 27 (4.8%) pts both PTA and BPG were not possible.</li> </ul>	<ul> <li>of Output Dynamic Pyrock consortience of Concept Pyrol (p&lt;0.001). CLI at presentation (tissue loss, HR: 8.1; p&lt;0.001), poor outflow (HR: 2; p=0.023), and renal failure (HR: 2.5; p=0.02) were associated with decreased survival.</li> <li>Proof of concept; poor quality</li> <li>The limb salvage rate was 93% after a mean follow-up of 1048±525 d (2.9±1.4 y). Transcutaneous oxygen tension significantly increased after 1 mo (18.1±11.2 vs. 39.6±15.1; p&lt;0.05). After 1 y, target-vessel re-stenosis had occurred in 42% of the non-amputated limbs, 9 pts (9%) had died because of medical conditions unrelated to PTA and 3 pts had undergone repeat PTA for recurrent CLI.</li> <li>Reasonable limb salvage</li> <li>Poor vessel patency at 1 y</li> <li>The BTK EVT was performed on 64 limbs. Technical success rate was 93.8% and limb salvage rate was 90.6%. 3 of 4 limbs with technical failure and 3 of 60 limbs with technical failure and 3 of 60 limbs with technical failure and 12 mo follow-up, respectively. Minor complications disappeared through the follow-up periods and there was no 30 d complication or systemic adverse events for the treated vessel.</li> </ul>	of distal hypass were associated with decreased natency

	Feiring AJ, et al. 2010 (342) <u>20378075</u>	lida O, et al. 2012 (341) <u>22051875</u>	Faglia E, et al. 2005. (340) <u>15878541</u>	
	<u>Study type</u> : Case series <u>Size</u> : n=105 pts	Study type: Retrospective analysis of BTK PTA: angiosome vs. non- angiosome <u>Size</u> : n=369 limbs from 329 consecutive pts	<u>Study type:</u> Case series <u>Size:</u> n=993 pts	
	Inclusion criteria: Infrapop DES for CLI Exclusion criteria: • Lack of CLI • No exclusions for other comorbidities	Inclusion <u>criteria</u> : CLI treated with endo <u>Exclusion criteria</u> : Unsuccessful recanalization of ≥1 vessel to the pedal arch	Inclusion criteria: CLI treated with endo Exclusion criteria: • Pts without DM • No stenosis >50%	
	<u>1° endpoint</u> : Major amputation and mortality <u>Results</u> : The 3 y cumulative incidence of amputation was 6±2%, survival was 71±5%, and amputation-free-survival was 68±5%	<u>1° endpoint</u> : Limb salvage <u>Results</u> : Freedom from major amputation at 18±16 mo was higher in the angiosome directed group 51%±8% vs. 28%±8%, p=0.008	<u>1° endpoint</u> : Limb salvage <u>Results</u> : 1.7% major amputation rate at variable follow-up of 26±15 mo. No comparator	0.74–0.87).
17.2	<ul> <li>Infrapop DES for CLI appears effective</li> <li>The mean pt age was 74±9 y. There were 228 DES implanted (83% Cypher [Cordis, Johnson &amp; Johnson, Warren, New Jersey], 17% Taxus [Boston Scientific, Maple Grove, Minnesota]). The number of stents per limb was 1.9±0.9, and 35% of limbs received overlapping DES (length of 60±13 mm). There were no procedural deaths, and 96% of pts were discharged within 24 h. The 3 y cumulative incidence of amputation was 68±5%. Only 12% of pts who died had a preceding major amputation. Rutherford category, age,</li> </ul>	<ul> <li>AFS higher in angiosome directed endo group</li> <li>During follow-up (mean, 18±16 mo), the overall limb salvage rate was 81% (300 of 369), death occurred in 36% (119 of 329), and the reintervention rate was 31% (114 of 369). After propensity score adjustment, the estimated (± standard error) rates for AFS (49%±8% vs. 29%±6%; p=0.0002), freedom from MALE (51%±8% vs. 28%±6%, p=0.001) were significantly higher in the direct group than in the indirect group for up to 4 y after the index procedure. After multivariable Cox proportional analysis, the independent factors associated with major amputation were hemoglobin A(1c) level (HR: 1.4; 95% CI: 1.1–1.9; p=0.006) and cilostazol administration (HR: 0.28; 95% CI: 0.11–0.70; p=0.006) in the direct group, and C-reactive protein level (HR: 1.2; 95% CI: 1.1–1.4; p=0.002) in the indirect group</li> </ul>	<ul> <li>PTA effective</li> <li>PTA was successful performed in 993 pts. 17 (1.7%) major amputations were carried out. 1 death and 33 nonfatal complications were observed. Mean follow-up was 26±15 mo. Clinical restenosis was observed in 87 pts. The 5 y primary patency was 88%, 95% CI 86-91%. During follow-up 119 (12.0%) pts died at a rate of 6.7% per y.</li> </ul>	was associated with age (HR: 1.05; p<0.001; 95% CI: 1.03– 1.07), Hx of cardiac disease (HR: 2.16; p<0.001; 95% CI: 1.53–3.06), dialysis (HR: 3.52; p<0.001; 95% CI: 2.08–5.97), absence of revascularization (HR: 1.68; p<0.001; 95% CI: 1.29–2.19) and impaired ejection fraction (HR: 1.08; p<0.001; 95% CI: 1.05–1.09).

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NIA	1° endpoint: Ischemic ulcer healing and limb salvage rates	Inclusion criteria: Infrapop intervention for CLI in pts with	<u>Study type:</u> Retrospective case	Acin F, et al. 2014 (345)
was associated with a higher rate of ISS. During clinical follow-up of 144 (91%) pts over a mean $31.1\pm20.3$ mo, there were 27 (18.8%) deaths, 4 (2.8%) amputations, and no bypass surgery. Clinical status improved in 92% of the pts with CLI and 77% of the pts suffering from claudication (p=0.022).	right; no comparator group			
83.8% at 60 mo. In-stent stenosis was predominantly observed in the first v after stent placement. Female gender	Results: Results in column to the	Exclusion criteria: Lack of infrapop stenosis	Size: n=158 pts	
• Technical success was achieved in all cases. The primary patency rates were 97.0% after 6 mo, 87.0% after 12 mo, and	restenosis; freedom from death, amputation, and bypass	for CLI	series	2012 (344) 22313195
<ul> <li>Proof of concept for infrapop DES</li> </ul>	1° endpoint: Angiographic binary	Inclusion criteria: Infrapop DES	Study type: Case	Werner M, et al.
<ul> <li>Infrapop DES for CLI appears effective</li> <li>In total, 103 pts were included in the analysis; 41 (75.6% with DM) were treated with a BMS (47 limbs; 77 lesions) and 62 (87.1% with DM) with an SES (75 limbs; 153 lesions). At 3 y, SES-treated lesions were associated with significantly better primary patency (HR: 4.81; 95% CI: 2.91–7.94; p&lt;0.001), reduced binary restenosis (HR: 0.38; 95% CI: 0.25–0.58; p&lt;0.001), and better repeat intervention-free survival (HR: 2.56; 95% CI: 1.30–5.00; p=0.006) vs. BMS-treated ones. No significant differences were identified between SESs and BMSs with regard to overall 3 y pt mortality (29.3% vs. 32.0%; p=0.205) and limb salvage (80.3% vs. 82.0%; p=0.507).</li> </ul>	<ol> <li><u>endpoint</u>: Primary clinical and angiographic endpoints included mortality, limb salvage, primary patency, binary angiographic reset intervention-free survival.</li> <li><u>Results</u>: At 3 y, SES-treated lesions were associated with significantly better primary patency (HR: 4.81; 95% CI: 2.91– 7.94; p&lt;0.001), reduced binary restenosis (HR: 0.38; 95% CI: 0.25–0.58; p&lt;0.001), and better repeat intervention-free survival (HR: 2.56; 95% CI: 1.30–5.00; p=0.006) vs. BMS-treated ones. No significant differences were identified between SESs and BMSs with regard to overall 3 y pt mortality (29.3% vs. 32.0%; p=0.205) and limb salvage (80.3% vs. 82.0%; p=0.507).</li> </ol>	Inclusion criteria: CLI treated with infrapop DES or BMS Exclusion criteria: • Hx of severe contrast allergy/hypersensitivity • Hypersensitivity to ASA and/or clopidogrel • Systemic coagulopathy or hypercoagulation disorders • ALI • Buerger disease • Deep vein thrombosis • Bifurcation and/or trifurcation lesions • Previous use of other DES (not SES) • Stenting indications after suboptimal and/or complicated balloon angioplasty • Elastic recoil Flow-limiting dissection • Residual stenosis >30%	<u>Study type:</u> Registry: Infrapop DES vs. BMS <u>Size</u> : n=103 pts	Siablis D, et al. 2009 (343) <u>19620014</u>
creatinine level, and dialysis (p≤0.001–0.04) were predictors of death but not amputation. Target limb revascularization occurred in 15% of pts, and repeat angiography in 35% of pts revealed a binary restenosis in 12%.				

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				2017) TI VI	2014 (349)	Krat MR at al							<u>+7100007</u>	220E0Z24	Kabra A, et al. 2013 (348)						23358605	2013 (347)	Fossacaca R, et al.				188/00/6	2008 (346)	<u></u>	Alexandrescu VA, et								24527215
	relationship	andiosome	treatment with	corios accossing CII	Retroenentive nace	Study type			<u>51ze</u> : n=64 pts		reiduorisiiib	relationship	treatment with		<u>Study type</u> : Prospective case	Size: II-ZUI pts		relationship	angiosome	treatment with	series assessing CLI	Retrospective case	Study type:	<u>Size</u> : n=98 pts	relationshin		treatment with	series assessing CI I	Retrospective case	Study type:	procedures; 92 pts	Size: n=101		relationship	and angiosome	of infrapop vessels	treatment with number	series assessing CLI
					intervention for CI in the	Inclusion criteria: Infrance									Inclusion criteria: Intrapop intervention for CLI in pts						DM .	intervention for CLI in pts with	Inclusion criteria: Infrapop						intervention for CI I in pts with	Inclusion criteria: Infrapop								DM
		INO UIIIEIEIICE DELWEEII	time to complete wound		- Complete wound booling and	10 ondpoint:	statistically significant (p=0.06)	and IR group (75%) was not	salvage in the DR group (84%)	significant (p=0.021). The limb	IR groups was statistically	ulcer healing between the DR and	The difference in the other of		<ul> <li>Ischemic ulcer healing and limb</li> </ul>	revasc	revasc vs. anglosome directed		Results: No difference in		1.6 and 12 mo	healing and limb salvage rates at	1° endpoint: Ischemic ulcer		described for endo for CLI	Nesules. Lillo salving and	Docuto: Limb colving and		healing and limb salvage rates	1° endpoint: Ischemic ulcer			wound	vessel was in angiosome of	vessels; no difference is single	vessel run-off and multiple	Results: No difference between 1	
100															Small study							difference	Higher TcPO ₂ in angiosome group but no clinical outcome							No comparator group								

			26391460	איז, פנ al. 2015 (353)				23822940	Usawa S, et al. 2013 (352)	)				<u>19179041</u>	Neville RF, et al. 2009 (351)						<u>24333196</u>	Lejay A, et al. 2014 (350)	
(1,527)	Size: n=13 studies	Study type: SR/MA of	ischemia	natural hx of untreated	for endo therapy)	<b>Size:</b> n=111 nte (n=57	with anglosome relationship	series assessing CLI	Study type: Retrospective case	Size: n=48 pts		relationship	treatment with	series assessing CLI	<u>Study type</u> : Retrospective case		Size: n=54 pts	<b>1</b>	relationship	angiosome	series assessing CLI	Study type: Retrospective case	Size: n=97 pts
<ul> <li>trainscutarieous O₂ pressure</li> <li>40 mmHg for ≥1 y.</li> <li>No revasc treatment.</li> </ul>	<ul> <li>Flat pulse volume recording</li> <li>Francourbaneous On processing</li> </ul>	<ul> <li>Rutherford class 4–6</li> <li>Or ankle pressure &lt;70 mm Hg,</li> </ul>	gangrene	Studies with pts. reporting rest     pain tissue loss ulter or					Inclusion criteria: CLI	-					Inclusion criteria: Infrapop bypass for CLI in pts							Inclusion criteria: Infrapop bypass for CLI in pts	
• worseried would of ulcet. 32 % (95% CI: 10%–62%)	<ul> <li>Major amputation rate: 22%</li> <li>(95% Cl: 2%–42%)</li> <li>Worsened wound or ulcer: 35%</li> </ul>	<ul> <li>All-cause mortality: 22% (95%</li> <li>Cl: 12%-33%)</li> </ul>	Results:	amputation, wound healing		for angiosome directed group	<ul> <li>Wound healing rate was faster</li> </ul>	who had angiosome or indirect	<ul> <li><u>1° endpoint</u>:</li> <li>Time to complete wound in pts</li> </ul>	between the 2 groups	no difference in time to healing	complete wound nealing ; among wounds that did heal there was	<ul> <li>Angiosome group had more</li> </ul>	time to complete wound	<ul> <li><u>1° endpoint:</u></li> <li>Complete wound healing and</li> </ul>	revasc	higher limb salvage at 1, 3, and 5	• Angiosome directed bypass had	group	limb salvage rates between	survival, primary patency, and	<ul> <li><u>1° endpoint:</u></li> <li>Median ulcer-healing time,</li> </ul>	
470				to improved medical care	Travel for rough "more reasonable the summer and such the block of the second s				Small study	-					Small study							Small study	

AAA indicates abdominal aortic aneurysm; ABF, aortobifemoral bypass; ABI, ankle-brachial index; AFS, amputation free survival; AIOD, aortoiliac occlusive disease; ALI, acute limb ischemia; ASA, aspirin; BMS, bare metal stent; BPG, bypass graft; BTK, below the knee; BPG, bypass graft; CI, confidence interval; CLI, critical limb ischemia; CTO, chronic total occlusion; DES, drug eluting stent; DM, diabetes mellitus; DR, direct revascularization; EVT, endovascular treatment; HR, hazard ratio; IR, indirect revascularization; MALE major adverse

Exclusion criteria: Revascularization treated arms

limb event; N/A, not applicable; OR, odds ratio; PTA, percutaneous angioplasty; pt, patient; R/PTAS, recanalization, percutaneous transluminal angioplasty, and stenting; RR, relative risk;

SES, self-expanding stents; and TcPO₂, transcutaneous oxygen pressure.

	<ul> <li>(78%) of 37 pts and 26 (77%) of 34 pts in groups I and II, respectively (p=0.6); however, 43% and 35% of the pts, respectively, still had symptoms.</li> <li>QoL improved significantly after</li> </ul>	biacement, Group i	excluded.		
	<ul> <li>II), respectively.</li> <li>Clinical success rates at 2 y were 29</li> </ul>	Comparator: Direct stent	pregnant or in whom	Size: n=279 pts	
	126 limbs and 10 (7%) of 136 limbs (group	group II	Women who were	Study type: RCT	
	rates were 119 (81%) of 149 limbs and 6 (4%) of 143 limbs (group I) vs_103 (82%) of	gradient >10 mm Hg	Exclusion criteria:	angioplasty	
	hemodynamic success and complication	residual mean pressure	angiography	placement after	<u>9643685</u>
	follow-up was 9.3 mo (range 3-24). Initial	placement in case of a	than 50%, proven by	delayed stent	1998 (221)
	done in 59 (43%) of the 136 pts. The mean	subsequent stent	artery stenosis of more	direct stent vs.	Tetteroo E, et al.
N/A	1º endpoint:	Intervention: Primary	on the basis of iliar-	<u>Aim</u> : To determine	Dutch Iliac Stent
	thrombolysis group.				
	amputations), as compared with 315 in the				
	551 open operative procedures (excluding				
	<ul> <li>At 6 mo the surgery group had undergone</li> </ul>				
	p=0.23.				
	• 1 y differences 95% CI: -12.9%-3.1%;		possibility.		
	p=0.43.		pregnancy was a		
	• 6 mo differences 95% CI: 10.5%–4.5%;		pregnant or in whom		
	group;		Women who were		
	rates of 74.8% and 69.9% in the surgery		Exclusion criteria:	Size: n=544 pts	
in the surgery group.	65.0% at 1 y, as compared with respective		l	2	
episodes of intracranial hemorrhage	urokinase group were 71.8% at 6 mo and		threatening ischemia	Multicenter	
was fatal. By contrast, there were no	<ul> <li>Amputation-free survival rates in the</li> </ul>		for reversible limb-	Study type : RCT	
urokinase group (1.6%), 1 of which	improvements in mean ABI.		that met the guidelines		
intracranial hemorrhage in the	<ul> <li>Both Tx groups had similar significant</li> </ul>	revascularization	d before randomization	intervention	
(p=0.005). There were 4 episodes of	dissolution of thrombus in 167 (67.9%).	Comparator: Surgical	bypass graft) within 14	prior endovascular	
surgery group (5.5%)	recanalization in 196 (79.7%) and complete		leg (native artery or	with and without	9545358
as compared with 14 pts in the	for 246 pts treated with urokinase, revealed	urokinase	embolic occlusion of a	revascularization	1998 (356)
pts in the urokinase group (12.5%)	<ul> <li>Final angiograms, which were available</li> </ul>	Thrombolysis with	Acute thrombotic or	surgical	Ouriel K, et al.
Major hemorrhane occurred in 32	1º and noint:	Intervention:	Inclusion criteria	Aim I F lysis vs	TOPAS
				Size: n=237 pts	
				Study type : RCT	
iysis and surgical groups.				Intervention	
مستمسط المدلسينية لالسب والمنا					

were lost to tonow-up after randomization (3 assigned angioplasty, 2 surgery); of these, 3 were lost (1 angioplasty, 2 surgery) during the first y of follow-up. 195 (86%) of 228 pts assigned to bypass surgery and 216 (96%) of 224 to				<u>Size</u> : n= 452 pts	
above the ankle or death). 7 pts		(N=228)	Exclusion criteria: N/A	Study type: RCT	
endpoint (amputation of trial leg	<u>Salety endpoint</u> : Mortainty	Comparator: Bypass			16325694
<ul> <li>The trial ran for 5.5 y, and follow-</li> </ul>		(N=224)	due to infrainguinal	Surgical bypass vs.	Adam DJ, et al.
<ul> <li>Equal outcomes</li> </ul>	1° endpoint: Amputation free survival	Intervention: PTA	Inclusion criteria: CLI	Aim: Infrainguinal	BASIL
	angioplasty group.				
	were able to walk significantly farther on a treadmill at 6 and 12 mo than those in the		Exclusion criteria: N/A	<u>Size</u> : n=104 pts	
	respectively (p=0.01). Pts in the stent group				
residuai stenosis, etc.	at 12 mo the rates on duplex	<b>Comparator:</b> Angloplasty	occlusion of the SFA	multicenter	(800) 0007
quantitative data on lumen diameter,	and 43% in the angioplasty group (p=0.05);		chronic limb ischemia		al.
done in all pts, resulting in lack of	on angiography was 24% in the stent group	expanding nitinol stent	Severe claudication or	vs. Angioplasty	Schillinger M, et
Angiographic follow-up was not		Intervention: Self-	Inclusion criteria:	Aim: Primary Stent	CRISP-US
	the groups. The frequency of major adverse events was similar at 1 y (4.9% vs. 5.9%).				
	improvement was also comparable between				
	respectively. Functional and hemodynamic			Size: n=203 pts	
	with the SMART stent and Wallstent,				
	(2.0% vs. 4.0%) III III 2 gloups. Filliary			multicenter	
	(2.0% vs. 0.0%), and revascularization			Chindry tripper DOT	
	5		Exclusion criteria: N/A	suboptimal PTA.	<u>15361558</u>
group (98.2% vs. 87.5%; p=0.002).		Comparator: Wall stent		Wallstent after	2004 (357)
was higher in the SMART stent	rate was equivalent for the SMART stent		Chronic limb ischemia	SMART stent vs.	Ponec D, et el.
The acute procedural success rate	1° endpoint: 9 mo composite end point	Intervention: Smart Stent	Inclusion criteria:	Aim: Compare	CRISP-US
	9%).				
	7% vs 4% respectively (95% CI -2% to				
	were similar at 71% vs. 70% (p=0.2),				
	follow-up. 2 y cumulative patency rates				
	difference between the groups during				
	intervention (p<0.05) but we found no				

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	procedure)	(N=1,050)		treatment efficacy,	
	MALE-POD (i.e., death within 30 d of	Comparator: Dissoo	Exclusion criteria: N/A	therapy in pts with	<u>25241324</u>
	Safety endpoint:	(n=1,050)	1 - -	vs. best surgical	2014 (360)
		Endovascular Tx	with CLI (R4-6)	best endovascular	Farber A, et al.
N/A	1° endpoint: MALE-free survival	Intervention:	Inclusion criteria: Pts	Aim: To compare	BEST-CLI
				<b>Size:</b> n=1,404 pts	
			bypass grafts	RCT	
			revisions of infrainguinal	blinded, phase III	
	survival, graft patency, and limb salvage		hypercoagulable state,	randomized, double	
Limb salvage: 88%	stenosis, amputation, index graft failure		Exclusion criteria: IC,	Prospective,	
Secondary patency: 80%	freedom from significant index graft			Study type:	
<ul> <li>Primary assisted patency: 77%</li> </ul>	Safety endpoint: All-cause graft failure,	(N=341)	placebo or E2F decoy	2	
<ul> <li>Primary patency at 1 y: 61%</li> </ul>		Comparator: Bypass	gratt randomized to	using EZF decoy	10010230
<ul> <li>5.2% early graft occlusion</li> </ul>	amputation	•	had autologous vein	bypass for CLI	2006 (359)
• 4.7% MI	failure resulting in revision or major	(/1/c=n)	with CLI (R4-6) who	stenosis in Surgical	Conte MS, et al.
• 2.7% 30 d mortality	1° endpoint: Nontechnical index graft	Intervention: PIA	Inclusion criteria: Pts	Aim: Reduce	PREVENIII
angioplasty-first strategy.			- - - -	-	
higher than those with an					
surgery-first strategy were about 1/3					
hospital costs associated with a					
strategies, but for the first y the					
quality of life between the 2					
1.07). No difference in health-related					
adjusted HR: 0.73; 95% CI: 0.49-					
HR: 1.07; 95% CI: 0.72–1.6;					
survival (48 vs. 60 pts; unadjusted					
significantly in amputation-free					
2 strategies did not differ					
without amputation. After 6 mo, the					
amputation, and 130 (29%) dead					
amputation, 36 (8%) dead after					
trial leg), 38 (8%) alive with					
were alive without amputation (of					
end of follow-up, 248 (55%) pts					
randomization, respectively. At the					
(3–16) and 6 (2–20) d after					
intervention at a median (IQR) of 6					
attempt at their allocated					
balloon angioplasty underwent an					

	Veves A, et al. 2002 (361) <u>12093340</u>	
	<u>Aim</u> : To compare a collagen and oxidized cellulose dressing to moistened gauze with regards to wound healing. <u>Study Type</u> : RCT <u>Size</u> : n=276 pts	outcomes, and cost in pts with CLI undergoing best open surgical or best endovascular revascularization Study type: A prospective, multicenter, RCT. CLI trial has a 2- cohort design. The first cohort (1,620 pts) evaluates outcomes in pts who have adequate single segment great saphenous vein. The second cohort (480 pts) will study pts who do not have adequate single segment great saphenous vein.
	Inclusion criteria: ≥8 y of age with a diabetic foot ulcer ≥30 d duration, Wagner grade 1–2, and an area of ≥1 cm² (greatest length × greatest width). Pts had adequate circulation with an oscillometer reading of the limb that had the target wound of ≥1 U and a wound that was debrided of	
	Intervention: Promogran, a wound dressing consisting of collagen and oxidized regenerated cellulose for diabetic plantar ulcers. <u>Comparator</u> : Moistened Gauze with secondary dressing.	
	<ul> <li><u>1° endpoint:</u></li> <li>Complete healing of the study ulcer (wound)</li> <li>After 12 wk of treatment, 51 (37.0%) Promogran treated pts had complete wound closure compared with 39 (28.3%) control pts, but this difference was not statistically significant (p=0.12).</li> <li>The difference in healing between Tx groups achieved borderline significance in the subgroup of pts with wounds of &lt;6 mo duration. In pts with ulcers &lt;6 mo duration,</li> </ul>	<ul> <li>Freedom from MI</li> <li>Freedom from stroke, freedom from reinterventions (major and minor) in index leg, number of reinterventions (major and freedom from clinical failure</li> <li>Freedom from CLI</li> <li>Freedom from all-cause mortality</li> <li>Freedom from hemodynamic failure.</li> </ul>
161	Limitations: Study did not standardize frequency of dressing changes.	

Biancari F and Juvonen T 2014 (60) <u>24491282</u>	Study Acronym (if applicable) Author Year	ABI indicates ankle- ratio; IC, intermitten applicable; PTA, pei <b>Evidence Table 42</b> .																						
<u>Aim</u> : Compare direct vs. indirect revascularization for wound healing and limb salvage.	Study Type/Design; Study Size	ABI indicates ankle-brachial index; CI, confider ratio; IC, intermittent claudication; IQR, interqu applicable; PTA, percutaneous angioplasty; pt, <b>Evidence Table 42. Nonrandomized Trials, (</b>																						
Inclusion criteria: Prospective and retrospective observational studies with surgical, endovascular, or hybrid revascularization.	Patient Population	nce interval; CLI, critical limb is artile range; LE, lower extremi patient; QoL, quality of life; R Observational Studies, and/o	foot.	<ul> <li>Multiple diabetic</li> </ul>	loading device	shoe gear or an off-	fitted with appropriate	ambulatory pt to be	<ul> <li>Onwiningness or</li> </ul>	components	of the dressing	hypersensitivity to any	wound healing, known	have interfered with	a condition that may	<ul> <li>A concurrent illness or</li> </ul>	had exposed bone	<ul> <li>A target wound that</li> </ul>	infection	<ul> <li>Clinical signs of</li> </ul>	Exclusion criteria:		tissue at enrollment.	necrotic/nonviable
Intervention: Indirect Revascularization <u>Comparator</u> : Direct Revascularization <u>1° endpoint</u> : The risk of unhealed wound was significantly lower after direct revascularization (HR:	Primary Endpoint and Results (include P value; OR or RR; and 95% Cl)	a di															groups.	seen in the satety measurements between	[19%]; p=0.83) groups. No differences were	pts healed in the Promogran (8/43	wounds <6 mo duration, similar numbers of	controls (p=0.056). In the group with	healed compared with 29 (33%) of 89	43 (45%) of 95 Promogran-treated pts
<ul> <li>Pooled limb salvage rates after direct and indirect revascularization were at 1 y 86.2% vs. 77.8% and at 2 y 84.9% vs. 70.1%, respectively.</li> <li>The analysis of 3 studies reporting only on pts with DM confirmed the benefit of direct revascularization in terms of limb salvage (HR:</li> </ul>	Summary/Conclusion Comment(s)	FP, femoral popliteal; HgbA1c, hemoglobin A1c; HR, hazard ), major adverse limb event perioperative death; N/A, not femoral artery; TBI, toe-brachial index; and tx, treatment. or Chronic CLI–Section 8.2.																sbetween	ences were	5	numbers of	with	of 89	ed pts

 $\ensuremath{\textcircled{O}}$  American Heart Association, Inc. and American College of Cardiology Foundation

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<ul> <li>Limitation: Study did not stratify pts with underlying renal disease. Wound care techniques were not completely standardized.</li> <li>N/A</li> </ul>				
<ul> <li>Limitation: Study did not stratify pts with underlying renal disease. Wound care techniques were not completely standardized.</li> <li>N/A</li> </ul>			endovascular	21802888
<ul> <li>Limitation: Study did not stratify pts with underlying renal disease. Wound care techniques were not completely standardized.</li> </ul>	Intervention: LE bypass post endovascular intervention.	Inclusion criteria: CLI (rest pain or tissue loss)	Aim: LE bypass with and without prior	Nolan BW, et al. 2011 (364)
(p=u.uzo4).			Study type: Restrospective Size: n=154 pts	
7%,	<ul> <li>88.8%, respectively (p=0.8940)</li> <li>Amputation-free survival at 48 mo was 67.2%, 69.7%, and 45.9%, respectively (p=0.3883)</li> </ul>		atter direct anglosome revascularization in pts with CLI.	
	<ul> <li>respectively (p=0.5168)</li> <li>Secondary patency rates were 86.0%, 84.7%, and</li> </ul>		healing and time to healing of tissue loss	
<u> </u>	<ul> <li>The primary patency rates at 1 y in the CPA, IPA, and NPA groups were 58.4%, 54.6%, and 63.8%,</li> </ul>		grafts and its direct impact on the rate of	
<ul> <li>Similarly, in the IPA group, 90% with DAR</li> </ul>	1° endpoint:		rates of distal bypass	
with DAR healed compared with 92% in the non-	infrapopliteal bypass.	Exclusion criteria: N/A	the amputation-free	<u>23523278</u>
<ul> <li>Tissue loss was present in 141 of the 167 bypasses. In the CPA group, 83% of tissue loss</li> </ul>	Intervention: Pts with a CPA, IPA, and NPA, all underwent	Inclusion criteria: CLI Rutherford Class 4–6	Aim: The effect of pedal arch quality on	Rashid H, et al. 2013 (363)
	•	atherosclerotic disease.	1	
	3 y 73%	for indications other than		
	Reversed vein cumulative patency	Exclusion criteria: Pts		
יסט ונערויסאסן נימווויס איזי ניסוסאסן	3 y 85%	necrosis.		
<ul> <li>Limitation: Study only examined cumulative natency not primary natency atc</li> </ul>	1 v 85%	rest pain or tissue	722	
3 y 87%	1º endpoint:	tissue necrosis or schemic	size: n=675 gratts, 582	
<ul> <li>Infrapopliteal in situ cumulative patency</li> </ul>		defined by the presence	2	
3 y 62%	Comparator: Reversed vein graft	and/or limb salvage,	observational study	3795391
<ul> <li>Reversed vein patency at 5 y 63%</li> <li>Infrancoliteal reversed vein cumulative natency</li> </ul>	Intervention: In Situ Vein Graft	Inclusion criteria: Disabling claudication	Study type: Retrospective	Fogle MA, et al. 1987 (362)
<u>,</u>	significantly lower risk of major amputation (HR: 0.44; 95% CI: 0.26–0.75; r ² : 62%; 8 studies included).			
	Direct revascularization was also associated with	not reporting 6 mo data.	Size: n=1,290 Legs	
0.48; 95% Cl: 0.31–0.75; r²: 0%; 4 studies included)	0.64; 95% Cl: 0.52–0.8; r²: 0%; 4 studies included) compared with indirect revascularization.	Exclusion criteria: Data	<u>Study type</u> : 9 Study Meta-Analysis	

		24418639	Uhl C, et al. 2014 (366)							Santo VJ, et al. 2014 (365) 24613692			
Study type:	endovascular intervention	without prior	<u>Aim</u> : Pedal bypass surgery with and			vein LE bypasses	Size: n=314 autologous	Study type: Retrospective	intervention	Aim: LE bypass with and without prior endovascular		<u>Size</u> : n=1,880 LE bypasses	Study type: Retrospective cohort analysis (10 Centers)
primary therapy or after prior endovascular	6), who then required	gangrene (Rutherford 4–	Inclusion criteria: CLI with rest pain. ulcers. or				Exclusion criteria: N/A	62% and 25%, respectively.	TASC II type D or type C	Inclusion criteria: CLI LEBs were performed for CLI, 71% for tissue loss.			
<ul> <li>Overall, primary patency at 1 y was 58.3%, and secondary patency was 61.3%.</li> </ul>	1° endpoint:	Comparator: Primary pedal bypass. BSF	Intervention: Pedal Bypass post intervention. PEI	<ul> <li>The 3 y amputation-free survival was 59% for NPEI pts vs. 52% for PEI pts (p=0.399). Median follow-up time was 323 d for NPEI pts (IQR: 83–918) vs. 463 d for PEI pts (IQR: 145–946; p=0.275).</li> </ul>	<ul> <li>The 1 y primary patency rate was 62% for NPEI pts vs. 59% for PEI pts (p=0.759).</li> <li>The 3 y limb salvage rate was 89% for NPEI pts vs. 92%</li> </ul>	<ul> <li>The 5 y limb salvage rate was 89%, and the 5 y amputation-free survival was 49%.</li> </ul>	<ul> <li>Overall, Primary patency rates at 1 y and 5 y were 61% and 45%.</li> </ul>	<ul> <li><u>1° endpoint:</u></li> <li>The 30-day mortality rate was 3.5%.</li> </ul>	Comparator: Primary LE bypass NPEI	Intervention: LE bypass post endovascular intervention. PEI	1 y major amputation and 1 y graft occlusion rates were significantly higher in pts who had prior iPVI than those without (31% vs. 20%; p=0.046 and 28% vs. 18%; p=0.009), similar to pts who had a prior ipsilateral bypass (1 y major amputation, 29% vs. 20%; p=0.022; 1 y graft occlusion, 33% vs. 18%; p=0.001).	Prior PVI or bypass did not alter 30 d MAE and 1 y mortality after the index bypass.	<u>1° endpoint</u> : Major amputation and graft occlusion at 1 y postoperatively. Secondary outcomes included in- hospital MAE, 1 y mortality, and composite 1 y MALE.
			N/A							N/A			

165				
	rate was 4.5% ABI improved from 0.49 + 0.19 to 0.96 +	severe claudication or CLI.		
	<ul> <li>Technical success was 100% Perionerative mortality</li> </ul>	endovascular therapy was		
	1° endpoint:	Clinical indication for		
		or iliac arteries (EIAs).		
	technique was performed in the last 4 pts.	extending to the common	Size: n=22 pts.	
	Teconstruction with the Y-guidewire configuration	and 16 infrarenal),		
	pts and the aortic birurcation	suprarenal, 4 Juxtarenal,	Retrospective	
	In 9 cases, covered stents in kissing conliguration in 9			
	in 0 apparent traine appropria	ontolico popueion (o	Study Tump	
	configuration were deployed	cm) TASC type D		
	January 2008–September 2014. BMSs in kissing	were long-segment (>10	of AIOD	26370748
	of AIOD from	Indication for treatment	endovascular treatment	2016 (368)
N/A	A total of 22 pts underwent total endovascular treatment	Inclusion criteria:	Aim: To evaluate	Kasemi H, et al.
	adjustment			
	<ul> <li>No significant difference in AFS after propensity score</li> </ul>			
	• Surgery: 95%, 77%, 75.3%			
	/5.3%			
	• LIMB SALVAGE: (1 y, 3 y, 5 y): Endo: 87%, 77%,			
			<u>azic</u> . 11–000 pts	
	• Mortality: (30 d, 1 y, 3 y): Endo: 5.1%, 24.3%, 41.1%		<b>Cito:</b> n=0.70 ntn	
	intervention		center	
	<ul> <li>Mortality, limb salvage, AFS, Freedom from repeat</li> </ul>		Ubservational single	
	<u>1° endpoint:</u>		Study type:	
		Exclusion criteria: N/A		<u>21195637</u>
	Comparator: Bypass (N=341)		bypass for CLI	2011 (367)
		Consecutive pts enrolled	pop PTA vs. surgical	al.
N/A	Intervention: PTA (N=517)	Inclusion criteria:	Aim: Compare Fem-	Korhonen M, et
	81.3% in BSF group (p=0.765).			
	<ul> <li>Survival rate at 1 y was 79.5% in PEI group and</li> </ul>			
	17.9%; p=0.547).			
	p=0.515) or graft occlusions within 30 d (19.4% vs.			
	impact on either limb salvage (82.3% vs. 71.6% at 1 y;			
	<ul> <li>Prior endovascular intervention had no significant</li> </ul>			
	was 73.5% vs. 48.6% (p=0.100).			
	48.3% in BSF group (p=0.409) and secondary patency			
	<ul> <li>Primary patency at 1 y was 67.0% in PEI group vs.</li> </ul>		pts	
	major amputation.		bypass operations in 71	
	those cases was futile and 78.6% of pts had to undergo	Exclusion criteria: N/A	Size: n=75 pedal	
	<ul> <li>Graft occlusion within 30 d was 18.7%. Revision in</li> </ul>			
	<ul> <li>Limb salvage was 76.8% and survival was 80.4%</li> </ul>	intervention.	Retrospective	

	Limb salvage was $81\% \pm 5\%$ at 5 y.			
	grafts (27%).			
	required during follow-up to maintain patency of the			
	<ul> <li>A high revision rate for stenosis or thrombosis was</li> </ul>			
	for secondary patency SP.	reconstructions.	Size: n=154 pts	
	for primary patency, $63\% \pm 5\%$ for PAP, and $65\% \pm 5\%$	failed previous	2	
	<ul> <li>Overall, 5 y cumulative patency rates were 44% ± 5%</li> </ul>	were performed after	Retrospective	
	1.2%.	<ul> <li>48% of bypass grafts</li> </ul>	Study Type:	
	grafts (11%), resulting in early amputation (<30 d) in	severe claudication.	2	
	Perioperative graft failure (<30 d) occurred in 18 bypass	21%); the rest were for	arterial bypass gratting	
		ulcer: 33%; gangrene:		
	<u>1° endpoint:</u>	salvage (rest pain: 36%;	composite vein grafts	
		were performed for limb	autogenous	111/4//6
	vein grafts were examined	<ul> <li>90% of the operations</li> </ul>	long-term results of	2001 (370)
N/A	Intervention: Infrainguinal bypasses using composite	Inclusion criteria:	Aim: To evaluate the	Chew DK, et al.
		surgery.		
		intra-abdominal vascular		
	and 34%, respectively.	nad previously undergone		
	complications, with mortality rates of 94%, 44%, 38%,	aneurysm, and pts who		
	dialysis and cardiac complications were the most lethal	ischemia, secondary		
	<ul> <li>Multiorgan failure, mesenteric ischemia, need for</li> </ul>		pincennies	
	1.4–3.1; p<0.001).	nypertension,	autuulliac uypass	
		byportopoion		
	5.8: n=0.035) and cardiac disease (OR: 2.1: 05% CI:	renovascular	aortobifemoral and 144	
	followed by renal insufficiency (OR 2.5: 95% CI 1.1-	ischemia, secondarv	Size: n=3.623	
	the most significant risk factor for 30-day mortality.	limb		
	• Gangrene (OR: 3.3; 95% Cl: 1.7–6.5; p=0.005) was	excluded pts with acute	Retrospective	
		Exclusion criteria: We	Study Type:	
	3.0–4.1) and the 30 d major complication rate remained			
	study period, but the 30 d mortality at 3.6% (95% CI:	chronic CLI	outcomes.	
	<ul> <li>The annual caseload fell from 323 to 106 during the</li> </ul>	due to chronic IC or	open aortic repair	
	<u>1° endpoint:</u>	performed in Denmark	endovascular repair on	<u>26115920</u>
		Bypass procedures	effect of growing	2015 (369)
N/A	Intervention: Open Bypass	Inclusion criteria:	Aim: To identify the	Bredahl K, et al.
		from the study.		
		occlusion were excluded		
		aortoiliac thromboembolic		
	at 3 y	vascular disease and		
	• The primary patency rate was 95.2% at 1 y and 90.5%	occlusive		
	was 39.5 mo (range, 5–80 mo).	with inflammatory		
	$\bullet$ 0.98 + 0.04 at the left side (n<0.01) Mean follow-un	Exclusion criteria: Pts		

point (if any); tations;	P value; OR Relevant 2º Endpoint (if any); Study Limitations;	Endpoint Results (Absolute Event Rates, P value; OR	Patient Population	Aim of Study; Study Type;	Study Acronym; Author;
or CLI-Section 8.2.3.	Evidence Table 44. Nonrandomized Trials, Observational Studies, and/or Registries for Would Healing Therapies for CLI-Section 8.2.3.	es, and/or Registries fo	ials, Observational Studi	4. Nonrandomized Tr	Evidence Table 4
	ntrolled trial; and RR, relative risk.	patient; RCT, randomized cc	CI indicates confidence interval; CLI, critical limb ischemia; ITT, intent to treat; pt, patient; RCT, randomized controlled trial;	e interval; CLI, critical limb	CI indicates confidence
conclusive evidence that prostanoids provided long-term benefit."	<b>1° Safety endpoint:</b> No effect on mortality (RR: 1.07; 95% CI: 0.65– 1.75); higher risk of adverse events (RR: 2.35; 95% CI: 1.99–2.78)			<u>Size</u> : n=2,724 pts from 20 randomized trials	
or 363, 36.4%) than overall (201 of 753, 26.7%)	decreased amputation rate (RR: 0.69; 95% CI: 0.52–0.93)	Comparator: Placebo d or other pharmacologic 0. control	assignment was not masked; withdrawal of ≥10% of study population:	randomized trials	
<ul> <li>if major only or total) in 9 of the trials</li> <li>Amputation rate of placebo group notably higher in iloprost studies (147</li> </ul>	Cl: 1.22–1.96) but no class effect on amputations (24.8 vs. 26.7%; RR: 0.89; 95% Cl: 0.76–1.04). Iloprost	prostaglandin E1, C prostacyclin, iloprost, au betaprost, cisaprost) 0.	ı: Trials	pts with CLI" <u>Study type</u> : Meta-	
<ul> <li>Adverse events included headache, flushing, nausea, vomiting, diarrhea</li> <li>"Amputation" not specifically defined</li> </ul>	1° endpoint: Decrease in rest pain relief (RR: 1.32; 95% CI: 1.10–1.57) and ulcer healing (RR: 1.54; 95%	luding	<u>iteria</u> : CLI nce of rescue ctive	<u>Aim</u> : Evaluation of the "effectiveness and safety of prostanoids in	Ruffolo AJ, et al. 2010 (371) <u>20091595</u>
Relevant 2º Endpoint (if any); Study Limitations; Adverse Events	Endpoint Results (Absolute Event Rates, P value; OR or RR; & 95% Cl)	Study Intervention (# patients) / Study Comparator (# patients)	Patient Population	Aim of Study; Study Type; Study Size (N)	Study Acronym; Author; Year Published
ABI indicates ankle-brachial index; AFS, amputation free survival; AIOD, aortoiliac occlusive disease; BMS, bare metal stent; BSF, bypass surgery as first-line treatment; CI, confidence interval; CLI, critical limb ischemia; CPA, complete pedal arch; DAR, direct angiosome revascularization; DM, diabetes mellitus; EIA, external iliac artery; HR, hazard ratio; IC, internittent claudication; IPA, incomplete pedal arch; iPVI, ipsilateral peripheral endovascular intervention; IQR, interquartile range; LEB, lower extremity bypass; LE, lower extremity; MAE, major adverse event; MALE, major adverse limb event; N/A, not applicable; NPA, no pedal arch; NPEI, no prior endovascular intervention; PAP, primary assisted patency; PEI, prior endovascular intervention; PTA, percutaneous angioplasty; pt, patient; PVI, peripheral endovascular intervention; SP, secondary patency; and TASC, TransAtlantic Inter-Society Consensus.  Evidence Table 43. RCT Comparing Prostanoids for End-Stage Peripheral Artery Disease–Section 8.2.3.	ABI indicates ankle-brachial index; AFS, amputation free survival; AIOD, aortoiliac occlusive disease; BMS, bare metal stent; BSF, bypass surgery as first-line treatment; CI, con interval; CLI, critical limb ischemia; CPA, complete pedal arch; DAR, direct angiosome revascularization; DM, diabetes mellitus; EIA, external iliac artery; HR, hazard ratio; IC, in claudication; IPA, incomplete pedal arch; iPVI, ipsilateral peripheral endovascular intervention; IQR, interquartile range; LEB, lower extremity bypass; LE, lower extremity; MAE, adverse event; MALE, major adverse limb event; N/A, not applicable; NPA, no pedal arch; NPEI, no prior endovascular intervention; PAP, primary assisted patency; PEI, prior endovascular intervention; PTA, percutaneous angioplasty; pt, patient; PVI, peripheral endovascular intervention; SP, secondary patency; and TASC, TransAttantic Inter-Society Consensus.  Evidence Table 43. RCT Comparing Prostanoids for End-Stage Peripheral Artery Disease–Section 8.2.3.	c occlusive disease; BMS, ba some revascularization; DM, some revascularization; DM, interquar ir intervention; IQR, interquar dal arch; NPEI, no prior endc heral endovascular interventi	ABI indicates ankle-brachial index; AFS, amputation free survival; AIOD, aortoiliac occlusive disease; BMS, bare metal stent; BSF, by interval; CLI, critical limb ischemia; CPA, complete pedal arch; DAR, direct angiosome revascularization; DM, diabetes mellitus; EIA, or claudication; IPA, incomplete pedal arch; iPVI, ipsilateral peripheral endovascular intervention; IQR, interquartile range; LEB, lower e adverse event; MALE, major adverse limb event; N/A, not applicable; NPA, no pedal arch; NPEI, no prior endovascular intervention; FTA, percutaneous angioplasty; pt, patient; PVI, peripheral endovascular intervention; SP, secondary pate Consensus. Evidence Table 43. RCT Comparing Prostanoids for End-Stage Peripheral Artery Disease–Section 8.2.3.	achial index; AFS, amputat nb ischemia; CPA, complet mplete pedal arch; iPVI, ip major adverse limb event; tion; PTA, percutaneous ar <b>3. RCT Comparing Pr</b>	ABI indicates ankle-bra interval; CLI, critical lin claudication; IPA, inco adverse event; MALE, endovascular interven Consensus.
	composite vein fared ary reconstructions (SP superior patency us vein composites (SP	<ul> <li>Primary reconstructions with composite vein fared significantly better than secondary reconstructions (SP 76% vs. 54% at 5 y; p&lt;0.01).</li> <li>Arm vein composites showed superior patency compared with greater saphenous vein composites (SP 79% vs. 61% at 5 y, p&lt;0.05).</li> </ul>	● F 56 79 79		

 ${\ensuremath{\mathbb G}}$  American Heart Association, Inc. and American College of Cardiology Foundation

Inclusion criteria: Pts with neuroischemic wounds treated at a signle institutional
Inclusion criteria: "All consecutive pts" with R5/6 CLI at a single hospital
Inclusion criteria: All diabetic foot operations 2006–2008 vs. 2008-2010
Exclusion criteria: N/A
with tissue loss who achieved complete wound healing after endovascular
Inclusion criteria: CLI pts
Exclusion criteria: N/A
1 of publication for details
revascularization"; see Table
Inclusion criteria: CLI "ineligible for

CLI indicates critical limb ischemia; IPC, intermittent pneumatic compression; and N/A, not applicable.

conditions (e.g., arthritis, overuse injuries, simple infections in nondiabetics, venous ulcers, minor trauma,

radiculopathy)

evaluated for benign

failure to heal (OR: 0.21; p<0.01; 95%

may help to minimize hospital

readmissions in the high-risk

population.

amputation prevention program

mandatory. A coordinated

therefore surveillance is

12 wk. Hindfood wounds predictive of

CI: 0.06-0.68).

and average time to full healing was

revascularizations. Complete wound healing observed in 59% of wounds,

Exclusion criteria: New pts

from March 2012–July 2013. Pts at highest risk for limb loss, defined as ischemic wounds (ischemic ulcer or gangrene) or diabetic foot

were prformed, with an equal distribution of endovascular and open

Results: 67% of wounds were present >6 wk before referral. A total of 151 podiatric and 86 vascular interventions

or ankle wounds can adversely influence the outcome. Healing

can be prolonged and a

neuroischemic wounds. Hindfoot

with limb threatening

substancial proportion of pts can

be expected to have a recurrence,

ulcers.

Size: n=91 limbs from 89 pts

amputation prevention clinic

onset of symptoms of ALI. Exclusion criteria:
-
Exclusion criteria: N/A
between 2003 and 2011 (ALI vs. CLI)
Inclusion criteria: All pts undergoing infrainguinal
Exclusion criteria: ALI
Inclusion criteria: All pts w ALI
Exclusion criteria: N/A
Inclusion criteria: All pts with ALI
Exclusion criteria: N/A

al.       Study type: Registry       Inclusion criteria: erripheral blood vessels       1° endpoint: Amputation and mortality exacularization         al.       Study type: Registry       Inclusion criteria: erripheral blood vessels       1° endpoint: Amputation and mortality exacularization         Size: n=200 pts       Enrolled in the FRIENDS registry       1° endpoint: Amputation and mortality exclusion criteria: • N/A         V/A       Exclusion criteria: • N/A       1° endpoint: Amputation and mortality exocutation of limb ischemia in pts with AL associated with much higher rates of first amputation (p= 0.002) and worse ampu free survival (p=0.037). No significant associated with AL was associated with progressive increased 30-day ambputation (p=0.028 trend)	vy angiography; HR, hazard ratio; LE, lower ext	ALI indicates acute limb ischemia; CI, confidence interval; CFA, common femoral artery; CLI, critical limb; CTA, computed tomography angiography; HR, hazard ratio; LE, lower extre	terval; CFA, common femoral a	ALI indicates acute limb ischemia; CI, confidence interval; CFA, common femoral a	ALI indicates acute limb
al.       Study type: Registry       Inclusion criteria: erroled in the peripheral blood vessels       10.5% of pts achieved successful revascularization         al.       Study type: Registry       Inclusion criteria: enclusion criteria: bischemia       1° endpoint: Amputation and mortality         Study type: Registry       Inclusion criteria: enclusion criteria: • Limb threatening ischemia       1° endpoint: Amputation and mortality         Mathematical Size: n=200 pts       Enrolled in the FRIENDS registry       1° endpoint: Amputation and mortality         • N/A       Exclusion criteria: • N/A       • N/A         • N/A       • N/A       • Increased duration of limb ischemia in pts with ALI was associations were observed in pts with CLI. • Increased 30-day ambputation (p=0.028 for		trend)			
al.       Study type: Registry       Inclusion criteria: erripheral blood vessels       1° endpoint: Amputation and mortality exacularization         al.       Study type: Registry       Inclusion criteria: erripheral blood vessels       1° endpoint: Amputation and mortality exacularization         Size: n=200 pts       Enrolled in the FRIENDS registry       1° endpoint: Amputation and mortality exacularization         VIA       Exclusion criteria: • N/A       1° endpoint: Amputation and mortality exacularization         with ALI was associated with progressively       • N/A		increased 30-day ambputation (p=0.028 for			
al.       Study type: Registry       Inclusion criteria: • Enrolled in the FRIENDS registry       Inclusion criteria: • Limb threatening • Enrolled in the FRIENDS registry       1° endpoint: Amputation and mortality • Unation of limb ischemia in pts with ALI was amputation (p= 0.002) and worse amputation • 3.2% of pts had a fatal outcome • 10.5% of pts had a fatal ou		with ALI was associated with progressively			
al.       Study type: Registry       Inclusion criteria: ischemia       1° endpoint: Amputation and mortality ischemia         al.       Study type: Registry       Inclusion criteria: ischemia       1° endpoint: Amputation and mortality ischemia         Size: n=200 pts       Exclusion criteria: ischemia       1° endpoint: Amputation and mortality ischemia         • Enrolled in the FRIENDS registry       Exclusion criteria: ischemia       1° endpoint: Amputation and mortality ischemia         • M/A       • N/A       • N/A		<ul> <li>Increased duration of limb ischemia in pts</li> </ul>			
al.       Study type: Registry       Inclusion criteria: • Enrolled in the FRIENDS registry       Inclusion criteria: • Enrolled in the FRIENDS registry       1º endpoint: Amputation and mortality • Study type: Registry         al.       Study type: Registry • Limb threatening ischemia       Inclusion criteria: • Enrolled in the FRIENDS registry       1º endpoint: Amputation and mortality • Duration of limb ischemia in pts with ALI was associated with much higher rates of first amputation (p= 0.002) and worse amputation- free survival (p=0.037). No significant		associations were observed in pts with CLI.	• N/A		
al.       Study type: Registry       Inclusion criteria: • Enrolled in the FRIENDS registry       Inclusion criteria: • Enrolled in the FRIENDS registry       1º endpoint: Amputation and mortality • Study type: Registry         al.       Study type: Registry • Limb threatening sichemia       Inclusion criteria: • Enrolled in the FRIENDS registry       1º endpoint: Amputation and mortality • Study type: Registry         al.       Study type: Registry • Limb threatening sichemia       Inclusion criteria: • Enrolled in the FRIENDS registry       1º endpoint: Amputation and mortality • Duration of limb ischemia in pts with ALI was amputation (p= 0.0002) and worse amputation.		free survival (p=0.037). No significant	Exclusion criteria:		
al.       Study type: Registry       Inclusion criteria: • Limb threatening ischemia       Inclusion criteria: • Enrolled in the FRIENDS registry       1º endpoint: Amputation and mortality • Study type: Registry         al.       Study type: Registry • Limb threatening ischemia       Inclusion criteria: • Limb threatening ischemia       1º endpoint: Amputation and mortality • Duration of limb ischemia in pts with ALI was associated with much higher rates of first		amputation (p= 0.0002) and worse amputation-			
al.       Study type: Registry       Inclusion criteria: • Enrolled in the       Inclusion criteria: • Enrolled in the       1° endpoint: Amputation and mortality • Duration of limb ischemia in pts with ALI was	deadly, even with limb revascularization	associated with much higher rates of first	FRIENDS registry		
al.       Study type: Registry       Inclusion criteria:       1° endpoint: Amputation and mortality         Size: n=200 pts       Inclusion criteria:       1° endpoint: Amputation and mortality         Results:       Results:	<ul> <li>Limb ALI episodes are extreamly</li> </ul>	<ul> <li>Duration of limb ischemia in pts with ALI was</li> </ul>	<ul> <li>Enrolled in the</li> </ul>		
al.       Study type: Registry       Inclusion criteria:       Inclusion criteria:       1° endpoint: Amputation and mortality         al.       Study type: Registry       Inclusion criteria:       1° endpoint: Amputation and mortality	of salvage	Results:	ischemia	Size: n=200 pts	25262269
Previous reconstructive       (73.7%) compared to a chronic lesion (26.3%);         procedures on blood       p<0.05	in ALI occur, the less likely the possibility		<ul> <li>Limb threatening</li> </ul>		2014 (381)
<ul> <li>(73.7%) compared to a chronic lesion (26.3%); p&lt;0.05</li> <li>86.2% of pts achieved successful revascularization</li> <li>3.2% of pts had mputating treatment ≤30 d.</li> <li>10.5% of pts had a fatal outcome</li> </ul>	<ul> <li>The longer lower extremity symptoms</li> </ul>	1° endpoint: Amputation and mortality	Inclusion criteria:	Study type: Registry	Duval S, et al.
<ul> <li>(73.7%) compared to a chronic lesion (26.3%); p&lt;0.05</li> <li>86.2% of pts achieved successful revascularization</li> <li>3.2% of pts had mputating treatment ≤30 d.</li> <li>10.5% of pts had a fatal outcome</li> </ul>	embolism)				
<ul> <li>(73.7%) compared to a chronic lesion (26.3%); p&lt;0.05</li> <li>86.2% of pts achieved successful revascularization</li> <li>3.2% of pts had mputating treatment ≤30 d.</li> <li>10.5% of pts had a fatal outcome</li> </ul>	the clause of ischemia (thrombosis or		peripheral blood vessels		
<ul> <li>ive (73.7%) compared to a chronic lesion (26.3%);</li> <li>p&lt;0.05</li> <li>e 86.2% of pts achieved successful revascularization</li> <li>- 3.2% of pts had mputating treatment ≤30 d.</li> </ul>	operative treatment in ALI, regardless of	<ul> <li>10.5% of pts had a fatal outcome</li> </ul>	aneurysmal disease of the		
<ul> <li>ive (73.7%) compared to a chronic lesion (26.3%);</li> <li>p&lt;0.05</li> <li>e 86.2% of pts achieved successful revascularization</li> </ul>	This demonstrates benefits of early	<ul> <li>3.2% of pts had mputating treatment ≤30 d.</li> </ul>	induced by trauma or		
<ul> <li>ive (73.7%) compared to a chronic lesion (26.3%);</li> <li>p&lt;0.05</li> <li>e 86.2% of pts achieved successful</li> </ul>	revascularization of LE achieved in 85%.	revascularization	ischemia had been		
ive (73.7%) compared to a chronic lesion (26.3%); p<0.05	<ul> <li>High success rate, with successful</li> </ul>	<ul> <li>86.2% of pts achieved successful</li> </ul>	vessels and where acute		
(73.7%) compared to a chronic lesion (26.3%);	decompensation	p<0.05	procedures on blood		
	chronic myocardiopathy and metabolic	(73.7%) compared to a chronic lesion (26.3%);	Previous reconstructive		

MI, myocardial infarction; N/A, not applicable; OR, odds ratio; and RR, relative risk. dremity;

## Evidence Table 46. Nonrandomized Trials, Observational studies, and/or Registries Comparing Evaluating Noninvasive Testing and Angiography for ALI–Section 9.1.

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			& 95% CI)	
Morris-Stiff G, et al.	Study type: Retrospective	Inclusion criteria: Pts	Results: Despite increased pre-operative	<ul> <li>Delay from symptom onset to</li> </ul>
2009 (382)	review comparing pts with	presenting with ALI during	(15% vs. 47%; p<0.05) and on-table imaging	surgery is a major determinant of
<u>19785938</u>	ALI from 2 time periods	specified time period	(0% vs. 16%; p<0.05) technical success did	outcome.
			not improve.	
	Size: n=205 pts	Exclusion criteria: N/A		
Londero LS, et al.	Study type: Prospective	Inclusion criteria: All	1° endpoint: 30 pts needed immediate	<ul> <li>If CT or MRA was used the</li> </ul>
2014 (383)	cross-sectional cohort		intervention. In the group of 14 pts who had	intervention was delayed by 3 h
25400690	study including all pts	Exclusion criteria: N/A	immediate operation, the median time from	<ul> <li>No clear delay to angiography, but</li> </ul>
	suspected with ALI		vascular evaluation to revascularization was	thrombolysis duration was longer
			324.5 (122–873) min and in the group of 8 pts	than surgery
	Size: n=42 pts		that went through an imaging procedure	

5.	complications in			IIrokinase vs. surgery Exclusion criteria. Pts were excluded	IIrokinase vs. surgerv	8201703
ary	cardiopulmonary	<ul> <li>Limb salvage 82% at 12 mo both</li> </ul>	directed urokinase		directed Intra-arterial	1994 (384)
	<ul> <li>Increased</li> </ul>	<u>1° endpoint:</u>	Intervention: Catheter	Inclusion criteria: ALI <7 d	Aim: Catheter	Ouriel K, et al.
vents	Adverse Events	95% ĆI)	(# patients)			Year Published
ations:	Study Limitations:	or RR; &	Study Comparator		Study Size (N)	Author:
Endpoint	Relevant 2° Endpoint	Endpoint Results (Absolute Event Rates. P value: OR	Study Intervention	Patient Population	Aim of Study; Study Type:	Study Acronym:
			ion 9.2.2.	Evidence Table 47. RCTs of Revascularization Strategy for ALI-Section 9.2.2.	47. RCTs of Revas	Evidence Table
sound arterial	۹M, duplex ultra	DSA, digital subtraction angiography; DUAM, duplex ultrasound arterial it; and RR, relative risk.	^r , computed tomography; [ ; OR, odds ratio; pt, patien	ALI indicates acute limb ischemia; CI, confidence interval; CLI, critical limb ischemia; CT, computed tomography; DSA, digital mapping; HR, hazard ratio; N/A, not applicable; MRA, magnetic resonance angiography; OR, odds ratio; pt, patient; and RR,	limb ischemia; Cl, confid rd ratio; N/A, not applicab	ALI indicates acute mapping; HR, haza
			(1686–8376) min.			
		Ibolysis was 5621	treated with arterial thrombolysis was 5621			
		4 pts, who were	revascularization among 4 pts, who			
		edian time for	revascularization. The median time			
		cialist assessment to	(494–1185) min from specialist assessment to			
		nedian delay was 822	before an operation the median delay was 822			

Study	Aim of Study;	Patient Population	Study Intervention	Endpoint Results	Relevant 2° Endpoint
Acronym;	Study Type;		(# patients) /	(Absolute Event Rates, P value; OR	(if any);
Author;	Study Size (N)		Study Comparator	or RR; &	Study Limitations;
Year Published			(# patients)	95% CI)	Adverse Events
Ouriel K, et al.	Aim: Catheter	Inclusion criteria: ALI <7 d	Intervention: Catheter	1° endpoint:	<ul> <li>Increased</li> </ul>
1994 (384)	directed Intra-arterial		directed urokinase	<ul> <li>Limb salvage 82% at 12 mo both</li> </ul>	cardiopulmonary
<u>8201703</u>	urokinase vs. surgery	Exclusion criteria: Pts were excluded		groups	complications in
		from study if they manifested a	Comparator: Surgery	<ul> <li>Survival 84% IAT vs. 58% surgerv</li> </ul>	surgery group 49% vs.
	Study type: RCT	contraindication to thrombolytic therapy,		at 12 mo. p=0.01	16%, p=0.001
		including one or more of the following: a		-	
	Size: n=57 pts IAT	major operative procedure within 14 d,			
	vs. n=57 pts surgery	active peptic ulcer disease, an			
		intracranial neoplasm, or a Hx of a			
		cerebrovascular accident. Pts were also			
		excluded if they had a contraindication			
		to operative revascularization; non-			
		ambulatory prior to ALI or Cr>2.5			
TOPAS	Aim: Catheter	Inclusion criteria: ALI ≤14 d	Intervention: Catheter	1° endpoint: 6 mo amputation free	N/A
Ouriel K, et al.	directed Intra-arterial		directered urokinase	survival 71.68 IAT vs. 74.8 surgery	
1998 (356)	urokinase vs. surgery	Exclusion criteria: pts ineligible for		p=0.43	
<u>9545358</u>		thrombolytics	Comparator: Surgery		
	Study type: RCT			Safety endpoint: Mortality at hospital	
				discharge 8.8 IAT vs. 5.9 surgery	
	Size: n=272 pts IAT			p=0.19	
	vs. n=272 pts surgery				

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Diffin DC and <u>Aim</u> : Kandarpa K 1996 (387) PIAT <u>8773976</u> tx for of 2 F of 2 F pts) c pts) c	Comerota AJ, et <u>Aim</u> : al. (2017 1996 (386) <u>8795509</u> <u>Stud</u> <u>Size</u> : pts) c pts) c	STILE <u>Aim</u> : C; Graor RA, et al. directed 1994 (385) tPA or u <u>8092895</u> surgery <u>Study t</u> n=112 p 144 pts
Aim: Review the risks and benefits of PIAT vs. SR as initial tx for ALLI Study type: Analysis of 2 RCTs of 2 RCTs Size: SR (n=1,051 pts) or PIAT (n=895 pts)	<u>Aim</u> : Surgery vs. CDT for occluded bypass grafts <u>Study type</u> : RCT <u>Size</u> : Surgery (n=46 pts) or CDT (n=78 pts) pts)	<u>Aim</u> : Catheter directed Intra-arterial tPA or urokinase vs. surgery <u>Study type</u> : RCT <u>Size</u> : n=137 pts tPA, n=112 pts UK, N= 144 pts surgery
Inclusion criteria: Published RCTs that compared PIAT with SR as the initial treatment of ALLI Exclusion criteria: Studied that included >1 disease category but did not specifically stratify results by category	Inclusion criteria: ALI <14 d or chronic ischemia >14 d Exclusion criteria: contra-indications to thrombolysis	<ul> <li>Inclusion criteria:</li> <li>18–90 y</li> <li>Signs or symptoms of worsening limb ischemia within the past 6 mo who required intervention</li> <li>Angiographically documented nonembolic arterial or bypass graft occlusion</li> <li>Exclusion criteria: infected grafts or contraindications to lytics</li> </ul>
Intervention: PIAT Comparator: SR	Intervention: CDT Comparator: Surgery	Intervention: Catheter directed urokinase or tPA <u>Comparator</u> : Surgery
<u>1° endpoint</u> : Limb salvage and mortality at 30 d and 6–12 mo	<ul> <li><u>1° endpoint</u>:</li> <li>A composite clinical outcome including death, amputation, ongoing/recurrent ischemia, and major morbidity was analyzed on an intent-to-treat basis at 30 d and 1 y.</li> <li>Acutely ischemic pts (0–14 d) randomized to lysis demonstrated a trend toward a lower major amputation rate at 30 d (p=0.074) and significantly at 1 y (p=0.026) compared with surgical pts, while those with &gt;14 d ischemia showed no difference in limb salvage but higher ongoing/recurrent ischemia in lytic pts (p&lt;0.001)</li> </ul>	<u>1° endpoint</u> : Composite clinical outcome (see page 255 of mansuscript) 22.6% surgery vs. 38.3% IAT, p=0.011
<ul> <li>Limb salvage rates at 30 d for PIAT vs. SR: 93%; vs. 89%</li> <li>Limb salvage rates at 6–12 mo for PIAT vs. SR: 89%; vs. 73%</li> <li>PIAT better limb- salvage rate and mortality than SR in the treatment of ALLI</li> </ul>	• For ALI <14 d CDT is similar to surgery	<ul> <li>Note: failure of catheter placement occurred in 28% of IAT group resulting in large failure rate</li> <li>Poor quality study</li> </ul>

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	Schrijver AM,, et al. 2011 (388) <u>PMC3033836</u>
	<u>Size</u> : n=60 pts
<ul> <li>Exclusion criteria:</li> <li>Isolated common femoral artery thrombosis</li> <li>Iocalized emboli (&lt;5 cm) or occlusions in the native femoropopliteal arteries</li> <li>Clinical complaints of ALI due to thrombosis of the femoropopliteal or femorocrural venous or prothetic bypass grafts &lt;1 wk and &gt;7 wk</li> <li>ALI class IIb and III Rutherford classification</li> <li>Antiplatelet therapy, anticoagulants, or thrombolytic drugs are contraindicated</li> <li>6 wk surger</li> <li>DBP &gt;110 mm HG, SBP &gt;200 mm Hg</li> <li>Current malignancy</li> <li>Hx of life-threatening reaction to contrast medium</li> <li>Uncorrected bleeding disorders</li> <li>Women with child-bearing potential not on contraceptives or currently breastfeeding</li> </ul>	Inclusion criteria: • Pts age >18 y and <85 y • Pts with thrombosed femoropopliteal or femorocrural native arteries or femoropopliteal or femorocrural venous or prosthetic bypass grafts with ischemic complaints between 1–7 wks • Pts with acute lower limb ischaemia dass I and IIa according to Rutherford classification • Pts understand the nature of the procedure and provide written informed consent
	Intervention: Standard thrombolysis <u>Comparator</u> : US- accelerated thrombolysis
	<u>1° endpoint</u> : Duration of catheter- directed thrombolysis needed for uninterrupted flow in the thrombosed infrainguinal native artery or bypass graft, with outflow through ≥1 crural artery
	• RCT comparing this technique to standard catheter-based thrombolytic therapy failed to demonstrate a difference in outcomes including bleeding despite a lower total amount of lytic delivered

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N/A	<u>1° endpoint:</u> Outcome of arterial bypass reconstruction in the setting of acute arterial ischemia	Inclusion criteria: Acute arterial ischemia and required an urgent/emergent lower-	Study type: Single institution retrospective cohort	Nypaver TJ, et al. 1998 (375) <u>9737621</u>
	<u>Results</u> : N/A	Exclusion criteria: N/A	Size: N/A	1997 (46) <u>9308598</u>
N/A	1° endpoint: Scoring Scheme for ALI	Inclusion criteria: N/A	Study type: Consensus document	Rutherford RB, et al.
	<ul> <li>with embolism</li> <li>Occlusion longer than 24 h (OR: 2.6; 95% CI: 1.1–7.6) was associated with death and amputation in the multivariate analysis</li> <li>Mortality 15 (18.1%)</li> <li>Amputation 24 (28.9%)</li> </ul>			
	Male gender, smoking, and comorbidities were more frequent among pts with thrombosis, and atrial fibrillation was more common among pts	<u>Exclusion criteria:</u> Stage I ischemia	Size: n=83 pts	
<ul> <li>Comorbidities were also more frequent among pts with thrombosis</li> </ul>	1° endpoint: Mortality and amputation Results:	Inclusion criteria: ALI, and etiology	Study type: Single institution prospective cohort (observational)	Fagundes C, et al. 2005 (389) <u>17315606</u>
Summary/Conclusion Comment(s)	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Patient Population	Study Type/Design; Study Size	Study Acronym; Author; Year Published
resentation of ALI–Section 9.2.2.	Evidence Table 48. Nonrandomized Trials, Observational Studies, and/or Registries of Clinical Presen	als, Observational Stud	8. Nonrandomized Tria	Evidence Table 4
rval; DBP, diastolic blood pressure; HR, hazard ratio; hx, intraarterial thrombolysis; pt, patient; RCT, randomized bolysis for Ischemia of the Lower Extremity; TOPAS,	directed thrombolysis; CI, confidence inte plicable; OR, odds ratio; PIAT, peripheral pressure; STILE, Surgery Versus Throm	9r-limb ischemia; CDT, cathe resonance imaging; N/A, no cularization; SBP, systolic blo A, tissue plasminogen activa	ALI indicates acute limb ischemia; ALLI, acute lower-limb ischemia; CDT, catheter- history; IAT, intra-arterial treatment; MRI, magnetic resonance imaging; N/A, not ap controlled trail; RR, relative risk; SR, surgical revascularization; SBP, systolic blood Thrombolysis or Peripheral Arterial Surgery; and tPA, tissue plasminogen activator	ALI indicates acute lim history; IAT, intra-artei controlled trail; RR, re Thrombolysis or Perip
		<ul> <li>study</li> <li>Life expectancy of &lt;1 mo</li> <li>Contraindication for MRI</li> </ul>	study • Life • Cor	

pregnancy
Hemodynamically unstable at the onset of the procedure
Pts who refuse treatment

Currently participating in another

Baril DT, et al. 2013 (379) <u>23714364</u>		10074242	de Donato G, et al. 2014 (378)	<u>19298933</u>	Eliason JL and Wakefield TW 2000 (301)		Shin HS, et al. 2013 (377) <u>24436594</u>	Fogarty TJ, et al. 1963 (390) <u>13945714</u>	
<u>Study type</u> : Registry review		Size: n=322 pts	Study type: Single institution cohort	<u>Size</u> : n=18 studies	<u>Study type</u> : Review article	onsecutive pts	Study type: Single institution	<u>Study type</u> : Descriptive <u>Size</u> : N/A	<u>Size</u> : n= 71
Inclusion criteria: All pts undergoing infrainguinal lower		Exclusion criteria: ALI from graft thrombosis	Inclusion criteria: All pts w ALI		Inclusion criteria: N/A	EXCIUSION CRIERIA: N/A	Inclusion criteria: All pts with ALI	Inclusion criteria: N/A Exclusion criteria: N/A	extremity arterial bypass reconstruction <u>Exclusion criteria</u> : N/A
<u>1° endpoint</u> : Major amputation and mortality <u>Results</u> : ALI predictor of both major amputation	<ul> <li>Limb salvage</li> <li>Overall survival rates</li> <li><u>Results</u>: Reduction in complications when hybrid techniques utilized as opposed to just thromboembolectomy</li> </ul>	<ul> <li>30 d mortality</li> <li>Primary and secondary patency reintervention rate</li> </ul>	<ul> <li>In-hospital complications</li> </ul>	<u>nesulis</u> . IVA	<u>1° endpoint</u> : N/A	<u>Kesuits</u> : N/A	<u>1° endpoint:</u> Limb salvage via novel surgical approach	<u>1° endpoint</u> : N/A <u>Results</u> : N/A	<ul> <li>Results:</li> <li>Mean duration of symptoms was 43 h (median 24), and mean time from hospital presentation to the operating room was 36 h (median 12)</li> <li>Death, limb loss, or both, were associated with a paralytic limb (p=0.001) and congestive heart failure (p=0.03)</li> </ul>
<ul> <li>Age and gender similar to CLI</li> <li>ALI less likely to be on ASA (63% vs. 75%; p&lt;0.0001) or a statin (55% vs. 68%;</li> </ul>		<ul> <li>30 d mortality 4.4%</li> <li>15% in hospital complications</li> <li>8 pts with complication from catheter</li> </ul>	<ul> <li>Thromboembolectomy alone in 35%</li> <li>45.5% via CFA</li> </ul>	<ul> <li>The majority of the lethal events associated with IR injury occur with acute lung injury as a prominent component of the multiple organ dysfunction syndrome</li> </ul>	<ul> <li>Compartment pressures are easily measured through multiple methods of</li> </ul>	<ul> <li>14% had a Hx of previous MI</li> <li>86% of pts with mixed thromboembolic disease</li> <li>Below knee exposure and 1 vessel runoff</li> </ul>	<ul> <li>CTA for Dx</li> <li>71% heart disease:</li> <li>57% atrial fibrillation</li> </ul>	<ul> <li>First description of embolectomy catheter</li> </ul>	

lower in the CDT and CDTA groups	was significantly	Combined endpoint of readmission and AFS	in combination	occlusion, or the class of ischemia, individually or	therapy treatment, pt characteristics, location of the required amp	Elective admission, no    No association between the choice of initial required a thi	Exclusion criteria: Results: required a se		Size: n=1,074 pts of nonembolic ALI and AFS • Initial treatm	onset of their symptoms complications, relief of ischemia, limb salvage,	2015 (392) institution review treated within 14 d of number and type of reinterventions, native vessel	Lurie F, et al. Study type: Multiple Inclusion criteria: Pts 1° endpoint: Clinical and technical outcomes, • The cause	intervention (	or a prior ipsi	Exclusion criteria: N/A ipsilateral byp	vs. 39%; p<0	(ALI vs. CLI) (49%	procedures   between 2003 and 2011   (HR: 1.41; Cl: 1.09–1.83; p=0.009 at 1 y • ALI more lii	Size: n=323 bypass extremity bypass (HR: 2.16; CI: 1.38–3.40; p=0.001) and mortality p<0.0001)
				ly or	required amputation, and 2 (1%) died	al required a third intervention, 5 (2.4%)	required a second intervention, 11 (5.4%)	primary success of 67.3%. 60 pts (29.7%)	<ul> <li>Initial treatment resulted in an overall</li> </ul>		native vessel in 115 pts (56.1%) and an	<ul> <li>The cause of ALI was an occluded</li> </ul>	intervention (41% vs. 29%; p=0.001)	or a prior ipsilateral percutaneous	ipsilateral bypass (33% vs. 24%; p=0.004)	vs. 39%; p<0 .0001), to have had a prior	(49%)	<ul> <li>ALI more likely to be current smokers</li> </ul>	

angiography; DUAM, duplex ultrasound arterial mapping; HR, hazard ratio; MI, myocardial infarction; MRA, magnetic resonance angiography; N/A, not applicable; NEJM, New England Journal of Medicine; NIS, National Inpatient Sample; OR, odds ratio; pt, patient; and RR, relative risk. catheter directed thrombolysis; CFA, common femoral artery; CI, confidence interval; CLI, critical limb ischemia; CTA, computed tomography angiography; DSA, digital subtraction angiography; DUAM, duplex ultrasound arterial mapping; HR, hazard ratio: MI. mvncardial infanction: MRA manadia manadia

Evidence Table 49. Nonrandomized Trials, Observational Studies, and/or Registries of Diagnostic Evaluation of the Cause of ALI–Section 9.2.2. (There is no literature specifically addressing the diagnostic work up for the cause of ALI. This large single-center series does give etiologies. Echocardiography and telemetry seem reasonable for those without underlying DAD. Economic for hyperconductive state specifically addressing the diagnostic work up for the cause of ALI. This large single-center series does give etiologies. Echocardiography and telemetry seem reasonable for those without underlying DAD. Economic for hyperconductive state specifically addressing to a second the second s

reasonable for those with	out underlying PAD. Focusec	l evaluation for hypercoagulable	reasonable for those without underlying PAD. Focused evaluation for hypercoagulable state seems reasonable in those with native artery thrombosis.)	ive artery thrombosis.)
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			& 95% Cl)	
Taha	Study type: Single	Inclusion criteria: ALI pts	1° endpoint: Technical success,	<ul> <li>Underlying cause of ALI retrieved from</li> </ul>
2015 (393)	center retrospective	cared for my vascular	incidence of postoperative	medical record, cause by percent: cardiac
25080883	review comparing open	surgeons. All with embolism	complications, length of hospital stay,	embolism 17.7; native artery thrombosis 26.2;
	and endovascular repair	or thrombosis as etiology.	loss of primary patency, loss of	failed stent 17.9; failed bypass graft 33.5;
	in ALI		assisted primary patency, and loss of	thrombosed peripheral aneurysm 4.7
		Exclusion criteria: Trauma	secondary patency as well as	
	Size: n=473 pts	as etiology of ALI, blue toe	amputation and mortality rates at 30 d	
		syndrome	and 1 v	

ALI indicates acute li	I mb ischemia: N/A, not applicable	ا <b>Kesuits:</b> ۱/A ALI indicates acute limb ischemia; N/A, not applicable; PAD, peripheral artery disease; and pt, patient	N/A	
Evidence Table :	50. Nonrandomized Trials	s, Observational Studies, and/or Re	Evidence Table 50. Nonrandomized Trials, Observational Studies, and/or Registries of Revascularization Strategy for ALI–Section 9.2.2.	· ALI–Section 9.2.2.
Study Acronym; Author; Year Published	Study Type/Design; Study Size	Patient Population	Primary Endpoint and Results (include P value; OR or RR; & 95% Cl)	Summary/Conclusion Comment(s)
Gupta R and Hennebry TA	Study type: Case series	Inclusion criteria: ALI <14 d treated with Trellis device	1° endpoint: Limb salvage=100%	<ul> <li>Proof of concept</li> <li>Level C data</li> </ul>
2012 (394)	Size: n=24 pts		Results: In hospital and 30 d mortality=4.16%	
<u>22511320</u>		Exclusion criteria: Vessel size less than 3 mm diameter or distal location or		
		contrast intolerance, as assessed by the		
		treating clinician's discretion		
Ansel GM, et al. 2008(395)	Study type: Case series	Inclusion criteria: ALI <14 d treated with pharmaco-mechanical	<u>1° endpoint:</u> Limb salvage	<ul> <li>Level C data</li> </ul>
<u>18726955</u>	Size: n=29 limbs treated in 119 pts	thrombectomy±catheter directed lysis	<b>Results:</b> In-hospital success with limb salvage was attained in 96.5% (n=55) with mortality of	
		Exclusion criteria: Pts felt to have	3.5% (n=2). 30 d limb salvage and mortality were	
		and evaluated pts with only arterial	mean 5 y follow-up (mean=62 mo), 3 pts have	
		thrombosis as the inciting event.	64.7%) are available Amountation free survival	
			was 94.7% (n=36/38) with long-term mortality rate of 29.6% (n=16/54).	
Byrne RM, et al. 2014 (396)	Study type: Case series	Inclusion criteria: ALI treated with PMT±CDT	<ol> <li><u>1° endpoint</u>: Technical success was achieved in</li> <li>83.8% of cases, with a 30 d mortality rate of 5.2%</li> </ol>	<ul> <li>Level C data</li> </ul>
24360240	Size: n=154 limbs were	1 - - -		
			15.0% (18.1% for CDT only, 11.3% for PMT; p=NS)	
Taha AG, et al. 2015 (393)	Study type: Retrospective comparison of endo vs. OR	Inclusion criteria: ALI	1° endpoint: Amputation and mortality at 1 y	<ul> <li>Equal amputation rates</li> <li>Endo had lower 30 d</li> </ul>
25080883	Size: n=154 limbs were	Exclusion criteria: Blue toe syndrome and acute ischemia secondary to trauma	<ul> <li>Overall amoutation rates were 13.5% (OR) vs.</li> </ul>	● I evel C data
	treated in 147 pts in the ER group, compared with 326	or dissection were excluded	6.5% (ER) at 30 d (p=0.023) and 19.6% (OR) vs. 13.0% (ER) at 1 y (p=0.074)	

Ascher et al. 1999 (402) <u>12712369</u>	Elmandy MG, et al. 2010 (401) <u>20934653</u>	Allie DE, et al. 2004 (400) <u>15558768</u>	Kasirajan K, et al. 2001 (399) <u>11287526</u>	Silva JA, et al. 1998 (398) <u>9863742</u>	2014 (397) <u>24933285</u>	Schernthaner MB, et al.
Study type: Retrospective, bypass for CLI performed using ultrasound alone or	<u>Study type</u> : Prospective <u>Size</u> : n=97 pts	<u>Study type</u> : Case series <u>Size</u> : n=49 pts	<u>Study type</u> : Retrospective analysis <u>Size</u> : n=86 pts (acute, n=65; subacute, n=21); acute <14 d; suacute 14 d– 4 mo	<u>Study type</u> : Case series <u>Size</u> : n=21 pts	CDT in pts with acute and subacute limb ischemia. <u>Size:</u> n=UAT was performed in 75 pts, and CDT was performed in 27 pts	limbs in 296 pts in the OR group <u>Study type:</u> Retrospective series; UAT and standard
Inclusion criteria: Need for infra inguinal arterial bypass	<b>Exclusion criteria:</b> Non traumatic ALI <b>Exclusion criteria:</b> Past Hx of peripheral arterial graft, traumatic limb ischemia, dissection, and thrombosis induced by vasospasm, arteritis, popliteal cyst, or entrapment.	Inclusion criteria: ALI treated with rheolytic thrombectomy catheter with thrombolytic solution priming agent Exclusion criteria: None reported	<u>Inclusion criteria</u> : ALI (acute or subacute) <u>Exclusion criteria</u> : None reported	Inclusion criteria: ALI ≤14 d treated with rheolytic thrombectomy Exclusion criteria: None reported	Exclusion criteria: None reported	Inclusion criteria: ALI or subacute limb ischemia
<u>1° endpoint</u> : Adequacy of ultrasound to diagnose stenosis	<ul> <li><u>1° endpoint</u>: Agreement with surgical determination of embolic or thrombotic</li> <li><u>Results</u>: <ul> <li>Clinical characteristics similar in embolic and thrombotic groups</li> <li>Greater difference in diameter of artery compared with contralateral artery diameter identified embolic etiology</li> </ul> </li> </ul>	<u>1° endpoint</u> : 30 d limb salvage=91% <u>Results</u> : No significant difference between power pulse with UK or TNK; however no comparator group using catheter directed lytic delivery	<u>1° endpoint</u> : Angiographic success=61.4% <u>Results</u> : 1 mo amputation and mortality rates were 11.6% and 9.3%	<u>1° endpoint</u> : Limb salvage <u>Results</u> : The overall 6 mo survival was 81% (17 pts), and limb salvage occurred in 16 of 18 limbs (89%) in the 17 pts	<ul> <li><u>Results:</u></li> <li>No difference in limb salvage</li> <li>Major and minor bleeding combined was lower:</li> <li>6.7% (UAT) vs. 22.2% (CDT) (p=0.025) despite</li> <li>no difference in lytic dose</li> </ul>	<ul> <li>30 d mortality rate was 13.2% (OR) and 5.4% (ER) (p=0.012)</li> <li>1° endpoint: Limb salvage</li> </ul>
• Duplex took 100 min angiography required in 2 pts due to arterial	<ul> <li>Duplex provided information on etiology that could guide treatment</li> </ul>	Proof of concept     Level C data	<ul> <li>Level C data</li> <li>Mixed population</li> </ul>	<ul> <li>Proof of concept</li> <li>Level C data</li> </ul>		<ul> <li>Pilot data – level C</li> </ul>

Schrijver AM, et al. 2012 (405) <u>21534002</u>		Leung DA, et al. 2015 (404) <u>26109628</u>	Lowery AJ, et al. 2007 (403) <u>17628263</u>	
<u>Study type</u> : Prospective cohort <u>Size</u> : n=21 consecutive pts	<u>Size</u> : n=283 pts	Study type: Rheolytic thrombectomy registry study	<u>Study type</u> : Prospective evaluation of US, MRA, DSA <u>Size</u> : n=465 pts	ultrasound + angiography
Inclusion criteria: Pts with aotrofemoral arterial thromboembolic obstructions Exclusion criteria: N/A	<u>Exclusion criteria</u> : N/A	Inclusion criteria: Pts with ALI undergoing treatment with the AngioJet System	Inclusion criteria: All pts with CLI being considered for endovascular revascularization Exclusion criteria: N/A	Exclusion criteria: Contrast allergy
<ol> <li><u>endpoint</u>: 30-d technical and clinical outcome of US-accelerated thrombolysis</li> <li><u>Results</u>: Complete thrombolysis (&gt;95% lysis of thrombus) was achieved in 20 pts; in 9 pts within 24 hours. Median ankle-brachial index (ABI) increased from 0.28 (range, 0-0.85) to 0.91</li> </ol>	<b>Results:</b> 83% achieved procedure success. 52% of procedures completed without the need for adjunctive CDT. 12-mo follow-up, 81% amputation free survival and 91% freedom from mortality, 91% freedom from bleeding requiring transfusion, 95% freedom from renal failure. Significantly better outcomes in pts without infrapopliteal involvement and those who underwent PMT without CDT. Higher rates of procedure success (p=0.021), 12-mo freedom from amputation free survival (p=0.028), and 12-mo freedom from amputation (p=0.01) in the PMT without CDT group	<u>1° endpoint</u> : Procedure success, 12-mo amputation free survival, 12-mo freedom from amputation	<u>1° endpoint</u> : Compared clinical pragmatism, hemodynamic outcomes, and cost-effectiveness when using DUAM alone compared to DSA or MRA as preoperative assessment <u>Results</u> : In the DUAM group, 43 lesions were identified and marked at the time of preoperative DUAM, all of which were treated at angioplasty. In the DSA group, 53 lesions identified appreoperatively were treated at angioplasty. In the MRA group, 58 lesions were identified as requiring treatment on the preoperative MRA. Only 50 of these required angioplasty.	Results: Adequate map by ultrasound alone in the majority of pts
• This feasibility study showed a high technical success rate of US- accelerated thrombolysis for aortofemoral arterial obstructions. US- accelerated thrombolysis		• PMT had more positive results as a first line treatment for ALI	<ul> <li>ALI</li> <li>US and DSA are reasonable, MRA may have overestimated stenosis</li> <li>Not clear if any pts had ALI</li> <li>Similar results from Hingorani and Soule, different from Cambria</li> </ul>	<ul><li>calcification</li><li>Not clear if any pts had</li></ul>

<ul> <li>Secc commo interve &gt;80% - &gt;80%</li></ul>	<ul> <li><u>1° endpoint</u>: Graft failures and time points</li> <li><u>Results:</u> <ul> <li>Occlusions-21.6%</li> <li>Salvage procedure-16% (40.5% done at 6 mo)</li> <li>56.6% occlusion preceded by stenosis</li> <li>Primary occlusions: 95.9% in the prosthetic group and 66.5% in the femorocrural group</li> <li>Twice as many stenosis in venous conduits than the prosthetic ones</li> <li><u>1° endpoint</u>: No. of evaluations and interventions to prevent graft occlusion after the threshold criteria based on existent literature (HVC defined as PSV &lt;45 cm/sec: an ABI decrease &gt;0.15)</li> </ul> </li> </ul>	Exclusion criteria:       None reported         Inclusion criteria:       Infrainguinal         lower limb grafts with duplex u/s         surveillance (0, 1, 3, 6, 12 and 18         mo)         Exclusion criteria:         None reported         every 3 mo for 1 y and every 6 mo         thereafter for another y         Exclusion criteria:         Locusion cri	Size:     n=69 pts     E       Study type:     In       Observational     In       Size:     n=212 grafts     In       (197 pts)     In     In       Study type:     In     In       Observational     In     In       Study type:     In     In	Carter A, et al. 2007 (410) <u>17980793</u> Westerband A, et al. 1997 (411) <u>9061138</u>
	<ol> <li><u>1° endpoint</u>: Graft failures and time points</li> <li><u>Results:</u> <ul> <li>Occlusions-21.6%</li> <li>Salvage procedure-16% (40.5% done at 6 m</li> <li>Transport of the prosthetic group and 66.5% in the femorocrural group</li> <li>Twice as many stenosis in venous conduits than the prosthetic ones</li> <li><u>1° endpoint</u>: No. of evaluations and interventition prevent graft occlusion after the threshold criteria based on existent literature (HVC define as PSV &gt;300 cm/sec and Vr &gt;3.5; LVC define</li> </ul> </li> </ol>	<u>Exclusion criteria</u> : None reported nolusion criteria: Infrainguinal ower limb grafts with duplex u/s surveillance (0, 1, 3, 6, 12 and 18 mo) <u>Exclusion criteria</u> : None reported every 3 mo for 1 y and every 6 mo hereafter for another y	9 pts onal 12 grafts onal onal 8 pts (101	Carter A, et al. 2007 (410) <u>17980793</u> Westerband A, et al. 1997 (411) <u>9061138</u>
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<u>о</u>	<ol> <li><u>1° endpoint</u>: Graft failures and time points</li> <li><u>Results:</u> <ul> <li>Occlusions-21.6%</li> <li>Salvage procedure-16% (40.5% done at 6 m</li> <li>Salvage procedure-16% (40.5% done at 6 m</li> <li>Salvage procedure-16% (40.5% done at 6 m</li> <li>The procedure proceded by stenosis</li> <li>Primary occlusions: 95.9% in the prosthetic group and 66.5% in the femorocrural group</li> <li>Twice as many stenosis in venous conduits than the prosthetic ones</li> <li><u>1° endpoint</u>: No. of evaluations and interventition to prevent graft occlusion after the threshold</li> </ul> </li> </ol>	<u>Exclusion criteria</u> : None reported <u>Inclusion criteria</u> : Infrainguinal ower limb grafts with duplex u/s surveillance (0, 1, 3, 6, 12 and 18 mo) <u>Exclusion criteria</u> : None reported <u>Inclusion criteria</u> : OFDS and ABI nol Svery 3 mo for 1 y and every 6 mo	9 pts onal	Carter A, et al. 2007 (410) <u>17980793</u> Westerband A, et al.
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	<ul> <li><u>1° endpoint</u>: Graft failures and time points</li> <li><u>Results:</u></li> <li>Occlusions-21.6%</li> <li>Salvage procedure-16% (40.5% done at 6 m</li> <li>Salvage procedure-16% (40.5% done at 6 m</li> <li>Primary occlusions: 95.9% in the prosthetic</li> </ul>	Exclusion criteria: None reported nclusion criteria: Infrainguinal ower limb grafts with duplex u/s surveillance (0, 1, 3, 6, 12 and 18 no) Exclusion criteria: None reported	9 pts onal 12 grafts	Carter A, et al. 2007 (410) <u>17980793</u>
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	oint: Graft failures and tim	<u>Exclusion criteria</u> : None reported <u>inclusion criteria</u> : Infrainguinal ower limb grafts with duplex u/s surveillance (0, 1, 3, 6, 12 and 18		Carter A, et al. 2007 (410) 17980793
	<u>1° endpoint:</u> Graft failures and time points	Exclusion criteria: None reported nclusion criteria: Infrainguinal ower limb grafts with duplex u/s		Carter A, et al. 2007 (410)
	<u>1° endpoint</u> : Graft failures and time points	Suplex u/s for 1y Suplex u/s for 1y <u>Exclusion criteria</u> : None reported <u>nclusion criteria</u> : Infrainguinal		Carter A, et al.
		Juplex u/s for 1y <u>Exclusion criteria</u> : None reported		
	<ul> <li>14% occluded</li> </ul>	Suplex u/s for 1y		
	<ul> <li>42% recurrent stenosis</li> </ul>	uplex u/s for 1y		
	<ul> <li>43% free of major stenosis/ bypass occlusion</li> </ul>	Inianiguna bypasses momores with	_	
Comment(s)     Secondary interventions are     common however such frequent	Results:	nfrainm linal hunaceae monitorad with	study	26482995
Comment(s)     Secondary interventions are		Pts with primary PTA for autologous	Retrospective cohort F	2016 (409)
Comment(s)	1° endpoint: Number of study interventions	Inclusion criteria:		Jongsma H, et al.
Comment(s)	& 95% CI)			Year Published
	(include P value; OR or RR;		Study Size	Author;
Summary/Conclusion	Primary Endpoint and Results	Patient Population	Study Type/Design;	Study Acronym;
inal Follow-Up-Section 10.	Evidence Table 52. Nonrandomized Trials, Observational Studies, and/or Registries for Longitudinal Foll	rials, Observational Studies, a	52. Nonrandomized Ti	Evidence Table (
ABI indicates ankle brachial index; CLI, critical limb ischemia; ePTFE, Polytetrafluoroethylene; IC, intermittent claudication; N/A, not applicable; NS, not significant; pt, patient; QoL, quality of life; and RCT, randomized controlled trial;	thylene; IC, intermittent claudication; N/A, not ap	mb ischemia; ePTFE, Polytetrafluoroe	ABI indicates ankle brachial index; CLI, critical lin of life; and RCT, randomized controlled trial;	ABI indicates ankle b of life; and RCT, rand
	Safety endpoint: N/A		<u>9126</u> . II- 130 pts	
SO NS.	patency results were also NS.		Cize: n=156 nte	
d secondary	(57% vs. 50%; NS) and secondary		Study type: Randomized	
routine group	intensive aroup vs. the routine aroup			
tency in the	Assisted primary cumulative eP incomposite graft patency in the		follow up.	
first 6mo		<u>Exclusion criteria</u> : N/A	graft patency as	
between visits was 6 wk for	56%; p<0.05)		femoropopliteal/crural	

Duplex surveillance with repair of	1° endpoint and results:	Inclusion criteria: Bypasses	Study type:	Stone PA, et al.
	88% for FT vs. 57% for FP (p = 0.04) -PPV was 95% FT vs. 65% FP (p = 0.04)	Exclusion criteria: No duplex surveillance, inadequate follow up (<3 mo)	bypasses)	
<ul> <li>Prosthetic grafts more prone to thrombosis.</li> </ul>	-20 raining grans were recorde. -Sensitivity of duplex correctly identifying failing graft:	surveillance protocol	Size: n= 66 pts (89 infrainguinal	<u>11665434</u>
<ul> <li>The surveillance and follow up management not shown to be</li> </ul>	<u>1° endpoint and results:</u> -22 thrombosed and 25 failing grafts	Inclusion criteria: Infringuinal prosthetic bypasses with Duplex	Study type: Observational	Calligaro KD, et al.
	95% CI: 1.9–19.2), use of warfarin (OR: 8.4; 95% CI: 2.1–34.5) and therapeutic warfarin (OR; 24.6%; CI: 5.7–106) to be independent predictors of patency.			
	<ul> <li>Distal anastomotic adjunct with below knee bypasses reduced graft thrombosis (35% with vs. 60% without) but no patency advantage.</li> <li>Multivariate analysis: low graft flow (OR: 6.1;</li> </ul>	Exclusion criteria: Cadaveric vein		
the thrombotic potential.	<ul> <li>At 3 y, treedom from limb loss was 75% and pt. survival was 75%.</li> </ul>	afterwards.		
<ul> <li>Early duplex scanning more important for diagnosing MGV and</li> </ul>	<ul> <li>NS differences noted between above knee and below knee grafts.</li> </ul>	bypass). Duplex surveillance at 1, 4	bypasses)	
detected stenoses.	respectively.	disabling claudication/and or popliteal	Size: n=121 pts (130	17920227
<ul> <li>Low graft flow endangered graft patency more frequently than development of duplex scan</li> </ul>	<ul> <li><u>1° endpoint and results:</u></li> <li>3y primary patency, assisted and secondary</li> <li>patency, results were 30% / 3%, and 50%</li> </ul>	Inclusion criteria: Pts with no usable saphenous veins. Lower limb ischemia (rest nain tissue loss	Study type: Observational	Brumberg RS, et al. 2007 (413)
	Jirans, ruenurying poliserioses in 46 bypasses. -29% of grafts diagnosed as failing by duplex scans were related to decrease in ABI >0.15.			
grafts and repair was successful	mitnin 6 mo of surveillance exam. GEV <45 cm/sec in 99 grafts resulted in arteriography in 75 cm/sec in 48 by proceeding and the supervisional sector of the supervisiona sector of the supervisiona sector of the supervisiona se	Exclusion criteria: None reported	ופעפוספט עפווו אומונס/	
Dupex surveillance identified graft-	• -2.1% of 280 grafts with GFV >45cm/sec failed	protocol	Size: n=292 pts (379	
be more reliable in the failing gratts than ABI	<ul> <li>Mean of 3.2 surveillance exams/ graft with a mean follow up was 21.5 mo.</li> </ul>	undergoing prospective surveillance	Observational	1990 (412) <u>2214034</u>
Duplex surveillance appeared to	1° endpoint and results:	Inclusion criteria: Infrainguinal	Study type:	Mills JL, et al.
	stenosed, 10 regressed spontaneously, 10 remained stable)			
	evision. -43 had stenosis (20 underwent revision, 2			
observational validation.	Results: -51 grafts didn't occlude and didn't require			

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Inclusion criteria. Dte with IC
Exclusion criteria: None reported
first y and every 6 mo thereafter.
at least 1 study documenting patent
Inclusion criteria: DU protocol with
Exclusion criteria: None reported
indefinitely after procedure.
mo. and then at 6 mo. intervals
angioplasty and stenting pts.
Inclusion criteria: Femoropoplitea
Exclusion criteria: None reported
aortolliac duplex surveillance protoco at <1 mo. 3 mo. and 6 mo. intervals
Iliac PTA and stents undergoing
-
Exclusion criteria: None reported
undergoing Duplex surveillance protocol

								111 consecutive pts	Size: n=142 limbs in	20853355	
								revascularization for CLI	Exclusion criteria: Pts with		
mm=23% vs. <100 mm=8%; p=0.041)	significantly so for intermediate lesions (100–200	and reintervention for longer lesions and	long-term patency with outpatient surveillance	<ul> <li>Compared to lesions &gt;100 mm, higher gain in</li> </ul>	secondary patency (HR=4.2; p=0.06)	<ul> <li>Lesions &gt;200 mm had higher trend in failed</li> </ul>	secondary patency (<5% incidence)	<ul> <li>Short and intermediate lesions had similar failed</li> </ul>	HR: 2.0; p=0.16 vs. >200 mm: HR=2.6; p=0.03)	had higher failed primary patency (100–200 mm;	
			artery lesions (>100 mm length).	initial treatment for longer temoral	long-term patency in pts receiving	intervention had a greater effect on	restenosis requiring repeat	<ul> <li>Outpatient surveillance for</li> </ul>	lesions (>100mm).	artery lesions was lower for long	

DU, duplex ultrasound; FP, femoropopliteal graft; FT, femorotibial graft; GFV, graft flow velocity; HVC, high-velocity criteria; IC, intermittent claudication; LCV; MGV; NPV, negative predictive value; NS, not significant; OR, odds ratio; PPV, positive predictive value; PSV, peak systolic velocities; PTA, percutaneous transluminal angioplasty; PTFE, polytetrafluoroethylene; pt, patient; PSV; u/s, ultrasound; ROC, receiver operating characteristic; and Vr, velocity ratio. nia;

## References

- ROSE GA. The diagnosis of ischaemic heart pain and intermittent claudication in field surveys. Bull World Health Organ. 1962;27:645-58
- 2 Leng GC, Fowkes FG. The Edinburgh Claudication Questionnaire: an improved version of the WHO/Rose Questionnaire for use in epidemiological surveys. J
- ω Criqui MH, Denenberg JO, Bird CE, et al. The correlation between symptoms and non-invasive test results in patients referred for peripheral arterial disease Clin Epidemiol. 1992;45:1101-9
- 4 Med. 1999;159:387-92. McDermott MM, Mehta S, Greenland P. Exertional leg symptoms other than intermittent claudication are common in peripheral arterial disease. Arch Intern testing. Vasc Med. 1996;1:65-71
- Ś McDermott MM, Greenland P, Liu K, et al. Leg symptoms in peripheral arterial disease: associated clinical characteristics and functional impairment. JAMA 2001;286:1599-606.
- Hirsch AT, Criqui MH, Treat-Jacobson D, et al. Peripheral arterial disease detection, awareness, and treatment in primary care. JAMA. 2001;286:1317-24
- .8 .7 .6 Khan NA, Rahim SA, Anand SS, et al. Does the clinical examination predict lower extremity peripheral arterial disease? JAMA. 2006;295:536-46
- Grøndal N, Sogaard R, Lindholt JS. Baseline prevalence of abdominal aortic aneurysm, peripheral arterial disease and hypertension in men aged 65-74 years from a population screening study (VIVA trial). Br J Surg. 2015;102:902-6.
- 9. Wassel CL, Loomba R, Ix JH, et al. Family history of peripheral artery disease is associated with prevalence and severity of peripheral artery disease: the San Diego Population Study. J Am Coll Cardiol. 2011;58:1386-92.
- 10. Clark CE, Taylor RS, Shore AC, et al. Association of a difference in systolic blood pressure between arms with vascular disease and mortality: a systematic review and meta-analysis. Lancet. 2012;379:905-14.
- 11. Singh S, Sethi A, Singh M, et al. Simultaneously measured inter-arm and inter-leg systolic blood pressure differences and cardiovascular risk stratification: a systemic review and meta-analysis. J Am Soc Hypertens. 2015;9:640-50.e12.
- 12 Shadman R, Criqui MH, Bundens WP, et al. Subclavian artery stenosis: prevalence, risk factors, and association with cardiovascular diseases. J Am Col Cardiol. 2004;44:618-23.
- 13. Cournot M, Boccalon H, Cambou JP, et al. Accuracy of the screening physical examination to identify subclinical atherosclerosis and peripheral arterial disease in asymptomatic subjects. J Vasc Surg. 2007;46:1215-21.
- 14. Armstrong DWJ, Tobin C, Matangi MF. The accuracy of the physical examination for the detection of lower extremity peripheral arterial disease. Can J Cardiol. 2010;26:e346-50.
- 15 Fowkes FG, Price JF, Stewart MC, et al. Aspirin for prevention of cardiovascular events in a general population screened for a low ankle brachial index: a randomized controlled trial. JAMA. 2010;303:841-8
- 16. Belch J, MacCuish A, Campbell I, et al. The Prevention of Progression of Arterial Disease and Diabetes (POPADAD) trial: factorial randomised placebo controlled trial of aspirin and antioxidants in patients with diabetes and asymptomatic peripheral arterial disease. BMJ 2008;337:a1840
- 17. McDermott MM, Liu K, Guralnik JM, et al. Home-based walking exercise intervention in peripheral artery disease: a randomized clinical trial. JAMA 2013;310:57-65.
- 19 18. Criqui MH, Vargas V, Denenberg JO, et al. Ethnicity and peripheral arterial disease: the San Diego Population Study. Circulation. 2005;112:2703-7.
- Examination Survey, 1999-2000. Circulation. 2004;110:738-43. Selvin E, Erlinger TP. Prevalence of and risk factors for peripheral arterial disease in the United States: results from the National Health and Nutrition
- 20. Guo X, Li J, Pang W, et al. Sensitivity and specificity of ankle-brachial index for detecting angiographic stenosis of peripheral arteries. Circ J. 2008;72:605-10

- 21. Aboyans V, Criqui MH, Abraham P, et al. Measurement and interpretation of the ankle-brachial index: a scientific statement from the American Heart Association. Circulation. 2012;126:2890-909.
- 22. Aboyans V, Ho E, Denenberg JO, et al. The association between elevated ankle systolic pressures and peripheral occlusive arterial disease in diabetic and nondiabetic subjects. J Vasc Surg. 2008;48:1197-203
- 23. disease. J Vasc Surg. 2006;44:531-6. Schröder F, Diehm N, Kareem S, et al. A modified calculation of ankle-brachial pressure index is far more sensitive in the detection of peripheral arterial
- 24. Premalatha G, Ravikumar R, Sanjay R, et al. Comparison of colour duplex ultrasound and ankle-brachial pressure index measurements in peripheral vascular disease in type 2 diabetic patients with foot infections. J Assoc Physicians India. 2002;50:1240-4.
- 25. Allen J, Oates CP, Henderson J, et al. Comparison of lower limb arterial assessments using color-duplex ultrasound and ankle/brachial pressure index measurements. Angiology. 1996;47:225-32.
- 26. 27. Fowkes FG, Murray GD, Butcher I, et al. Ankle brachial index combined with Framingham Risk Score to predict cardiovascular events and mortality: a meta-Lijmer JG, Hunink MG, van den Dungen JJ, et al. ROC analysis of noninvasive tests for peripheral arterial disease. Ultrasound Med Biol. 1996;22:391-8
- analysis. JAMA. 2008;300:197-208.
- 28. Fowkes FG, Murray GD, Butcher I, et al. Development and validation of an ankle brachial index risk model for the prediction of cardiovascular events. Eur J Prev Cardiol. 2014;21:310-20.
- 29. Diehm C, Allenberg JR, Pittrow D, et al. Mortality and vascular morbidity in older adults with asymptomatic versus symptomatic peripheral artery disease Circulation. 2009;120:2053-61.
- 30. Lin JS, Olson CM, Johnson ES, et al. The ankle-brachial index for peripheral artery disease screening and cardiovascular disease prediction among asymptomatic adults: a systematic evidence review for the U.S. Preventive Services Task Force. Ann Intern med. 2013;333-41.
- 31. Alahdab F, Wang AT, Elraiyah TA, et al. A systematic review for the screening for peripheral arterial disease in asymptomatic patients. J Vasc Surg 2015;61:42S-53S.
- 32. Body Composition (Health ABC) study. Vascular. 2014;22:142-8. Hiramoto JS, Katz R, Ix JH, et al. Sex differences in the prevalence and clinical outcomes of subclinical peripheral artery disease in the Health, Aging, and
- 33. . brachial index as a prognostic marker of cardiovascular disease. Ann Vasc Surg. 2010;24:985-93. Bundó M, Muñoz L, Pérez C, et al. Asymptomatic peripheral arterial disease in type 2 diabetes patients: a 10-year follow-up study of the utility of the ankle
- 34. Atherosclerosis. 2012;220:407-12. Tsivgoulis G, Bogiatzi C, Heliopoulos I, et al. Low ankle-brachial index predicts early risk of recurrent stroke in patients with acute cerebral ischemia.
- 35. disease. Am J Cardiol. 2012;110:197-202. Bouisset F, Bongard V, Ruidavets JB, et al. Prognostic usefulness of clinical and subclinical peripheral arterial disease in men with stable coronary heart
- 36. Sen S, Lynch DR, Kaltsas E, et al. Association of asymptomatic peripheral arterial disease with vascular events in patients with stroke or transient ischemic attack. Stroke. 2009;40:3472-7.
- 37. Ratanakorn D, Keandoungchun J, Tegeler CH. Prevalence and association between risk factors, stroke subtypes, and abnormal ankle brachial index in acute ischemic stroke. J Stroke Cerebrovasc Dis. 2012;21:498-503.
- 38. Cardiol. 2016;67:630-40. Ramos R, García-Gil M, Comas-Cufí M, et al. Statins for prevention of cardiovascular events in a low-risk population with low ankle brachial index. J Am Coll
- 39. Jiménez M, Dorado L, Hernández-Pérez M, et al. Ankle-brachial index in screening for asymptomatic carotid and intracranial atherosclerosis. Atherosclerosis. 2014;233:72-5.
- 40. McDermott MM, Fried L, Simonsick E, et al. Asymptomatic peripheral arterial disease is independently associated with impaired lower extremity functioning the Women's Health and Aging Study. Circulation. 2000;101:1007-12
- 41. McDermott MM, Liu K, Greenland P, et al. Functional decline in peripheral arterial disease: associations with the ankle brachial index and leg symptoms JAMA. 2004;292:453-61.
- 42. McDermott MM, Liu K, Ferrucci L, et al. Physical performance in peripheral arterial disease: a slower rate of decline in patients who walk more. Ann Intern med. 2006;144:10-20.
- 43. McDermott MM, Ferrucci L, Liu K, et al. Leg symptom categories and rates of mobility decline in peripheral arterial disease. J Am Geriatr Soc. 2010;58:1256-
- 44 McDermott MM, Applegate WB, Bonds DE, et al. Ankle brachial index values, leg symptoms, and functional performance among community-dwelling older men and women in the Lifestyle Interventions and Independence for Elders Study. J Am Heart Assoc. 2013;2:e000257
- 45 Niazi K, Khan TH, Easley KA. Diagnostic utility of the two methods of ankle brachial index in the detection of peripheral arterial disease of lower extremities Catheter Cardiovasc Interv. 2006;68:788-92.
- 46. Rutherford RB, Baker JD, Ernst C, et al. Recommended standards for reports dealing with lower extremity ischemia: revised version. J Vasc Surg 1997;26:517-3
- 47. stratification. Ann Vasc Surg. 2014;28:18-27. Eslahpazir BA, Allemang MT, Lakin RO, et al. Pulse volume recording does not enhance segmental pressure readings for peripheral arterial disease
- 48. 50. 51. 52. Ouriel K, McDonnell AE, Metz CE, et al. Critical evaluation of stress testing in the diagnosis of peripheral vascular disease. Surgery. 1982;91:686-93
  - Aerden D, Massaad D, von KK, et al. The ankle--brachial index and the diabetic foot: a troublesome marriage. Ann Vasc Surg. 2011;25:770-7
  - Park SC, Choi CY, Ha YI, et al. Utility of toe-brachial index for diagnosis of peripheral artery disease. Arch Plast Surg. 2012;39:227-31.
  - Weinberg I, Giri J, Calfon MA, et al. Anatomic correlates of supra-normal ankle brachial indices. Catheter Cardiovasc Interv. 2013;81:1025-30
- Suominen V, Rantanen T, Venermo M, et al. Prevalence and risk factors of PAD among patients with elevated ABI. Eur J Vasc Endovasc Surg. 2008;35:709-14
- Wagener JS, Hendricker C. Intra-subject variability of noninvasive oxygen measurements. Chest. 1987;92:1047-9
- 5<u>4</u>. Tsai FW, Tulsyan N, Jones DN, et al. Skin perfusion pressure of the foot is a good substitute for toe pressure in the assessment of limb ischemia. J Vasc Surg 2000;32:32-6.
- 55. noninvasive diagnostic methods. J Vasc Surg. 2008;47:318-23. Yamada T, Ohta T, Ishibashi H, et al. Clinical reliability and utility of skin perfusion pressure measurement in ischemic limbs—comparison with other
- 56. Bosanquet DC, Glasbey JC, Williams IM, et al. Systematic review and meta-analysis of direct versus indirect angiosomal revascularisation of infrapopliteal arteries. Eur J Vasc Endovasc Surg. 2014;48:88-97.
- Carter SA. Clinical measurement of systolic pressures in limbs with arterial occlusive disease. JAMA. 1969;207:1869-74
- 57. Carter SA, Tate RB. Value of toe pulse waves in addition to systolic pressures in the assessment of the severity of peripheral arterial disease and critical limb ischemia. J Vasc Surg. 1996;24:258-65.
- 59. Cardiovasc Surg (Torino). 1983;24:43-8. Ramsey DE, Manke DA, Sumner DS. Toe blood pressure. A valuable adjunct to ankle pressure measurement for assessing peripheral arterial disease. J
- 60. Surg. 2014;47:517-22. Biancari F, Juvonen T. Angiosome-targeted lower limb revascularization for ischemic foot wounds: systematic review and meta-analysis. Eur J Vasc Endovasc
- 61. Vincent DG, Salles-Cunha SX, Bernhard VM, et al. Noninvasive assessment of toe systolic pressures with special reference to diabetes mellitus. J Cardiovasc Surg (Torino). 1983;24:22-8.
- 62. Mahe G, Pollak AW, Liedl DA, et al. Discordant diagnosis of lower extremity peripheral artery disease using American Heart Association postexercise guidelines. Medicine (Baltimore ). 2015;94:e1277
- 63 Nicolaï SP, Viechtbauer W, Kruidenier LM, et al. Reliability of treadmill testing in peripheral arterial disease: a meta-regression analysis. J Vasc Surg 2009;50:322-9
- 64. Laing SP, Greenhalgh RM. Standard exercise test to assess peripheral arterial disease. Br Med J. 1980;280:13-6.
- 65 Raines JK, Darling RC, Buth J, et al. Vascular laboratory criteria for the management of peripheral vascular disease of the lower extremities. Surgery 1976;79:21-9.

- 66. Sumner DS, Strandness DE. The relationship between calf blood flow and ankle blood pressure in patients with intermittent claudication. Surgery. 1969;65:763-71
- 67. Castronuovo JJ, Adera HM, Smiell JM, et al. Skin perfusion pressure measurement is valuable in the diagnosis of critical limb ischemia. J Vasc Surg 1997;26:629-37.
- 68 Biotteau E, Mahe G, Rousseau P, et al. Transcutaneous oxygen pressure measurements in diabetic and non-diabetic patients clinically suspected of severe limb schemia: a matched paired retrospective analysis. Int Angiol. 2009;28:479-83
- 69. critical limb ischemia. Vasc Med. 2015;20:23-9. Bunte MC, Jacob J, Nudelman B, et al. Validation of the relationship between ankle-brachial and toe-brachial indices and infragenicular arterial patency in
- 20 Stein R, Hriljac I, Halperin JL, et al. Limitation of the resting ankle-brachial index in symptomatic patients with peripheral arterial disease. Vasc Med 2006;11:29-33
- 71. Shishehbor MH, Hammad TA, Zeller T, et al. An analysis of IN.PACT DEEP randomized trial on the limitations of the societal guidelines-recommended hemodynamic parameters to diagnose critical limb ischemia. J Vasc Surg. 2016;63:1311-7.
- 72. body magnetic resonance angiography in the elderly. Acta Radiol. 2008;49:143-9. Wikström J, Hansen T, Johansson L, et al. Ankle brachial index < 0.9 underestimates the prevalence of peripheral artery occlusive disease assessed with whole-
- 73. brachial index. J Vasc Surg. 2009;50:330-4. Wikström J, Hansen T, Johansson L, et al. Lower extremity artery stenosis distribution in an unselected elderly population and its relation to a reduced ankle-
- 74. Clairotte C, Retout S, Potier L, et al. Automated ankle-brachial pressure index measurement by clinical staff for peripheral arterial disease diagnosis in nondiabetic and diabetic patients. Diabetes Care. 2009;32:1231-6.
- 75. lower extremity arterial disease. J Magn Reson Imaging. 2013;37:1427-35. Burbelko M, Augsten M, Kalinowski MO, et al. Comparison of contrast-enhanced multi-station MR angiography and digital subtraction angiography of the
- 76. Shareghi S, Gopal A, Gul K, et al. Diagnostic accuracy of 64 multidetector computed tomographic angiography in peripheral vascular disease. Catheter Cardiovasc Interv. 2010;75:23-31.
- 77. de Vries SO, Hunink MG, Polak JF. Summary receiver operating characteristic curves as a technique for meta-analysis of the diagnostic performance of duplex ultrasonography in peripheral arterial disease. Acad Radiol. 1996;3:361-9.
- 78. Ota H, Takase K, Igarashi K, et al. MDCT compared with digital subtraction angiography for assessment of lower extremity arterial occlusive disease importance of reviewing cross-sectional images. AJR Am J Roentgenol. 2004;182:201-9
- 79. He C, Yang JG, Li YM, et al. Comparison of lower extremity atherosclerosis in diabetic and non-diabetic patients using multidetector computed tomography. BMC Cardiovasc Disord. 2014;14:125.
- 80. angiography. Catheter Cardiovasc Interv. 2013;82:E516-E521. Philip F, Shishehbor MH, Desai MY, et al. Characterization of internal pudendal artery atherosclerosis using aortography and multi-detector computed
- 81. Kayhan A, Palabiyik F, Serinsöz S, et al. Multidetector CT angiography versus arterial duplex USG in diagnosis of mild lower extremity peripheral arterial disease: is multidetector CT a valuable screening tool? Eur J Radiol. 2012;81:542-6.
- 82 Joshi SB, Mendoza DD, Steinberg DH, et al. Ultra-low-dose intra-arterial contrast injection for iliofemoral computed tomographic angiography. JACC
- 83. Mesurolle B, Qanadli SD, El HM, et al. Occlusive arterial disease of abdominal aorta and lower extremities: comparison of helical CT angiography with Cardiovasc Imaging. 2009;2:1404-11.
- 84. Romano M, Mainenti PP, Imbriaco M, et al. Multidetector row CT angiography of the abdominal aorta and lower extremities in patients with peripheral arterial occlusive disease: diagnostic accuracy and interobserver agreement. Eur J Radiol. 2004;50:303-8. transcatheter angiography. Clin Imaging. 2004;28:252-60.
- 28 Martin ML, Tay KH, Flak B, et al. Multidetector CT angiography of the aortoiliac system and lower extremities: a prospective comparison with digital subtraction angiography. AJR Am J Roentgenol. 2003;180:1085-91.

- 86. Andreucci M, Solomon R, Tasanarong A. Side effects of radiographic contrast media: pathogenesis, risk factors, and prevention. Biomed Res Int. 2014;2014:741018.
- 87. Stacul F, van der Molen AJ, Reimer P, et al. Contrast induced nephropathy: updated ESUR Contrast Media Safety Committee guidelines. Eur Radiol 2011;21:2527-41.
- 88 Meyer BC, Klein S, Krix M, et al. Comparison of a standard and a high-concentration contrast medium protocol for MDCT angiography of the lower limb arteries. Rofo. 2012;184:527-34.
- 89. diagnostic accuracy in comparison with standard DSA. Eur Radiol. 2006;16:137-46 Fraioli F, Catalano C, Napoli A, et al. Low-dose multidetector-row CT angiography of the infra-renal aorta and lower extremity vessels: image quality and
- 90. Met R, Bipat S, Legemate DA, et al. Diagnostic performance of computed tomography angiography in peripheral arterial disease: a systematic review and meta-analysis. JAMA. 2009;301:415-24.
- 91. Cardiovasc Med (Hagerstown). 2007;8:337-41. Favaretto E, Pili C, Amato A, et al. Analysis of agreement between Duplex ultrasound scanning and arteriography in patients with lower limb artery disease. J
- 92. Kau T, Eicher W, Reiterer C, et al. Dual-energy CT angiography in peripheral arterial occlusive disease-accuracy of maximum intensity projections in clinical routine and subgroup analysis. Eur Radiol. 2011;21:1677-86.
- 93. controlled trials. BMC Med Imaging. 2011;11:12. McCullough PA, Capasso P. Patient discomfort associated with the use of intra-arterial iodinated contrast media: a meta-analysis of comparative randomized
- 94. Sultan S, Chua BY, Hamada N, et al. Preoperative vascular screening in the presence of aortic, carotid and peripheral pathology for patients undergoing their first arterial intervention: 18 month follow-up. Int Angiol. 2013;32:281-90.
- 96. 95. Giugliano G, Laurenzano E, Rengo C, et al. Abdominal aortic aneurysm in patients affected by intermittent claudication: prevalence and clinical predictors comparing the yield between patients with manifest atherosclerosis and patients with risk factors for atherosclerosis only. J Vasc Surg. 2003;37:1226-33 Kurvers HA, van der Graaf Y, Blankensteijn JD, et al. Screening for asymptomatic internal carotid artery stenosis and aneurysm of the abdominal aorta:
- 97. Barba A, Estallo L, Rodríguez L, et al. Detection of abdominal aortic aneurysm in patients with peripheral artery disease. Eur J Vasc Endovasc Surg BMC Surg. 2012;12(suppl 1):S17. 2005;30:504-8.
- 98. significant coronary artery disease. JACC Cardiovasc Interv. 2013;6:1303-13 Lee JY, Lee SW, Lee WS, et al. Prevalence and clinical implications of newly revealed, asymptomatic abnormal ankle-brachial index in patients with
- 99 Moyer VA. Screening for peripheral artery disease and cardiovascular disease risk assessment with the ankle-brachial index in adults: U.S. Preventive Services Task Force recommendation statement. Ann Intern Med. 2013;159:342-8.
- 100. McFalls EO, Ward HB, Moritz TE, et al. Coronary-artery revascularization before elective major vascular surgery. N Engl J Med. 2004;351:2795-804
- 101 Olin JW, Melia M, Young JR, et al. Prevalence of atherosclerotic renal artery stenosis in patients with atherosclerosis elsewhere. Am J Med. 1990;88:46N-51N
- 102. Leertouwer TC, Pattynama PM, van den Berg-Huysmans A. Incidental renal artery stenosis in peripheral vascular disease: a case for treatment? Kidney Int. 2001;59:1480-3.
- 103. 104 Catalano M, Born G, Peto R. Prevention of serious vascular events by aspirin amongst patients with peripheral arterial disease: randomized, double-blind trial Hansen KJ, Edwards MS, Craven TE, et al. Prevalence of renovascular disease in the elderly: a population-based study. J Vasc Surg. 2002;36:443-51
- 105. Horrocks M, Horrocks EH, Murphy P, et al. The effects of platelet inhibitors on platelet uptake and restenosis after femoral angioplasty. Int Angiol J Intern Med. 2007;261:276-84.
- 106. Minar E, Ahmadi A, Koppensteiner R, et al. Comparison of effects of high-dose and low-dose aspirin on restenosis after femoropopliteal percutaneous [997;16:101-6.
- 107. CAPRIE Steering Committee. A randomised, blinded, trial of clopidogrel versus aspirin in patients at risk of ischaemic events (CAPRIE). Lancet 1996;348:1329-39. transluminal angioplasty. Circulation. 1995;91:2167-73

- 108. Cacoub PP, Bhatt DL, Steg PG, et al. Patients with peripheral arterial disease in the CHARISMA trial. Eur Heart J. 2009;30:192-201
- 109. Bhatt DL, Flather MD, Hacke W, et al. Patients with prior myocardial infarction, stroke, or symptomatic peripheral arterial disease in the CHARISMA trial. J Am Coll Cardiol. 2007;49:1982-8.
- 110. Circulation. 2010;121:2575-83. Berger PB, Bhatt DL, Fuster V, et al. Bleeding complications with dual antiplatelet therapy among patients with stable vascular disease or risk factors for vascular disease: results from the Clopidogrel for High Atherothrombotic Risk and Ischemic Stabilization, Management, and Avoidance (CHARISMA) trial
- 111. Cassar K, Ford I, Greaves M, et al. Randomized clinical trial of the antiplatelet effects of aspirin-clopidogrel combination versus aspirin alone after lower limb angioplasty. Br J Surg. 2005;92:159-65.
- 112. Belch JJ, Dormandy J, CASPAR Writing CommitteeBiasi GM, et al. Results of the randomized, placebo-controlled Clopidogrel and Acetylsalicylic Acid in Bypass Surgery for Peripheral Arterial Disease (CASPAR) trial. J Vasc Surg. 2010;52:825-33.
- 113. Tepe G, Bantleon R, Brechtel K, et al. Management of peripheral arterial interventions with mono or dual antiplatelet therapy-the MIRROR study: a randomised and double-blinded clinical trial. Eur Radiol. 2012;22:1998-2006.
- 114. 115. Strobl FF, Brechtel K, Schmehl J, et al. Twelve-month results of a randomized trial comparing mono with dual antiplatelet therapy in endovascularly treated Bonaca MP, Scirica BM, Creager MA, et al. Vorapaxar in patients with peripheral artery disease: results from TRA2°P-TIMI 50. Circulation. 2013;127:1522-9
- 116. Collaborative overview of randomised trials of antiplatelet therapy--II: Maintenance of vascular graft or arterial patency by antiplatelet therapy. Antiplatelet patients with peripheral artery disease. J Endovasc Ther. 2013;20:699-706. Trialists' Collaboration. BMJ. 1994;308:159-68.
- 117. Antithrombotic Trialists' Collaboration. Collaborative meta-analysis of randomised trials of antiplatelet therapy for prevention of death, myocardial infarction, and stroke in high risk patients. BMJ. 2002;324:71-86.
- 118. Morrow DA, Braunwald E, Bonaca MP, et al. Vorapaxar in the secondary prevention of atherothrombotic events. N Engl J Med. 2012;366:1404-13
- 119. Bohula EA, Aylward PE, Bonaca MP, et al. Efficacy and safety of vorapaxar with and without a thienopyridine for secondary prevention in patients with
- 120. (TRA2°P-TIMI 50). Circulation. 2016;997-1005. Bonaca MP, Gutierrez JA, Creager MA, et al. Acute limb ischemia and outcomes with vorapaxar in patients with peripheral artery disease: results from the previous myocardial infarction and no history of stroke or transient ischemic attack: results from TRA 2°P-TIMI 50. Circulation. 2015;132:1871-9. Trial to Assess the Effects of Vorapaxar in Preventing Heart Attack and Stroke in patients With Atherosclerosis-Thrombolysis in Myocardial Infarction 50
- 121. Jones WS, Dolor RJ, Hasselblad V, et al. Comparative effectiveness of endovascular and surgical revascularization for patients with peripheral artery disease and critical limb ischemia: systematic review of revascularization in critical limb ischemia. Am Heart J. 2014;167:489-98.e7
- 122. Katsanos K, Spiliopoulos S, Saha P, et al. Comparative Efficacy and Safety of Different Antiplatelet Agents for Prevention of Major Cardiovascular Events and Leg Amputations in Patients with Peripheral Arterial Disease: A Systematic Review and Network Meta-Analysis. PLoS One. 2015;10:e0135692
- 123. Magnani G, Bonaca MP, Braunwald E, et al. Efficacy and safety of vorapaxar as approved for clinical use in the United States. J Am Heart Assoc 2015;4:e001505.
- 124 randomized trials. JAMA. 2009;301:1909-19. Berger JS, Krantz MJ, Kittelson JM, et al. Aspirin for the prevention of cardiovascular events in patients with peripheral artery disease: a meta-analysis of
- 125. symptomatic peripheral arterial disease. J Vasc Surg. 2015;62:157-65 Armstrong EJ, Anderson DR, Yeo KK, et al. Association of dual-antiplatelet therapy with reduced major adverse cardiovascular events in patients with
- 126. Heart Protection Study Collaborative Group. Randomized trial of the effects of cholesterol-lowering with simvastatin on peripheral vascular and other major vascular outcomes in 20,536 people with peripheral arterial disease and other high-risk conditions. J Vasc Surg. 2007,45:645-54
- 127. Mohler ER3, Hiatt WR, Creager MA. Cholesterol reduction with atorvastatin improves walking distance in patients with peripheral arterial disease Circulation. 2003;108:1481-6
- 128. Hiatt WR, Hirsch AT, Creager MA, et al. Effect of niacin ER/lovastatin on claudication symptoms in patients with peripheral artery disease. Vasc Med 2010;15:171-9.

- 129. Giri J, McDermott MM, Greenland P, et al. Statin use and functional decline in patients with and without peripheral arterial disease. J Am Coll Cardiol 2006;47:998-1004
- 130. West AM, Anderson JD, Meyer CH, et al. The effect of ezetimibe on peripheral arterial atherosclerosis depends upon statin use at baseline. Atherosclerosis 2011;218:156-62
- 131 Stoekenbroek RM, Boekholdt SM, Fayyad R, et al. High-dose atorvastatin is superior to moderate-dose simvastatin in preventing peripheral arterial disease Heart. 2015;101:356-62
- 133. 132. Kumbhani DJ, Steg PG, Cannon CP, et al. Statin therapy and long-term adverse limb outcomes in patients with peripheral artery disease: insights from the Aung PP, Maxwell HG, Jepson RG, et al. Lipid-lowering for peripheral arterial disease of the lower limb. Cochrane Database Syst Rev. 2007;CD000123
- 134 Circ Cardiovasc Interv. 2013;6:694-700. Vogel TR, Dombrovskiy VY, Galiñanes EL, et al. Preoperative statins and limb salvage after lower extremity revascularization in the Medicare population REACH registry. Eur Heart J. 2014;35:2864-72.
- 135. survival in patients with critical limb ischemia. J Am Coll Cardiol. 2014;63:682-90. Westin GG, Armstrong EJ, Bang H, et al. Association between statin medications and mortality, major adverse cardiovascular event, and amputation-free
- 136. Feringa HH, Karagiannis SE, van Waning VH, et al. The effect of intensified lipid-lowering therapy on long-term prognosis in patients with peripheral arterial disease. J Vasc Surg. 2007;45:936-43.
- 137. Ostergren J, Sleight P, Dagenais G, et al. Impact of ramipril in patients with evidence of clinical or subclinical peripheral arterial disease. Eur Heart J 2004;25:17-24
- 138. Yusuf S, Sleight P, Pogue J, et al. Effects of an angiotensin-converting-enzyme inhibitor, ramipril, on cardiovascular events in high-risk patients. The Heart Outcomes Prevention Evaluation Study Investigators. N Engl J Med. 2000;342:145-53.
- 139. Yusuf S, Teo KK, Pogue J, et al. Telmisartan, rampril, or both in patients at high risk for vascular events. N Engl J Med. 2008;358:1547-59
- 140. Bavry AA, Anderson RD, Gong Y, et al. Outcomes Among hypertensive patients with concomitant peripheral and coronary artery disease: findings from the INternational VErapamil-SR/Trandolapril STudy. Hypertension. 2010;55:48-53
- 141. of findings from the VALUE trial. J Hypertens. 2006;24:2163-8. Zanchetti A, Julius S, Kjeldsen S, et al. Outcomes in subgroups of hypertensive patients treated with regimens based on valsartan and amlodipine: an analysis
- 142. Diehm C, Pittrow D, Lawall H. Effect of nebivolol vs. hydrochlorothiazide on the walking capacity in hypertensive patients with intermittent claudication. J Hypertens. 2011;29:1448-56.
- 143. Espinola-Klein C, Weisser G, Jagodzinski A, et al. b-Blockers in patients with intermittent claudication and arterial hypertension: results from the nebivolol or metoprolol in arterial occlusive disease trial. Hypertension. 2011;58:148-54.
- 144. Paravastu SC, Mendonca DA, Da Silva A. Beta blockers for peripheral arterial disease. Cochrane Database Syst Rev. 2013;CD005508
- 145. ALLHAT Officers and Coordinators for the ALLHAT Collaborative Research Group. Major outcomes in high-risk hypertensive patients randomized to
- angiotensin-converting enzyme inhibitor or calcium channel blocker vs diuretic: The Antihypertensive and Lipid-Lowering Treatment to Prevent Heart Attack Trial (ALLHAT). JAMA. 2002;288:2981-97
- 146. Feringa HH, van Waning VH, Bax JJ, et al. Cardioprotective medication is associated with improved survival in patients with peripheral arterial disease. J Am Coll Cardiol. 2006;47:1182-7
- 147. Sleight P. The HOPE Study (Heart Outcomes Prevention Evaluation). J Renin Angiotensin Aldosterone Syst. 2000; 1:18-20
- 148. Rigotti NA, Regan S, Levy DE, et al. Sustained care intervention and postdischarge smoking cessation among hospitalized adults: a randomized clinical trial JAMA. 2014;312:719-28.
- 149. Rigotti NA, Pipe AL, Benowitz NL, et al. Efficacy and safety of varenicline for smoking cessation in patients with cardiovascular disease: a randomized trial Circulation. 2010;121:221-9.
- 150. Hennrikus D, Joseph AM, Lando HA, et al. Effectiveness of a smoking cessation program for peripheral artery disease patients: a randomized controlled trial. J Am Coll Cardiol. 2010;56:2105-12

- 151. Tonstad S, Farsang C, Klaene G, et al. Bupropion SR for smoking cessation in smokers with cardiovascular disease: a multicentre, randomised study. Eur Heart J. 2003;24:946-55
- 152. 153. Stead LF, Buitrago D, Preciado N, et al. Physician advice for smoking cessation. Cochrane Database Syst Rev. 2013;CD000165
- analysis. BMJ. 2012;344:e2856. Prochaska JJ, Hilton JF. Risk of cardiovascular serious adverse events associated with varenicline use for tobacco cessation: systematic review and meta-
- 154. Mills EJ, Thorlund K, Eapen S, et al. Cardiovascular events associated with smoking cessation pharmacotherapies: a network meta-analysis. Circulation 2014;129:28-41.
- 155. Clair C, Rigotti NA, Porneala B, et al. Association of smoking cessation and weight change with cardiovascular disease among adults with and without diabetes. JAMA. 2013;309:1014-21.
- 156. Hoel AW, Nolan BW, Goodney PP, et al. Variation in smoking cessation after vascular operations. J Vasc Surg. 2013;57:1338-44
- 157. Selvarajah S, Black JH3, Malas MB, et al. Preoperative smoking is associated with early graft failure after infrainguinal bypass surgery. J Vasc Surg 2014;59:1308-14.
- 158. Armstrong EJ, Wu J, Singh GD, et al. Smoking cessation is associated with decreased mortality and improved amputation-free survival among patients with symptomatic peripheral artery disease. J Vasc Surg. 2014;60:1565-71.
- 159. smokers. Atherosclerosis. 2013;229:273-6. Lu L, Mackay DF, Pell JP. Association between level of exposure to secondhand smoke and peripheral arterial disease: cross-sectional study of 5,686 never
- 160. Circulation. 2012;126:2177-83. Tan CE, Glantz SA. Association between smoke-free legislation and hospitalizations for cardiac, cerebrovascular, and respiratory diseases: a meta-analysis
- 161. Dormandy JA, Charbonnel B, Eckland DJ, et al. Secondary prevention of macrovascular events in patients with type 2 diabetes in the PROactive Study (PROspective pioglitAzone Clinical Trial In macroVascular Events): a randomised controlled trial. Lancet. 2005;366:1279-89
- 162. Singh S, Armstrong EJ, Sherif W, et al. Association of elevated fasting glucose with lower patency and increased major adverse limb events among patients with diabetes undergoing infrapopliteal balloon angioplasty. Vasc Med. 2014;19:307-14
- 163. angioplasty for critical limb ischemia. Diabetes Care. 2010;33:2538-42. Takahara M, Kaneto H, Iida O, et al. The influence of glycemic control on the prognosis of Japanese patients undergoing percutaneous transluminal
- 164. Resnick HE, Lindsay RS, McDermott MM, et al. Relationship of high and low ankle brachial index to all-cause and cardiovascular disease mortality: the Strong Heart Study. Circulation. 2004;109:733-9.
- 165. Anand S, Yusuf S, Xie C, et al. Oral anticoagulant and antiplatelet therapy and peripheral arterial disease. N Engl J Med. 2007;357:217-27
- 166. Efficacy of oral anticoagulants compared with aspirin after infrainguinal bypass surgery (The Dutch Bypass Oral Anticoagulants or Aspirin Study): a randomised trial. Lancet. 2000;355:346-51
- 167. Johnson WC, Williford WO, Department of Veterans Affairs Cooporative Study #362. Benefits, morbidity, and mortality associated with long-term administration of oral anticoagulant therapy to patients with peripheral arterial bypass procedures: a prospective randomized study. J Vasc Surg. 2002;35:413-
- 168. Sarac TP, Huber TS, Back MR, et al. Warfarin improves the outcome of infrainguinal vein bypass grafting at high risk for failure. J Vasc Surg. 1998;28:446-
- 169. Antonicelli R, Sardina M, Scotti A, et al. Randomized trial of the effects of low-dose calcium-heparin in patients with peripheral arterial disease and claudication. Italian CAP Study Group. Am J Med. 1999;107:234-9.
- 170. Alonso-Coello P, Bellmunt S, McGorrian C, et al. Antithrombotic therapy in peripheral artery disease: Antithrombotic Therapy and Prevention of Thrombosis. 9th ed: American College of Chest Physicians Evidence-Based Clinical Practice Guidelines. Chest. 2012;141:e669S-90S
- 171. Bedenis R, Lethaby A, Maxwell H, et al. Antiplatelet agents for preventing thrombosis after peripheral arterial bypass surgery. Cochrane Database Syst Rev. 2015;CD000535

- 172. Cosmi B, Conti E, Coccheri S. Anticoagulants (heparin, low molecular weight heparin and oral anticoagulants) for intermittent claudication. Cochrane Database Syst Rev. 2001;CD001999.
- 173. Bedenis R, Stewart M, Cleanthis M, et al. Cilostazol for intermittent claudication. Cochrane Database Syst Rev. 2014;CD003748
- 174. 175 Goldenberg NA, Krantz MJ, Hiatt WR. L-Carnitine plus cilostazol versus cilostazol alone for the treatment of claudication in patients with peripheral artery Dawson DL, Cutler BS, Hiatt WR, et al. A comparison of cilostazol and pentoxifylline for treating intermittent claudication. Am J Med. 2000; 109:523-30
- 176. Warner CJ, Greaves SW, Larson RJ, et al. Cilostazol is associated with improved outcomes after peripheral endovascular interventions. J Vasc Surg disease: a multicenter, randomized, double-blind, placebo-controlled trial. Vasc Med. 2012;17:145-54. 2014;59:1607-14.
- 177. lida O, Yokoi H, Soga Y, et al. Cilostazol reduces angiographic restenosis after endovascular therapy for femoropopliteal lesions in the Sufficient Treatment of Peripheral Intervention by Cilostazol study. Circulation. 2013;127:2307-15
- 178. Salhiyyah K, Senanayake E, Abdel-Hadi M, et al. Pentoxifylline for intermittent claudication. Cochrane Database Syst Rev. 2012;1:CD005262
- 179. Villarruz MV, Dans A, Tan F. Chelation therapy for atherosclerotic cardiovascular disease. Cochrane Database Syst Rev. 2002;CD002785
- 180. Khandanpour N, Loke YK, Meyer FJ, et al. Homocysteine and peripheral arterial disease: systematic review and meta-analysis. Eur J Vasc Endovasc Surg 2009;38:316-22.
- 181. 182. Lonn E, Yusuf S, Arnold MJ, et al. Homocysteine lowering with folic acid and B vitamins in vascular disease. N Engl J Med. 2006;354:1567-77
- B12 in high-risk patients: the Heart Outcomes Prevention Evaluation (HOPE)-2 trial. Can J Cardiol. 2006;22:47-53. Lonn E, Held C, Arnold JM, et al. Rationale, design and baseline characteristics of a large, simple, randomized trial of combined folic acid and vitamins B6 and
- 183. Study. Eur Heart J. 2004;25:25-31. Gurfinkel EP, Leo de la Fuente R, Mendiz O, et al. Flu vaccination in acute coronary syndromes and planned percutaneous coronary interventions (FLUVACS)
- 184. Ciszewski A, Bilinska ZT, Brydak LB, et al. Influenza vaccination in secondary prevention from coronary ischaemic events in coronary artery disease FLUCAD study. Eur Heart J. 2008;29:1350-8.
- 185. Davis MM, Taubert K, Benin AL, et al. Influenza vaccination as secondary prevention for cardiovascular disease: a science advisory from the American Heart Association/American College of Cardiology. J Am Coll Cardiol. 2006;48:1498-502.
- 186. Murphy TP, Cutlip DE, Regensteiner JG, et al. Supervised exercise, stent revascularization, or medical therapy for claudication due to aortoiliac peripheral artery disease: the CLEVER study. J Am Coll Cardiol. 2015;65:999-1009.
- 187. six-month outcomes from the claudication: exercise versus endoluminal revascularization (CLEVER) study. Circulation. 2012;125:130-Murphy TP, Cutlip DE, Regensteiner JG, et al. Supervised exercise versus primary stenting for claudication resulting from aortoiliac peripheral artery disease:
- 188. McDermott MM, Guralnik JM, Criqui MH, et al. Home-based walking exercise in peripheral artery disease: 12-month follow-up of the GOALS randomized trial. J Am Heart Assoc. 2014;3:e000711.
- 189. arterial disease: a randomized controlled trial. Diabetes Care. 2011;34:2174-9. Collins TC, Lunos S, Carlson T, et al. Effects of a home-based walking intervention on mobility and quality of life in people with diabetes and peripheral
- 190. Gardner AW, Parker DE, Montgomery PS, et al. Efficacy of quantified home-based exercise and supervised exercise in patients with intermittent claudication. a randomized controlled trial. Circulation. 2011;123:491-8.
- 191 Saxton JM, Zwierska I, Blagojevic M, et al. Upper- versus lower-limb aerobic exercise training on health-related quality of life in patients with symptomatic peripheral arterial disease. J Vasc Surg. 2011;53:1265-73
- 192. claudication. Vasc Med. 2009;14:203-13. Treat-Jacobson D, Bronas UG, Leon AS. Efficacy of arm-ergometry versus treadmill exercise training to improve walking distance in patients with
- 193. Mika P, Konik A, Januszek R, et al. Comparison of two treadmill training programs on walking ability and endothelial function in intermittent claudication. Int 1 Cardiol. 2013;168:838-42
- 194. Fakhry F, Rouwet EV, den Hoed PT, et al. Long-term clinical effectiveness of supervised exercise therapy versus endovascular revascularization for intermittent claudication from a randomized clinical trial. Br J Surg. 2013;100:1164-71.

- 195. Mazari FA, Gulati S, Rahman MN, et al. Early outcomes from a randomized, controlled trial of supervised exercise, angioplasty, and combined therapy in intermittent claudication. Ann Vasc Surg. 2010;24:69-79.
- 196 Fakhry F, Spronk S, van der Laan L, et al. Endovascular revascularization and supervised exercise for peripheral artery disease and intermittent claudication: a randomized clinical trial. JAMA. 2015;314:1936-44
- 197 Guidon M, McGee H. One-year effect of a supervised exercise programme on functional capacity and quality of life in peripheral arterial disease. Disabil Rehabil. 2013;35:397-404.
- 198 Gardner AW, Parker DE, Montgomery PS, et al. Step-monitored home exercise improves ambulation, vascular function, and inflammation in symptomatic patients with peripheral artery disease: a randomized controlled trial. J Am Heart Assoc. 2014;3:e001107
- 199 Langbein WE, Collins EG, Orebaugh C, et al. Increasing exercise tolerance of persons limited by claudication pain using polestriding. J Vasc Surg 2002;35:887-93.
- 200. with intermittent claudication. J Vasc Surg. 2000;31:662-9 Walker RD, Nawaz S, Wilkinson CH, et al. Influence of upper- and lower-limb exercise training on cardiovascular function and walking distances in patients
- 201. Pilz M, Kandioler-Honetz E, Wenkstetten-Holub A, et al. Evaluation of 6- and 12-month supervised exercise training on strength and endurance parameters in patients with peripheral arterial disease. Wien Klin Wochenschr. 2014;126:383-9.
- 202. Mays RJ, Rogers RK, Hiatt WR, et al. Community walking programs for treatment of peripheral artery disease. J Vasc Surg. 2013;58:1678-87
- 203. Crane M, Werber B. Critical pathway approach to diabetic pedal infections in a multidisciplinary setting. J Foot Ankle Surg. 1999;38:30-3.
- 204 approach? Diabet Med. 1995;12:770-6. Larsson J, Apelqvist J, Agardh CD, et al. Decreasing incidence of major amputation in diabetic patients: a consequence of a multidisciplinary foot care team
- 205. Armstrong DG, Bharara M, White M, et al. The impact and outcomes of establishing an integrated interdisciplinary surgical team to care for the diabetic foot Diabetes Metab Res Rev. 2012;28:514-8.
- 206 Chung J, Modrall JG, Ahn C, et al. Multidisciplinary care improves amputation-free survival in patients with chronic critical limb ischemia. J Vasc Surg 2015;61:162-9
- 207 Canavan RJ, Unwin NC, Kelly WF, et al. Diabetes- and nondiabetes-related lower extremity amputation incidence before and after the introduction of better organized diabetes foot care: continuous longitudinal monitoring using a standard method. Diabetes Care. 2008;31:459-63
- 208. Williams DT, Majeed MU, Shingler G, et al. A diabetic foot service established by a department of vascular surgery: an observational study. Ann Vasc Surg 2012;26:700-6
- 209 Diabetes Care. 2005;28:248-53. Driver VR, Madsen J, Goodman RA. Reducing amputation rates in patients with diabetes at a military medical center: the limb preservation service model
- 211. 210. Vartanian SM, Robinson KD, Ofili K, et al. Outcomes of neuroischemic wounds treated by a multidisciplinary amputation prevention service. Ann Vasc Surg. Wrobel JS, Charns MP, Diehr P, et al. The relationship between provider coordination and diabetes-related foot outcomes. Diabetes Care. 2003;26:3042-7
- 2015;29:534-42.
- 212. 213. Gardner SE, Hillis SL, Frantz RA. Clinical signs of infection in diabetic foot ulcers with high microbial load. Biol Res Nurs. 2009;11:119-28
- Lipsky BA, Berendt AR, Cornia PB, et al. 2012 Infectious Diseases Society of America clinical practice guideline for the diagnosis and treatment of diabetic foot infections. Clin Infect Dis. 2012;54:e132-73.
- 214. Pickwell K, Siersma V, Kars M, et al. Predictors of lower-extremity amputation in patients with an infected diabetic foot ulcer. Diabetes Care. 2015;38:852-7
- 215. analysis. Clin Infect Dis. 2008;47:519-27 Dinh MT, Abad CL, Safdar N. Diagnostic accuracy of the physical examination and imaging tests for osteomyelitis underlying diabetic foot ulcers: meta-
- 216. Prompers L, Schaper N, Apelqvist J, et al. Prediction of outcome in individuals with diabetic foot ulcers: focus on the differences between individuals with and without peripheral arterial disease. The EURODIALE Study. Diabetologia. 2008;51:747-55
- 217. Rogers LC, Andros G, Caporusso J, et al. Toe and flow: essential components and structure of the amputation prevention team. J Vasc Surg. 2010;52:23S-7S

- 218. Society for Vascular Surgery and the American Podiatric Medical Association. J Vasc Surg. 2010;51:1504-6. Sumpio BE, Armstrong DG, Lavery LA, et al. The role of interdisciplinary team approach in the management of the diabetic foot: a joint statement from the
- 220. 219. Wrobel JS, Robbins JM, Charns MP, et al. Diabetes-related foot care at 10 Veterans Affairs medical centers: must do's associated with successful Fitzgerald RH, Mills JL, Joseph W, et al. The diabetic rapid response acute foot team: 7 essential skills for targeted limb salvage. Eplasty. 2009;9:e15
- 221. Tetteroo E, van der Graaf Y, Bosch JL, et al. Randomised comparison of primary stent placement versus primary angioplasty followed by selective stent microsystems. Jt Comm J Qual Patient Saf. 2006;32:206-13. placement in patients with iliac-artery occlusive disease. Dutch Iliac Stent Trial Study Group. Lancet. 1998;351:1153-9
- 222. iliac artery disease: The Dutch Iliac Stent Trial Study. Radiology. 2004;232:491-8. Klein WM, van der Graaf Y, Seegers J, et al. Long-term cardiovascular morbidity, mortality, and reintervention after endovascular treatment in patients with
- 223. Bosch JL, Hunink MG. Meta-analysis of the results of percutaneous transluminal angioplasty and stent placement for aortoiliac occlusive disease. Radiology 1997;204:87-96.
- 224. Kashyap VS, Pavkov ML, Bena JF, et al. The management of severe aortoiliac occlusive disease: endovascular therapy rivals open reconstruction. J Vasc Surg 2008;48:1451-7, 1457.
- 225 stenting. Circulation. 2007;115:2745-9. Schillinger M, Sabeti S, Dick P, et al. Sustained benefit at 2 years of primary femoropopliteal stenting compared with balloon angioplasty with optional
- 226 up to 10 cm in length: the Femoral Artery Stenting Trial (FAST). Circulation. 2007;116:285-92. Krankenberg H, Schlüter M, Steinkamp HJ, et al. Nitinol stent implantation versus percutaneous transluminal angioplasty in superficial femoral artery lesions
- 227. Laird JR, Katzen BT, Scheinert D, et al. Nitinol stent implantation versus balloon angioplasty for lesions in the superficial femoral artery and proximal popliteal artery: twelve-month results from the RESILIENT randomized trial. Circ Cardiovasc Interv. 2010;3:267-76.
- 228 Dick P, Wallner H, Sabeti S, et al. Balloon angioplasty versus stenting with nitinol stents in intermediate length superficial femoral artery lesions. Catheter Cardiovasc Interv. 2009;74:1090-5.
- 229 Tepe G, Laird J, Schneider P, et al. Drug-coated balloon versus standard percutaneous transluminal angioplasty for the treatment of superficial femoral and popliteal peripheral artery disease: 12-month results from the IN.PACT SFA randomized trial. Circulation. 2015;131:495-502
- 230. eluting balloon in peripheral intervention for the superficial femoral artery). JACC Cardiovasc Interv. 2013;6:1295-302. Listro F, Grotti S, Porto I, et al. Drug-eluting balloon in peripheral intervention for the superficial femoral artery: the DEBATE-SFA randomized trial (drug
- 231 Scheinert D, Duda S, Zeller T, et al. The LEVANT I (Lutonix paclitaxel-coated balloon for the prevention of femoropopliteal restenosis) trial for Interv. 2014;7:10-9. femoropopliteal revascularization: first-in-human randomized trial of low-dose drug-coated balloon versus uncoated balloon angioplasty. JACC Cardiovasc
- 232 Werk M, Albrecht T, Meyer DR, et al. Paclitaxel-coated balloons reduce restenosis after femoro-popliteal angioplasty: evidence from the randomized PACIFIER trial. Circ Cardiovasc Interv. 2012;5:831-40.
- 233. Lammer J, Zeller T, Hausegger KA, et al. Heparin-bonded covered stents versus bare-metal stents for complex femoropopliteal artery lesions: the randomized femoral artery occlusive disease). J Am Coll Cardiol. 2013;62:1320-7 VIASTAR trial (Viabahn endoprosthesis with PROPATEN bioactive surface [VIA] versus bare nitinol stent in the treatment of long lesions in superficial
- 234. Geraghty PJ, Mewissen MW, Jaff MR, et al. Three-year results of the VIBRANT trial of VIABAHN endoprosthesis versus bare nitinol stent implantation for complex superficial femoral artery occlusive disease. J Vasc Surg. 2013;58:386-95
- 235. Saxon RR, Dake MD, Volgelzang RL, et al. Randomized, multicenter study comparing expanded polytetrafluoroethylene-covered endoprosthesis placement with percutaneous transluminal angioplasty in the treatment of superficial femoral artery occlusive disease. J Vasc Interv Radiol. 2008;19:823-37
- 236. superficial femoral arterial occlusive disease. J Vasc Surg. 2007;45:10-6 Kedora J, Hohmann S, Garrett W, et al. Randomized comparison of percutaneous Viabahn stent grafts vs prosthetic femoral-popliteal bypass in the treatment of
- 237. Dake MD, Ansel GM, Jaff MR, et al. Paclitaxel-eluting stents show superiority to balloon angioplasty and bare metal stents in femoropopliteal disease: twelvemonth Zilver PTX randomized study results. Circ Cardiovasc Interv. 2011;4:495-504

- 238. Dake MD, Ansel GM, Jaff MR, et al. Durable clinical effectiveness with paclitaxel-eluting stents in the femoropopliteal artery: 5-year results of the Zilver PTX Randomized Trial. Circulation. 2016;133:1472-83.
- 239 term results from the SIROCCO trial. J Endovasc Ther. 2006;13:701-10. Duda SH, Bosiers M, Lammer J, et al. Drug-eluting and bare nitinol stents for the treatment of atherosclerotic lesions in the superficial femoral artery: long-
- 240. Tepe G, Zeller T, Albrecht T, et al. Local delivery of paclitaxel to inhibit restenosis during angioplasty of the leg. N Engl J Med. 2008;358:689-99
- 241 Dippel EJ, Makam P, Kovach R, et al. Randomized controlled study of excimer laser atherectomy for treatment of femoropopuliteal in-stent restenosis: initial results from the EXCITE ISR trial (EXCImer Laser Randomized Controlled Study for Treatment of FemoropopliTEal In-Stent Restenosis). JACC Cardiovasc Interv. 2015;8:92-101.
- 242. arterial segments: the COBRA trial. J Am Coll Cardiol. 2012;60:1352-9 Banerjee S, Das TS, Abu-Fadel MS, et al. Pilot trial of cryoplasty or conventional balloon post-dilation of nitinol stents for revascularization of peripheral
- 243. Endovasc Surg. 1996;12:167-72. Whyman MR, Fowkes FG, Kerracher EM, et al. Randomised controlled trial of percutaneous transluminal angioplasty for intermittent claudication. Eur J Vasco
- 244. trial. J Vasc Surg. 1997;26:551-7. Whyman MR, Fowkes FG, Kerracher EM, et al. Is intermittent claudication improved by percutaneous transluminal angioplasty? A randomized controlled
- 245. Perkins JM, Collin J, Creasy TS, et al. Reprinted article "Exercise training versus angioplasty for stable claudication. Long and medium term results of a prospective, randomised trial". Eur J Vasc Endovasc Surg. 2011;42(suppl 1):S41-5.
- 246. exercise training-randomized controlled trial. Radiology. 2009;250:586-95. Spronk S, Bosch JL, den Hoed PT, et al. Intermittent claudication: clinical effectiveness of endovascular revascularization versus supervised hospital-based
- 247. Spronk S, Bosch JL, den Hoed PT, et al. Cost-effectiveness of endovascular revascularization compared to supervised hospital-based exercise training in patients with intermittent claudication: a randomized controlled trial. J Vasc Surg. 2008;48:1472-80
- 249. 248. Gelin J, Jivegård L, Taft C, et al. Treatment efficacy of intermittent claudication by surgical intervention, supervised physical exercise training compared to no Taft C, Karlsson J, Gelin J, et al. Treatment efficacy of intermittent claudication by invasive therapy, supervised physical exercise training compared to no treatment in unselected randomised patients, I: one year results of functional and physiological improvements. Eur J Vasc Endovasc Surg. 2001;22:107-13.
- 250. Hobbs SD, Marshall T, Fegan C, et al. The constitutive procoagulant and hypofibrinolytic state in patients with intermittent claudication due to infrainguinal treatment in unselected randomised patients, II: one-year results of health-related quality of life. Eur J Vasc Endovasc Surg. 2001;22:114-23
- 251. Nylaende M, Abdelnoor M, Stranden E, et al. The Oslo Balloon Angioplasty versus Conservative Treatment study (OBACT)—the 2-years results of a single disease significantly improves with percutaneous transluminal balloon angioplasty. J Vasc Surg. 2006;43:40-6.
- centre, prospective, randomised study in patients with intermittent claudication. Eur J Vasc Endovasc Surg. 2007;33:3-12.
- 252. Greenhalgh RM, Belch JJ, Brown LC, et al. The adjuvant benefit of angioplasty in patients with mild to moderate intermittent claudication (MIMIC) managed arterial disease. Eur J Vasc Endovasc Surg. 2008;36:680-8 by supervised exercise, smoking cessation advice and best medical therapy: results from two randomised trials for stenotic femoropopliteal and aortoiliac
- 253. Kruidenier LM, Nicolai SP, Rouwet EV, et al. Additional supervised exercise therapy after a percutaneous vascular intervention for peripheral arterial disease a randomized clinical trial. J Vasc Interv Radiol. 2011;22:961-8.
- 254. intermittent claudication due to femoropopliteal arterial disease. Br J Surg. 2012;99:39-48. Mazari FA, Khan JA, Carradice D, et al. Randomized clinical trial of percutaneous transluminal angioplasty, supervised exercise and combined treatment for
- 255. treatment for intermittent claudication—a prospective randomised trial. Eur J Vasc Endovasc Surg. 2011;42:220-7. Nordanstig J, Gelin J, Hensäter M, et al. Walking performance and health-related quality of life after surgical or endovascular invasive versus non-invasive
- 256. Nordanstig J, Taft C, Hensäter M, et al. Improved quality of life after 1 year with an invasive versus a noninvasive treatment strategy in claudicants: one-year results of the Invasive Revascularization or Not in Intermittent Claudication (IRONIC) Trial. Circulation. 2014;130:939-47
- 257. Malgor RD, Alahdab F, Elraiyah TA, et al. A systematic review of treatment of intermittent claudication in the lower extremities. J Vasc Surg. 2015;61:54S-

- 258. Vemulapalli S, Dolor RJ, Hasselblad V, et al. Comparative effectiveness of medical therapy, supervised exercise, and revascularization for patients with intermittent claudication: a network meta-analysis. Clin Cardiol. 2015;38:378-86.
- 259. 260. McPhail IR, Spittell PC, Weston SA, et al. Intermittent claudication: an objective office-based assessment. J Am Coll Cardiol. 2001;37:1381-5
- Schulte KL, Pilger E, Schellong S, et al. Primary self-expanding nitinol stenting vs balloon angioplasty with optional bailout stenting for the treatment of infrapopliteal artery disease in patients with severe intermittent claudication or critical limb ischemia (EXPAND Study). J Endovasc Ther. 2015;22:690-7
- 261. 262. Sakamoto Y, Hirano K, Iida O, et al. Five-year outcomes of self-expanding nitinol stent implantation for chronic total occlusion of the superficial femoral and Scheinert D, Scheinert S, Sax J, et al. Prevalence and clinical impact of stent fractures after femoropopliteal stenting. J Am Coll Cardiol. 2005;45:312-5.
- 263 Feinglass J, McCarthy WJ, Slavensky R, et al. Functional status and walking ability after lower extremity bypass grafting or angioplasty for intermittent proximal popliteal artery. Catheter Cardiovasc Interv. 2013;82:E251-E256.
- 264 Giugliano G, Di SL, Perrino C, et al. Effects of successful percutaneous lower extremity revascularization on cardiovascular outcome in patients with claudication: results from a prospective outcomes study. J Vasc Surg. 2000;31:93-103. peripheral arterial disease. Int J Cardiol. 2013;167:2566-71.
- 265. or open surgery for symptomatic lower limb atherosclerotic disease. Eur J Cardiovasc Nurs. 2008;7:247-56. Koivunen K, Lukkarinen H. One-year prospective health-related quality-of-life outcomes in patients treated with conservative method, endovascular treatment
- 266. Pell JP, Lee AJ. Impact of angioplasty and arterial reconstructive surgery on the quality of life of claudicants. The Scottish Vascular Audit Group. Scott Med J 997;42:47-8.
- 267 arterial disease. J Vasc Surg. 2006;44:296-302. Kalbaugh CA, Taylor SM, Blackhurst DW, et al. One-year prospective quality-of-life outcomes in patients treated with angioplasty for symptomatic peripheral
- 268 Sachs T, Pomposelli F, Hamdan A, et al. Trends in the national outcomes and costs for claudication and limb threatening ischemia: angioplasty vs bypass graft J Vasc Surg. 2011;54:1021-31.
- 269 Shammas NW, Shammas GA, Dippel EJ, et al. Predictors of distal embolization in peripheral percutaneous interventions: a report from a large peripheral vascular registry. J Invasive Cardiol. 2009;21:628-31.
- 270. Matsi PJ, Manninen HI. Complications of lower-limb percutaneous transluminal angioplasty: a prospective analysis of 410 procedures on 295 consecutive patients. Cardiovasc Intervent Radiol. 1998;21:361-6.
- 271. Linni K, Ugurluoglu A, Hitzl W, et al. Bioabsorbable stent implantation vs. common femoral artery endarterectomy: early results of a randomized trial. J Endovasc Ther. 2014;21:493-502.
- 272 Gabrielli R, Rosati MS, Vitale S, et al. Randomized controlled trial of remote endarterectomy versus endovascular intervention for TransAtlantic Inter-Society Consensus II D femoropopliteal lesions. J Vasc Surg. 2012;56:1598-605
- 273. Gisbertz SS, Tutein Nolthenius RP, de Borst GJ, et al. Remote endarterectomy versus supragenicular bypass surgery for long occlusions of the superficial femoral artery: medium-term results of a randomized controlled trial (the REVAS trial). Ann Vasc Surg. 2010;24:1015-23
- 274 van Det RJ, Vriens BH, van der Palen J, et al. Dacron or ePTFE for femoro-popliteal above-knee bypass grafting: short- and long-term results of a multicentre randomised trial. Eur J Vasc Endovasc Surg. 2009;37:457-63.
- 275. Gisbertz SS, Ramzan M, Tutein Nolthenius RP, et al. Short-term results of a randomized trial comparing remote endarterectomy and supragenicular bypass
- 276 Ricco JB, Probst H. Long-term results of a multicenter randomized study on direct versus crossover bypass for unilateral illac artery occlusive disease. J Vasc surgery for long occlusions of the superficial femoral artery [the REVAS trial]. Eur J Vasc Endovasc Surg. 2009;37:68-76
- 277. Jensen LP, Lepäntalo M, Fossdal JE, et al. Dacron or PTFE for above-knee femoropopliteal bypass. a multicenter randomised study. Eur J Vasc Endovasc Surg. 2007;34:44-9 Surg. 2008;47:45-53.
- 278 AbuRahma AF, Robinson PA, Holt SM. Prospective controlled study of polytetrafluoroethylene versus saphenous vein in claudicant patients with bilateral above knee femoropopliteal bypasses. Surgery. 1999;126:594-602

- 279. Green RM, Abbott WM, Matsumoto T, et al. Prosthetic above-knee femoropopliteal bypass grafting: five-year results of a randomized trial. J Vasc Surg. 2000;31:417-25
- 280. Johnson WC, Lee KK. Comparative evaluation of externally supported Dacron and polytetrafluoroethylene prosthetic bypasses for femorofemoral and axillofemoral arterial reconstructions. Veterans Affairs Cooperative Study #141. J Vasc Surg. 1999;30:1077-83
- 281 Klinkert P, Schepers A, Burger DH, et al. Vein versus polytetrafluoroethylene in above-knee femoropopliteal bypass grafting: five-year results of a randomized controlled trial. J Vasc Surg. 2003;37:149-55.
- 282 Veith FJ, Gupta SK, Ascer E, et al. Six-year prospective multicenter randomized comparison of autologous saphenous vein and expanded polytetrafluoroethylene grafts in infrainguinal arterial reconstructions. J Vasc Surg. 1986;3:104-14.
- Nguyen BN, Amdur RL, Abugideiri M, et al. Postoperative complications after common femoral endarterectomy. J Vasc Surg. 2015;61:1489-94
- 283. 284. Lo RC, Bensley RP, Dahlberg SE, et al. Presentation, treatment, and outcome differences between men and women undergoing revascularization or amputation for lower extremity peripheral arterial disease. J Vasc Surg. 2014;59:409-18.
- 285 Surg. 2014;48:27-33. Siracuse JI, Gill HL, Schneider DB, et al. Assessing the perioperative safety of common femoral endarterectomy in the endovascular era. Vasc Endovascular
- 286. Aihara H, Soga Y, Mii S, et al. Comparison of long-term outcome after endovascular therapy versus bypass surgery in claudication patients with Trans-Atlantic Inter-Society Consensus-II C and D femoropopliteal disease. Circ J. 2014;78:457-64.
- 287. Bouff M, Azghari A, Belahda K, et al. Subintimal recanalization plus stenting or bypass for management of claudicants with femoro-popliteal occlusions. Eur J Vasc Endovasc Surg. 2013;46:347-52
- 289 288 Jones WS, Schmit KM, Vemulapalli S, et al. Treatment Strategies for Patients With Peripheral Artery Disease. Comparative Effectiveness Review No. 118. Sachwani GR, Hans SS, Khoury MD, et al. Results of iliac stenting and aortofemoral grafting for iliac artery occlusions. J Vasc Surg. 2013;57:1030-7
- 290 Antoniou GA, Chalmers N, Georgiadis GS, et al. A meta-analysis of endovascular versus surgical reconstruction of femoropopliteal arterial disease. J Vasc The Duke Evidence-based Practice Center under Contract No 290-2007-10066-I. 2013; Available at: ttp://www.effectivehealthcare.ahrq.gov/ehc/products/368/1415/Peripheral-Artery-Disease-Treatment-130301.pdf. Accessed September 25, 2016
- 291 Malgor RD, Ricotta JJ, Bower TC, et al. Common femoral artery endarterectomy for lower-extremity ischemia: evaluating the need for additional distal limb revascularization. Ann Vasc Surg. 2012;26:946-56. Surg. 2013;57:242-53.
- 292. Simons JP, Schanzer A, Nolan BW, et al. Outcomes and practice patterns in patients undergoing lower extremity bypass. J Vasc Surg. 2012;55:1629-36
- 293. Siracuse JJ, Giles KA, Pomposelli FB, et al. Results for primary bypass versus primary angioplasty/stent for intermittent claudication due to superficial femoral artery occlusive disease. J Vasc Surg. 2012;55:1001-7.
- 294 Kakkos SK, Haurani MJ, Shepard AD, et al. Patterns and outcomes of aortofemoral bypass grafting in the era of endovascular interventions. Eur J Vasc Endovasc Surg. 2011;42:658-66.
- 295. Simó G, Banga P, Darabos G, et al. Stent-assisted remote iliac artery endarterectomy: an alternative approach to treating combined external iliac and common femoral artery disease. Eur J Vasc Endovasc Surg. 2011;42:648-55
- 296. Eugster T, Marti R, Gurke L, et al. Ten years after arterial bypass surgery for claudication: venous bypass is the primary procedure for TASC C and D lesions. World J Surg. 2011;35:2328-31.
- 297. Piazza M, Ricotta JJ, Bower TC, et al. Iliac artery stenting combined with open femoral endarterectomy is as effective as open surgical reconstruction for severe thac and common femoral occlusive disease. J Vasc Surg. 2011;54:402-11.
- 298. Surg. 2010;39:597-603. Derksen WJ, Gisbertz SS, Hellings WE, et al. Predictive risk factors for restenosis after remote superficial femoral artery endarterectomy. Eur J Vasc Endovasc
- 299 Br J Surg. 2010;97:344-8 Koscielny A, Putz U, Willinek W, et al. Case-control comparison of profundaplasty and femoropopliteal supragenicular bypass for peripheral arterial disease

- 300. Ballotta E, Gruppo M, Mazzalai F, et al. Common femoral artery endarterectomy for occlusive disease: an 8-year single-center prospective study. Surgery. 2010;147:268-74.
- 301. disease. Ann Vasc Surg. 2010;24:4-13. Burke CR, Henke PK, Hernandez R, et al. A contemporary comparison of aortofemoral bypass and aortoiliac stenting in the treatment of aortoiliac occlusive
- 302 Twine CP, McLain AD. Graft type for femoro-popliteal bypass surgery. Cochrane Database Syst Rev. 2010;CD001487
- 303. atherosclerotic disease. Ann Vasc Surg. 2009;23:764-9. Chiesa R, Marone EM, Tshomba Y, et al. Aortobifemoral bypass grafting using expanded polytetrafluoroethylene stretch grafts in patients with occlusive
- 304 Al-Khoury G, Marone L, Chaer R, et al. Isolated femoral endarterectomy: impact of SFA TASC classification on recurrence of symptoms and need for additional intervention. J Vasc Surg. 2009;50:784-9.
- 305. Goodney PP, Likosky DS, Cronenwett JL, et al. Predicting ambulation status one year after lower extremity bypass. J Vasc Surg. 2009;49:1431-9
- 306. Chang RW, Goodney PP, Baek JH, et al. Long-term results of combined common femoral endarterectomy and iliac stenting/stent grafting for occlusive disease. J Vasc Surg. 2008;48:362-7.
- 307. Jaquinandi V, Picquet J, Bouye P, et al. High prevalence of proximal claudication among patients with patent aortobifemoral bypasses. J Vasc Surg 2007;45:312-8.
- 308 Fowkes F, Leng GC. Bypass surgery for chronic lower limb ischaemia. Cochrane Database Syst Rev. 2008;CD002000
- 310309 year follow-up. J Cardiovasc Surg (Torino ). 2006;47:385-91. Rosenthal D, Martin JD, Smeets L, et al. Remote superficial femoral artery endarterectomy and distal aSpire stenting: results of a multinational study at three-Pereira CE, Albers M, Romiti M, et al. Meta-analysis of femoropopliteal bypass grafts for lower extremity arterial insufficiency. J Vasc Surg. 2006;44:510-7.
- 311. Martin JD, Hupp JA, Peeler MO, et al. Remote endarterectomy: lessons learned after more than 100 cases. J Vasc Surg. 2006;43:320-6
- 312. Surgery. 2002;131:S269-74. Mori E, Komori K, Kume M, et al. Comparison of the long-term results between surgical and conservative treatment in patients with intermittent claudication.
- 313. Archie JP. Femoropopliteal bypass with either adequate ipsilateral reversed saphenous vein or obligatory polytetrafluoroethylene. Ann Vasc Surg. 1994;8:475-
- 314. Hunink MG, Wong JB, Donaldson MC, et al. Patency results of percutaneous and surgical revascularization for femoropopliteal arterial disease. Med Decis Making. 1994;14:71-81.
- 315. Schweiger H, Klein P, Lang W. Tibial bypass grafting for limb salvage with ringed polytetrafluoroethylene prostheses: results of primary and secondary procedures. J Vasc Surg. 1993;18:867-74
- 316. Baldwin ZK, Pearce BJ, Curi MA, et al. Limb salvage after infrainguinal bypass graft failure. J Vasc Surg. 2004;39:951-7
- 317 Leng GC, Lee AJ, Fowkes FG, et al. Incidence, natural history and cardiovascular events in symptomatic and asymptomatic peripheral arterial disease in the general population. Int J Epidemiol. 1996;25:1172-81.
- 318. Kannel WB, Skinner JJ, Schwartz MJ, et al. Intermittent claudication. Incidence in the Framingham Study. Circulation. 1970;41:875-83
- 319. Kannel WB, Shurtleff D. The natural history of arteriosclerosis obliterans. Cardiovasc Clin. 1971;3:37-52
- 320. TILLGREN C. Obliterative Arterial Disease of the Lower Limbs. II. A Study of the Course of the Disease. Acta Med Scand. 1965;178:103-19
- 321. 322. Jelnes R, Gaardsting O, Hougaard Jensen K, et al. Fate in intermittent claudication: outcome and risk factors. Br Med J (Clin Res Ed). 1986;293:1137-40.
- Bloor K. Natural history of arteriosclerosis of the lower extremities: Hunterian lecture delivered at the Royal College of Surgeons of England on 22nd April 1960. Ann R Coll Surg Engl. 1961;28:36-52.
- 323. 324. Dormandy J, Mahir M, Ascady G, et al. Fate of the patient with chronic leg ischaemia. A review article. J Cardiovasc Surg (Torino ). 1989;30:50-7
- Gandini R, Del GC, Merolla S, et al. Treatment of chronic SFA in-stent occlusion with combined laser atherectomy and drug-eluting balloon angioplasty in patients with critical limb ischemia: a single-center, prospective, randomized study. J Endovasc Ther. 2013;20:805-14.
- 325 Zeller T, Baumgartner I, Scheinert D, et al. Drug-eluting balloon versus standard balloon angioplasty for infrapopliteal arterial revascularization in critical limb ischemia: 12-month results from the IN.PACT DEEP randomized trial. J Am Coll Cardiol. 2014;64:1568-76

- 326. Scheinert D, Katsanos K, Zeller T, et al. A prospective randomized multicenter comparison of balloon angioplasty and infrapopliteal stenting with the
- 327. Katsanos K, Spiliopoulos S, Diamantopoulos A, et al. Wound healing outcomes and health-related quality-of-life changes in the ACHILLES trial: 1-year results from a prospective randomized controlled trial of infrapopliteal balloon angioplasty versus sirolimus-eluting stenting in patients with ischemic sirolimus-eluting stent in patients with ischemic peripheral arterial disease: 1-year results from the ACHILLES trial. J Am Coll Cardiol. 2012;60:2290-5. peripheral arterial disease. JACC Cardiovasc Interv. 2016;9:259-67.
- 328 Adam DJ, Beard JD, Cleveland T, et al. Bypass versus angioplasty in severe ischaemia of the leg (BASIL): multicentre, randomised controlled trial. Lancet 2005;366:1925-34.
- 329 angioplasty-first revascularisation strategy for severe limb ischaemia due to infrainguinal disease. The Bypass versus Angioplasty in Severe Ischaemia of the Bradbury AW, Adam DJ, Bell J, et al. Multicentre randomised controlled trial of the clinical and cost-effectiveness of a bypass-surgery-first versus a balloon Leg (BASIL) trial. Health Technol Assess. 2010;14:1-210, iii-iv.
- 330. survival by treatment received. J Vasc Surg. 2010;51:18S-31S. Bradbury AW, Adam DJ, Bell J, et al. Bypass versus Angioplasty in Severe Ischaemia of the Leg (BASIL) trial: Analysis of amputation free and overall
- 331. Fanelli F, Cannavale A, Boatta E, et al. Lower limb multilevel treatment with drug-eluting balloons: 6-month results from the DEBELLUM randomized trial. J Endovasc Ther. 2012;19:571-80.
- 332. Rosenfield K, Jaff MR, White CJ, et al. Trial of a paclitaxel-coated balloon for femoropopliteal artery disease. N Engl J Med. 2015;373:145-53
- 333. Bosiers M, Scheinert D, Peeters P, et al. Randomized comparison of everolimus-eluting versus bare-metal stents in patients with critical limb ischemia and infrapopliteal arterial occlusive disease. J Vasc Surg. 2012;55:390-8.
- 334. Rastan A, Tepe G, Krankenberg H, et al. Sirolimus-eluting stents vs. bare-metal stents for treatment of focal lesions in infrapopliteal arteries: a double-blind, multi-centre, randomized clinical trial. Eur Heart J. 2011;32:2274-81.
- 335. Siablis D, Kitrou PM, Spiliopoulos S, et al. Paclitaxel-coated balloon angioplasty versus drug-eluting stenting for the treatment of infrapopliteal long-segment arterial occlusive disease: the IDEAS randomized controlled trial. JACC Cardiovasc Interv. 2014;7:1048-56.
- 336 Tepe G, Schnorr B, Albrecht T, et al. Angioplasty of femoral-popliteal arteries with drug-coated balloons: 5-year follow-up of the THUNDER trial. JACC Cardiovasc Interv. 2015;8:102-8.
- 337. Ferraresi R, Centola M, Ferlini M, et al. Long-term outcomes after angioplasty of isolated, below-the-knee arteries in diabetic patients with critical limb ischaemia. Eur J Vasc Endovasc Surg. 2009;37:336-42.
- 338. Park SW, Kim JS, Yun IJ, et al. Clinical outcomes of endovascular treatments for critical limb ischemia with chronic total occlusive lesions limited to belowthe-knee arteries. Acta Radiol. 2013;54:785-9
- 339. Faglia E, Clerici G, Clerissi J, et al. Early and five-year amputation and survival rate of diabetic patients with critical limb ischemia: data of a cohort study of 564 patients. Eur J Vasc Endovasc Surg. 2006;32:484-90.
- 340. Faglia E, Dalla PL, Clerici G, et al. Peripheral angioplasty as the first-choice revascularization procedure in diabetic patients with critical limb ischemia
- 341. lida O, Soga Y, Hirano K, et al. Long-term results of direct and indirect endovascular revascularization based on the angiosome concept in patients with critical prospective study of 993 consecutive patients hospitalized and followed between 1999 and 2003. Eur J Vasc Endovasc Surg. 2005;29:620-7
- 342 Feiring AJ, Krahn M, Nelson L, et al. Preventing leg amputations in critical limb ischemia with below-the-knee drug-eluting stents: the PaRADISE limb ischemia presenting with isolated below-the-knee lesions. J Vasc Surg. 2012;55:363-70
- 343. Siablis D, Karnabatidis D, Katsanos K, et al. Infrapopliteal application of sirolimus-eluting versus bare metal stents for critical limb ischemia: analysis of longterm angiographic and clinical outcome. J Vasc Interv Radiol. 2009;20:1141-50 (PReventing Amputations using Drug eluting StEnts) trial. J Am Coll Cardiol. 2010;55:1580-9
- 344. angiographic follow-up. J Endovasc Ther. 2012;19:12-9 Werner M, Schmidt A, Freyer M, et al. Sirolimus-eluting stents for the treatment of infrapopliteal arteries in chronic limb ischemia: long-term clinical and
- 345. Acin F, Varela C, López de Maturana I, et al. Results of infrapopliteal endovascular procedures performed in diabetic patients with critical limb ischemia and tissue loss from the perspective of an angiosome-oriented revascularization strategy. Int J Vasc Med. 2014;2014:270539.

- 346. Alexandrescu VA, Hubermont G, Philips Y, et al. Selective primary angioplasty following an angiosome model of reperfusion in the treatment of Wagner 1-4 diabetic foot lesions: practice in a multidisciplinary diabetic limb service. J Endovasc Ther. 2008;15:580-93
- 347. Fossaceca R, Guzzardi G, Cerini P, et al. Endovascular treatment of diabetic foot in a selected population of patients with below-the-knee disease: is the angiosome model effective? Cardiovasc Intervent Radiol. 2013;36:637-44.
- 348 Kabra A, Suresh KR, Vivekanand V, et al. Outcomes of angiosome and non-angiosome targeted revascularization in critical lower limb ischemia. J Vasc Surg 2013;57:44-9.
- 349 revascularization for critical limb ischemia. J Vasc Surg. 2014;59:121-8. Kret MR, Cheng D, Azarbal AF, et al. Utility of direct angiosome revascularization and runoff scores in predicting outcomes in patients undergoing
- 350. diabetic patients with critical limb ischemia. Ann Vasc Surg. 2014;28:983-9 Lejay A, Georg Y, Tartaglia E, et al. Long-term outcomes of direct and indirect below-the-knee open revascularization based on the angiosome concept in
- 351 Neville RF, Attinger CE, Bulan EJ, et al. Revascularization of a specific angiosome for limb salvage: does the target artery matter? Ann Vasc Surg 2009;23:367-73
- 352. Osawa S, Terashi H, Tsuji Y, et al. Importance of the six angiosomes concept through arterial-arterial connections in CLI. Int Angiol. 2013;32:375-85
- 353. Abu Dabrh AM, Steffen MW, Undavalli C, et al. The natural history of untreated severe or critical limb ischemia. J Vasc Surg. 2015;62:1642-51.
- 354 Abidia A, Laden G, Kuhan G, et al. The role of hyperbaric oxygen therapy in ischaemic diabetic lower extremity ulcers: a double-blind randomised-controlled trial. Eur J Vasc Endovasc Surg. 2003;25:513-8
- 355 results of a prospective randomized trial. The STILE Investigators. Surgery versus Thrombolysis for Ischemia of the Lower Extremity. J Vasc Surg Weaver FA, Comerota AJ, Youngblood M, et al. Surgical revascularization versus thrombolysis for nonembolic lower extremity native artery occlusions: 1996;24:513-21
- 356 Ouriel K, Veith FJ, Sasahara AA. A comparison of recombinant urokinase with vascular surgery as initial treatment for acute arterial occlusion of the legs Thrombolysis or Peripheral Arterial Surgery (TOPAS) Investigators. N Engl J Med. 1998;338:1105-11.
- 357. Ponec D, Jaff MR, Swischuk J, et al. The Nitinol SMART stent vs Wallstent for suboptimal iliac artery angioplasty: CRISP-US trial results. J Vasc Interv Radiol. 2004;15:911-8.
- 358. Schillinger M, Sabeti S, Loewe C, et al. Balloon angioplasty versus implantation of nitinol stents in the superficial femoral artery. N Engl J Med 2006;354:1879-88.
- 359 Conte MS, Bandyk DF, Clowes AW, et al. Results of PREVENT III: a multicenter, randomized trial of edifoligide for the prevention of vein graft failure in lower extremity bypass surgery. J Vasc Surg. 2006;43:742-51
- 360. Vasc Interv Radiol. 2014;17:221-4. Farber A, Rosenfield K, Menard M. The BEST-CLI trial: a multidisciplinary effort to assess which therapy is best for patients with critical limb ischemia. Tech
- 361. management of diabetic foot ulcers. Arch Surg. 2002;137:822-7 Veves A, Sheehan P, Pham HT. A randomized, controlled trial of Promogran (a collagen/oxidized regenerated cellulose dressing) vs standard treatment in the
- 362 Fogle MA, Whittemore AD, Couch NP, et al. A comparison of in situ and reversed saphenous vein grafts for infrainguinal reconstruction. J Vasc Surg 1987;5:46-52.
- 363 bypass outcome. J Vasc Surg. 2013;57:1219-26. Rashid H, Slim H, Zayed H, et al. The impact of arterial pedal arch quality and angiosome revascularization on foot tissue loss healing and infrapopliteal
- 364 Nolan BW, De Martino RR, Stone DH, et al. Prior failed ipsilateral percutaneous endovascular intervention in patients with critical limb ischemia predicts poor outcome after lower extremity bypass. J Vasc Surg. 2011;54:730-5.
- 365. Santo VJ, Dargon P, Azarbal AF, et al. Lower extremity autologous vein bypass for critical limb ischemia is not adversely affected by prior endovascular procedure. J Vasc Surg. 2014;60:129-35.
- 366 Uhl C, Hock C, Betz T, et al. Pedal bypass surgery after crural endovascular intervention. J Vasc Surg. 2014;59:1583-7

- 367. Korhonen M, Biancari F, Söderström M, et al. Femoropopliteal balloon angioplasty vs. bypass surgery for CLI: a propensity score analysis. Eur J Vasc Endovasc Surg. 2011;41:378-84
- 368 Kasemi H, Marino M, Dionisi CP, et al. Seven-year approach evolution of the aortoiliac occlusive disease endovascular treatment. Ann Vasc Surg 2016;30:277-85
- 369 Bredahl K, Jensen LP, Schroeder TV, et al. Mortality and complications after aortic bifurcated bypass procedures for chronic aortoiliac occlusive disease. J Vasc Surg. 2015;62:75-82.
- 370. 371. Ruffolo AJ, Romano M, Ciapponi A. Prostanoids for critical limb ischaemia. Cochrane Database Syst Rev. 2010;CD006544. Chew DK, Conte MS, Donaldson MC, et al. Autogenous composite vein bypass graft for infrainguinal arterial reconstruction. J Vasc Surg. 2001;33:259-64
- Moran PS, Teljeur C, Harrington P, et al. A systematic review of intermittent pneumatic compression for critical limb ischaemia. Vasc Med. 2015;20:41-50
- 372. 373. endovascular therapy. J Vasc Surg. 2015;61:951-9. Kobayashi N, Hirano K, Nakano M, et al. Prognosis of critical limb ischemia patients with tissue loss after achievement of complete wound healing by
- 374. Rutherford RB. Acute limb ischemia: Clinical assessment and standards for reporting. Semin Vasc Surg. 1992;5:4-10.
- 375. Nypaver TJ, Whyte BR, Endean ED, et al. Nontraumatic lower-extremity acute arterial ischemia. Am J Surg. 1998;176:147-52
- 376. FOGARTY TJ, CRANLEY JJ. Catheter Technic for Arterial Embolectomy. Ann Surg. 1965;161:325-30.
- 377. Shin HS, Kyoung KH, Suh BJ, et al. Acute limb ischemia: surgical thromboembolectomy and the clinical course of arterial revascularization at ankle. Int J Angiol. 2013;22:109-14.
- 378 acute lower limb ischemia. J Vasc Surg. 2014;59:729-36. de Donato G, Setacci F, Sirignano P, et al. The combination of surgical embolectomy and endovascular techniques may improve outcomes of patients with
- 379 Baril DT, Patel VI, Judelson DR, et al. Outcomes of lower extremity bypass performed for acute limb ischemia. J Vasc Surg. 2013;58:949-56
- 380. 381. Duval S, Keo HH, Oldenburg NC, et al. The impact of prolonged lower limb ischemia on amputation, mortality, and functional status: the FRIENDS registry Manojlovic V, Popovic V, Nikolic D, et al. Analysis of associated diseases in patients with acute critical lower limb ischemia. Med Pregl. 2013;66:41-5
- 382. Morris-Stiff G, D'Souza J, Raman S, et al. Update experience of surgery for acute limb ischaemia in a district general hospital-are we getting any better? Ann R Am Heart J. 2014;168:577-87
- Coll Surg Engl. 2009;91:637-40.
- 383. delay. World J Emerg Surg. 2014;9:56. Londero LS, Nørgaard B, Houlind K. Patient delay is the main cause of treatment delay in acute limb ischemia: an investigation of pre- and in-hospital time
- 384 Ouriel K, Shortell CK, DeWeese JA, et al. A comparison of thrombolytic therapy with operative revascularization in the initial treatment of acute peripheral arterial ischemia. J Vasc Surg. 1994;19:1021-30.
- 385 Results of a prospective randomized trial evaluating surgery versus thrombolysis for ischemia of the lower extremity. The STILE trial. Ann Surg 1994;220:266-8.
- 386. Comerota AJ, Weaver FA, Hosking JD, et al. Results of a prospective, randomized trial of surgery versus thrombolysis for occluded lower extremity bypass grafts. Am J Surg. 1996;172:105-12
- 387. Diffin DC, Kandarpa K. Assessment of peripheral intraarterial thrombolysis versus surgical revascularization in acute lower-limb ischemia: a review of limb-
- 388. Schrijver AM, Reijnen MM, van Oostayen JA, et al. Dutch randomized trial comparing standard catheter-directed thrombolysis versus ultrasound-accelerated salvage and mortality statistics. J Vasc Interv Radiol. 1996;7:57-63.
- 389 Fagundes C, Fuchs FD, Fagundes A, et al. Prognostic factors for amputation or death in patients submitted to vascular surgery for acute limb ischemia. Vasc thrombolysis for thromboembolic infrainguinal disease (DUET): design and rationale. Trials. 2011;12:20 Health Risk Manag. 2005;1:345-9.
- 390. FOGARTY TJ, CRANLEY JJ, KRAUSE RJ, et al. A method for extraction of arterial emboli and thrombi. Surg Gynecol Obstet. 1963;116:241-4
- 391 Eliason JL, Wakefield TW. Metabolic consequences of acute limb ischemia and their clinical implications. Semin Vasc Surg. 2009;22:29-33

- 392. settings. J Vasc Surg. 2015;61:138-46. Lurie F, Vaidya V, Comerota AJ. Clinical outcomes and cost-effectiveness of initial treatment strategies for nonembolic acute limb ischemia in real-life clinical
- 393. Taha AG, Byrne RM, Avgerinos ED, et al. Comparative effectiveness of endovascular versus surgical revascularization for acute lower extremity ischemia. J Vasc Surg. 2015;61:147-54
- 394 Gupta R, Hennebry TA. Percutaneous isolated pharmaco-mechanical thrombolysis-thrombectomy system for the management of acute arterial limb ischemia: 30-day results from a single-center experience. Catheter Cardiovasc Interv. 2012;80:636-43
- 395 salvage and survival results from a single center series. Catheter Cardiovasc Interv. 2008;72:325-30. Ansel GM, Botti CF, Silver MJ. Treatment of acute limb ischemia with a percutaneous mechanical thrombectomy-based endovascular approach: 5-year limb
- 396. 397. Byrne RM, Taha AG, Avgerinos E, et al. Contemporary outcomes of endovascular interventions for acute limb ischemia. J Vasc Surg. 2014;59:988-95
- Schernthaner MB, Samuels S, Biegler P, et al. Ultrasound-accelerated versus standard catheter-directed thrombolysis in 102 patients with acute and subacute limb ischemia. J Vasc Interv Radiol. 2014;25:1149-56.
- 398 Silva JA, Ramee SR, Collins TJ, et al. Rheolytic thrombectomy in the treatment of acute limb-threatening ischemia: immediate results and six-month follow-up of the multicenter AngioJet registry. Possis Peripheral AngioJet Study AngioJet Investigators. Cathet Cardiovasc Diagn. 1998;45:386-93
- 399 Kasirajan K, Gray B, Beavers FP, et al. Rheolytic thrombectomy in the management of acute and subacute limb-threatening ischemia. J Vasc Interv Radiol 2001;12:413-21.
- 400 Allie DE, Hebert CJ, Lirtzman MD, et al. Novel simultaneous combination chemical thrombolysis/rheolytic thrombectomy therapy for acute critical limb schemia: the power-pulse spray technique. Catheter Cardiovasc Interv. 2004;63:512-22.
- 401 Elmahdy MF, Ghareeb MS, Baligh EE, et al. Value of duplex scanning in differentiating embolic from thrombotic arterial occlusion in acute limb ischemia Cardiovasc Revasc Med. 2010;11:223-6.
- 402. Ascher E, Hingorani A, Markevich N, et al. Acute lower limb ischemia: the value of duplex ultrasound arterial mapping (DUAM) as the sole preoperative imaging technique. Ann Vasc Surg. 2003;17:284-9.
- 403. Lowery AJ, Hynes N, Manning BJ, et al. A prospective feasibility study of duplex ultrasound arterial mapping, digital-subtraction angiography, and magnetic resonance angiography in management of critical lower limb ischemia by endovascular revascularization. Ann Vasc Surg. 2007;21:443-51
- 404. Leung DA, Blitz LR, Nelson T, et al. Rheolytic pharmacomechanical thrombectomy for the management of acute limb ischemia: results from the PEARL Registry. J Endovasc Ther. 2015;22:546-57
- 405. Schrijver AM, Reijnen MM, van Oostayen JA, et al. Initial results of catheter-directed ultrasound-accelerated thrombolysis for thromboembolic obstructions of the aortofemoral arteries: a feasibility study. Cardiovasc Intervent Radiol. 2012;35:279-85
- 406. Schrijver A, Vos J, Hoksbergen AW, et al. Ultrasound-accelerated thrombolysis for lower extremity ischemia: multicenter experience and literature review. J Cardiovasc Surg (Torino ). 2011;52:467-76.
- 407. Ihlberg L, Luther M, Albäck A, et al. Does a completely accomplished duplex-based surveillance prevent vein-graft failure? Eur J Vasc Endovasc Surg [999;18:395-400.
- 408 study. J Vasc Surg. 1995;21:26-33. Lundell A, Lindblad B, Bergqvist D, et al. Femoropopliteal-crural graft patency is improved by an intensive surveillance program: a prospective randomized
- 409. Jongsma H, Bekken JA, van Buchem F, et al. Secondary interventions in patients with autologous infrainguinal bypass grafts strongly improve patency rates. J
- 410. Carter A, Murphy MO, Halka AT, et al. The natural history of stenoses within lower limb arterial bypass grafts using a graft surveillance program. Ann Vasc Surg. 2007;21:695-703. Vasc Surg. 2016;63:385-90.
- 411. Westerband A, Mills JL, Kistler S, et al. Prospective validation of threshold criteria for intervention in infrainguinal vein grafts undergoing duplex surveillance Ann Vasc Surg. 1997;11:44-8
- 412. Mills JL, Harris EJ, Taylor LM, et al. The importance of routine surveillance of distal bypass grafts with duplex scanning: a study of 379 reversed vein grafts. J Vasc Surg. 1990;12:379-86

- 413. Brumberg RS, Back MR, Armstrong PA, et al. The relative importance of graft surveillance and warfarin therapy in infrainguinal prosthetic bypass failure. J Vasc Surg. 2007;46:1160-6.
- 414. Calligaro KD, Doerr K, McAffee-Bennett S, et al. Should duplex ultrasonography be performed for surveillance of femoropopliteal and femorotibial arterial prosthetic bypasses? Ann Vasc Surg. 2001;15:520-4.
- 415. Stone PA, Armstrong PA, Bandyk DF, et al. Duplex ultrasound criteria for femorofemoral bypass revision. J Vasc Surg. 2006;44:496-502
- 416. Back MR, Novotney M, Roth SM, et al. Utility of duplex surveillance following iliac artery angioplasty and primary stenting. J Endovasc Ther. 2001;8:629-37.
- 417. Baril DT, Marone LK. Duplex evaluation following femoropopliteal angioplasty and stenting: criteria and utility of surveillance. Vasc Endovascular Surg. 2012;46:353-7.
- 419. 418. Troutman DA, Madden NJ, Dougherty MJ, et al. Duplex ultrasound diagnosis of failing stent grafts placed for occlusive disease. J Vasc Surg. 2014;60:1580-4
- Connors G, Todoran TM, Engelson BA, et al. Percutaneous revascularization of long femoral artery lesions for claudication: patency over 2.5 years and impact of systematic surveillance. Catheter Cardiovasc Interv. 2011;77:1055-62.

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Blue Cross/Blue Shield Association- Medical Advisory Panel to the Technology	<ul> <li>Abbott Vascular</li> <li>Corindus Vascular Robotics</li> <li>St. Jude Medical</li> </ul>	None	None	None	None	None	arch 2015) Consultant
None	None	None	None	None	None	None	Speakers Bureau
None	None	None	None	None	<ul> <li>Summit Doppler Systems</li> <li>Zin Medical</li> </ul>	None	Ownership/ Partnership/ Principal
<ul> <li>Johns Hopkins Medical Institutions</li> </ul>	<ul> <li>Atrium Medical</li> <li>Bard</li> <li>iDEV Technologies</li> <li>Lutonix</li> <li>PLC Medical Systems</li> <li>Prairie Education and Research Cooperative</li> </ul>	None	None	None	<ul> <li>AstraZeneca</li> <li>Theravasc</li> </ul>	<ul> <li>Progeria Research Foundation*</li> </ul>	Personal Research
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M. Eileen Walsh	University of Toledo, College of Nursing—Professor	None	None	None	None	<ul> <li>American Nurses Credentialing Center†</li> <li>Journal of Vascular Nursing (Editorial Board)†</li> </ul>	None
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DSMB indicates data safety monitoring board; IAC, Intersocietal Accreditation Commission; NHLBI, National Heart, Lung, and Blood Institute; NIH, National Institute of Health; and PCORI, Patient-Centered Outcomes Research Institute.