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ESC Guidelines on the diagnosis and treatment of peripheral artery diseases

- ¹⁰ Document covering atherosclerotic disease of extracranial carotid and vertebral, mesenteric, renal, upper and lower extremity arteries
- ¹⁵ The Task Force on the Diagnosis and Treatment of Peripheral Artery Diseases of the European Society of Cardiology (ESC)

Endorsed by: the European Stroke Organisation (ESO) $_{\scriptscriptstyle 20}$

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pheral artery disease • Carotid artery disease • Vertebral artery disease • Upper extremity artery
ase • Mesenteric artery disease • Renal artery disease • Lower extremity artery disease • Multisite
ry disease

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www.escardio.org/guidelines		

Abbreviations and acronyms

2D 3D ABI	two-dimensional three-dimensional ankle–brachial index	325
ACAS	Asymptomatic Carotid Atherosclerosis Study	
ACCF	American College of Cardiology Foundation	
ACE	angiotensin-converting enzyme	
ACS	acute coronary syndrome	330
ACST	Asymptomatic Carotid Surgery Trial	
ALI	acute limb ischaemia	
ASTRAL	Angioplasty and Stenting for Renal Artery Lesions	
	trial	
BASIL	Bypass versus Angioplasty in Severe Ischaemia of the Leg	335
BOA	Dutch Bypass Oral Anticoagulants or Aspirin	
CABG	coronary artery bypass grafting	
CAD	coronary artery disease	
CAPRIE	Clopidogrel versus Aspirin in Patients at Risk for	340
	Ischaemic Events	510
CAPTURE	Carotid ACCULINK/ACCUNET Post Approval	
	Trial to Uncover Rare Events	

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	CARP	Coronary Artery Revascularization Prophylaxis
	CAS	carotid artery stenting
345	CASPAR	Clopidogrel and Acetylsalicylic Acid in Bypass
		Surgery for Peripheral Arterial Disease
	CASS	Coronary Artery Surgery Study
	CAVATAS	CArotid and Vertebral Artery Transluminal Angio-
		plasty Study
350	CEA	carotid endarterectomy
	CHARISMA	Clopidogrel for High Atherothrombotic Risk and
		Ischaemic Stabilization, Management and
		Avoidance
	CI	confidence interval
355	CLEVER	Claudication: Exercise Versus Endoluminal
555	CLLVLIN	Revascularization
	CLI	critical limb ischaemia
	CORAL	
	CORAL	Cardiovascular Outcomes in Renal Atherosclero-
2/0		tic Lesions
360	COURAGE	Clinical Outcomes Utilization Revascularization
	6D.6	and Aggressive Drug Evaluation
	CPG	Committee for Practice Guidelines
	CREST	Carotid Revascularization Endarterectomy vs.
	CT	Stenting Trial
365	CT	computed tomography
	CTA	computed tomography angiography
	CVD	cardiovascular disease
	DECREASE-V	0 1
	DRASTIC	Dutch Renal Artery Stenosis Intervention Coop-
370		erative Study
	DSA	digital subtraction angiography
	DUS	duplex ultrasound/duplex ultrasonography
	EACTS	European Association for Cardio-Thoracic Surgery
	EAS	European Atherosclerosis Society
375	ECST	European Carotid Surgery Trial
	EPD	embolic protection device
	ESC	European Society of Cardiology
	ESH	European Society of Hypertension
	ESRD	end-stage renal disease
380	EUROSCORE	European System for Cardiac Operative Risk
		Evaluation
	EVA-3S	Endarterectomy Versus Angioplasty in Patients
		with Symptomatic Severe Carotid Stenosis
	EXACT	Emboshield and Xact Post Approval Carotid Stent
385		Trial
	GALA	General Anaesthesia versus Local Anaesthesia for
		Carotid Surgery
	GFR	glomerular filtration rate
	GRACE	Global Registry of Acute Coronary Events
390	Hb _{A1c}	glycated haemoglobin
	HDL	high-density lipoprotein
	HOPE	Heart Outcomes Prevention Evaluation
	HR	hazard ratio
	IC	intermittent claudication
395	ICSS	International Carotid Stenting Study
	IMT	intima-media thickness
	ITT	intention to treat

LDL	low-density lipoprotein	400
LEAD	lower extremity artery disease	
MACCEs	major adverse cardiac and cerebrovascular events	
MDCT	multidetector computed tomography	
MONICA	Monitoring of Trends and Determinants in Cardio-	
	vascular Disease	405
MRA	magnetic resonance angiography	
MRI	magnetic resonance imaging	
NASCET	North American Symptomatic Carotid Endarter-	
	ectomy Trial	
ONTARGET	Ongoing Telmisartan Alone and in Combination	410
	with Ramipril Global Endpoint Trial	
OR	odds ratio	
PAD	peripheral artery diseases	
PARTNERS	Peripheral Arterial Disease Awareness, Risk, and	
	Treatment: New Resources for Survival	415
PCI	percutaneous coronary intervention	
PET	positron emission tomography	
PRO-CAS	Predictors of Death and Stroke in CAS	
PTA	percutaneous transluminal angioplasty	
RAAS	renin–angiotensin–aldosterone system	420
RADAR	Randomized, Multicentre, Prospective Study Com-	
	paring Best Medical Treatment Versus Best	
	Medical Treatment Plus Renal Artery Stenting in	
	Patients With Haemodynamically Relevant Athero-	
	sclerotic Renal Artery Stenosis	425
RAS	renal artery stenosis	
RCT	randomized controlled trial	
REACH	Reduction of Atherothrombosis for Continued	
	Health	
RR	risk ratio	430
SAPPHIRE	Stenting and Angioplasty with Protection in	
	Patients at High Risk for Endarterectomy	
SCAI	Society for Cardiovascular Angiography and	
	Interventions	
SIR	Society of Interventional Radiology	435
SPACE	Stent-Protected Angioplasty versus Carotid	
	Endarterectomy	
SPARCL	Stroke Prevention by Aggressive Reduction in	
STAR	Cholesterol Levels Study Stent Placement in Patients With Atherosclerotic	440
STAR		440
SSYLVIA	Renal Artery Stenosis and Impaired Renal Function	
SSTEVIA	Stenting of Symptomatic Atherosclerotic Lesions in the Vertebral or Intracranial Arteries	
SVMB	Society for Vascular Medicine and Biology	
TASC	TransAtlantic Inter-Society Consensus	445
TIA	transient ischaemic attack	СНТ
UEAD	upper extremity artery disease	
VA	vertebral artery	
7/7	ver teor at ar ter y	

1. Preamble

Guidelines summarize and evaluate all available evidence, at the time of the writing process, on a particular issue with the aim of assisting physicians in selecting the best management strategies

for an individual patient, with a given condition, taking into account the impact on outcome, as well as the risk-benefit ratio of particular diagnostic or therapeutic means. Guidelines are no substitutes

- ⁴⁶⁰ but are complements for textbooks and cover the ESC Core Curriculum topics. Guidelines and recommendations should help the physicians to make decisions in their daily practice. However, the final decisions concerning an individual patient must be made by the responsible physician(s).
- A large number of Guidelines have been issued in recent years by the European Society of Cardiology (ESC) as well as by other societies and organizations. Because of the impact on clinical practice, quality criteria for the development of guidelines have been established in order to make all decisions transparent to the
- 470 user. The recommendations for formulating and issuing ESC Guidelines can be found on the ESC website (http://www. escardio.org/guidelines-surveys/esc-guidelines/about/Pages/rules-writ ing.aspx). ESC Guidelines represent the official position of the ESC on a given topic and are regularly updated.
- 475 Members of this Task Force were selected by the ESC to represent professionals involved with the medical care of patients with this pathology. Selected experts in the field undertook a comprehensive review of the published evidence for diagnosis, management, and/or prevention of a given condition according to ESC
- 480 Committee for Practice Guidelines (CPG) policy. A critical evaluation of diagnostic and therapeutic procedures was performed including assessment of the risk-benefit ratio. Estimates of expected health outcomes for larger populations were included, where data exist. The level of evidence and the strength of rec-
- 485 ommendation of particular treatment options were weighed and graded according to pre-defined scales, as outlined in *Tables 1* and 2.

The experts of the writing and reviewing panels filled in declarations of interest forms of all relationships which might be perceived as real or potential sources of conflicts of interest. These forms were compiled into one file and can be found on the ESC website (http://www.escardio.org/guidelines). Any changes in declarations of interest that arise during the writing period must 515 be notified to the ESC and updated. The Task Force received its entire financial support from the ESC without any involvement from the healthcare industry.

The ESC CPG supervises and coordinates the preparation of new Guidelines produced by Task Forces, expert groups, 520 or consensus panels. The Committee is also responsible for the endorsement process of these Guidelines. The ESC Guidelines undergo extensive review by the CPG and external experts. After appropriate revisions, it is approved by all the experts involved in the Task Force. The finalized document is 525 approved by the CPG for publication in the *European Heart Journal*.

The task of developing Guidelines covers not only the integration of the most recent research, but also the creation of educational tools and implementation programmes for the 530 recommendations. To implement the guidelines, condensed pocket guidelines versions, summary slides, booklets with essential messages, and electronic version for digital applications (smartphones, etc.), are produced. These versions are abridged and, thus, if needed, one should always refer to the full text version 535 which is freely available on the ESC website. The National Societies of the ESC are encouraged to endorse, translate, and implement the ESC Guidelines. Implementation programmes are needed because it has been shown that the outcome of disease may be favourably influenced by the thorough application of clinical 540 recommendations.

Surveys and registries are needed to verify that real-life daily practice is in keeping with what is recommended in the guidelines, thus completing the loop between clinical research, writing of Guidelines, and implementing them into clinical practice. 545

The Guidelines do not, however, override the individual responsibility of health professionals to make appropriate decisions in the circumstances of the individual patients, in consultation with that

Classes of recommendations	Definition Suggested wording to use	
Class I	Evidence and/or general agreement that a given treatment or procedure is beneficial, useful, effective.	Is recommended/is indicated
Class II	Conflicting evidence and/or a divergence of opinion about the usefulness/efficacy of the given treatment or procedure.	
Class Ila	Weight of evidence/opinion is in favour of usefulness/efficacy.	Should be considered
Class IIb	Usefulness/efficacy is less well established by evidence/opinion.	May be considered
Class III	Evidence or general agreement that the given treatment or procedure is not useful/effective, and in some cases may be harmful.	Is not recommended

Table I Classes of recommendations

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Table 2 Levels of evidence

575	Level of Evidence A	Data derived from multiple randomized clinical trials or meta-analyses.
	Level of Evidence B	Data derived from a single randomized clinical trial or large non-randomized studies.
580	Level of Evidence C	Consensus of opinion of the experts and/ or small studies, retrospective studies, registries.

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patient, and, where appropriate and necessary, the patient's guardian or carer. It is also the health professional's responsibility to verify the rules and regulations applicable to drugs and devices at the time of prescription.

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2. Introduction

Cardiovascular diseases (CVDs) are the leading cause of death and disability in Europe, posing a great social and economic burden. Coronary artery disease (CAD) is the cause of death in a large percentage of individuals, but stroke, renal failure, and complications from severe ischaemia of the lower extremities also contribute to an adverse prognosis.

Since atherosclerosis is a systemic disease, physicians must 600 appreciate the importance of detecting atherosclerosis in other vascular beds in order to establish the correct treatment to prevent organ damage. As shown recently by the Reduction of Atherothrombosis for Continued Health (REACH) Registry, a substantial percentage of patients with chronic CAD have associated cerebro-605 vascular disease, lower extremity artery disease (LEAD), or both.¹

This is the first document produced by the ESC addressing different aspects of peripheral artery diseases (PAD). This task has been undertaken because an increasing proportion of patients with heart disease need to be assessed for vascular problems in

610 other territories, both symptomatic and asymptomatic, that may affect their prognosis and treatment strategy. It is also recognized that patients with PAD will probably die from CAD.²

In this document the term PAD is used to include all vascular sites, including carotid, vertebral, upper extremity, mesenteric,

renal, and lower extremity vessels. Diseases of the aorta are not 615 covered.

Although different disease processes may cause PAD, the Task Force decided to focus on atherosclerosis. Other aetiologies, specific for different vascular territories, are mentioned but not 620 discussed.

Atherosclerosis in the peripheral arteries is a chronic, slowly developing condition causing narrowing of the arteries. Depending on the degree of narrowing at each vascular site, a range of severity of symptoms may occur, while many patients will remain asympto-

matic throughout their life. Occasionally acute events occur, often 625 associated with thrombosis and/or embolism and/or occlusion of a major artery.

In the first section of this document the general issues are addressed, whereas the detailed clinical presentations are covered in specific sections for each vascular site. Special emphasis 630 is put on multisite artery disease (e.g. patients with CAD plus disease in another vascular bed), addressing most common aspects from a diversity of complex clinical scenarios encountered in clinical practice. Finally, major gaps in evidence are identified, which may hopefully stimulate new research. 635

These guidelines are the result of a close collaboration between physicians from many different areas of expertise: cardiology, vascular surgery, vascular medicine/angiology, neurology, radiology, etc., who have worked together with the aim of providing the medical community with the data to facilitate clinical decision 640 making in patients with PAD.

3. General aspects

This section covers the epidemiology of PAD and associated risk 645 factors, as well as aspects of diagnosis and treatment common to all specific vascular sites.

3.1 Epidemiology

The epidemiology of LEAD has been investigated in many 650 countries, including several in Europe. In a recent study in a population aged 60-90 years in Sweden, the prevalence of LEAD was 18% and that of intermittent claudication was 7%.³ Typically, one-third of all LEAD patients in the community are symptomatic. The prevalence of critical limb ischaemia (CLI) is very much less— 655 0.4% in those over 60 years of age in the Swedish study.³ The estimated annual incidence of CLI ranges from 500 to 1000 new cases per 1 million population, with a higher incidence among patients with diabetes.

The frequency of LEAD is strongly age related: uncommon 660 before 50 years, rising steeply at older ages. In a recent study in Germany the prevalence of symptomatic and asymptomatic LEAD in men aged 45-49 years was 3.0%, rising to 18.2% in those aged 70-75 years. Corresponding rates for women were 2.7% and 10.8%.⁴ Prevalence rates between men and women are 665 inconsistent. There is, however, some suggestion of an equilibration between the sexes with increasing age. Incidence rates are less often reported, but also show a strong relationship with age. In the Framingham Study, the incidence of intermittent claudication in men rose from 0.4 per 1000 aged 35-45 years to 6 per 670 1000 aged 65 years and older.⁵ The incidence in women was around half that in men, but was more similar at older ages.

The annual incidence of major amputations is between 120 and 500 per million in the general population, of which approximately equal numbers are above and below the knee. The prognosis for 675 such patients is poor. Two years following a below-knee amputation, 30% are dead, 15% have an above-knee amputation, 15% have a contralateral amputation, and only 40% have full mobility.⁶

Future trends in the epidemiology of LEAD are difficult to predict due to changes in risk factors in the population, especially 680 tobacco smoking and diabetes, and due to the increased survival from CAD and stroke, allowing LEAD to become manifest. Limited evidence on trends during the past few decades has suggested a decline in the incidence of intermittent claudication.

- In 50-year-old Icelandic men the incidence decreased from 1.7 per 1000 in 1970 to 0.6 per 1000 in 1984,⁷ whereas in the Framingham Study, the incidence decreased from 282 per 100 000 person-years in 1950–1959 to 225 per 100 000 person-years in 1990–1999.⁸ In the Rotterdam Study of elderly people over 55 years of age, a
- reduction in lumen diameter of the right internal carotid artery from 16% to 49% was found in 3%, whereas severe stenosis (≥50% reduction) was found in 1.4%.⁹ Likewise in the Tromso Study of the general population over 50 years of age, the prevalence of carotid stenosis was 4.2% in men, which was significantly
- higher than in women (2.7%) (P = 0.001).¹⁰ Minor degrees of stenosis are much more common. In the Cardiovascular Health Study in subjects >65 years of age, 75% of men and 62% of women had carotid plaques,¹¹ and in the Framingham Study in men aged 75 years, >40% had stenosis >10%.⁸
- Renal artery disease has been found frequently in post-mortem studies, but evidence on prevalence in the general population is limited. In the Cardiovascular Health Study of an elderly population with mean age 77 years, the prevalence of renal artery disease, defined as stenosis reducing arterial diameter by \geq 60% or occlu-
- sion, was 9.1% in men and 5.5% in women.¹² However, much information on the prevalence of renal artery disease has been derived from studies of patients undergoing coronary angiography or abdominal aortography in which the renal arteries have been imaged. A systematic review of such studies found that between
- 710 10% and 50% of patients had renal artery stenosis (RAS) depending on the risk group being examined.¹³ Owing to the selection of patients for such studies, the prevalences were likely to be much higher than those found in the general population.

Chronic symptomatic mesenteric artery disease is found rarely in clinical practice although at times is under/misdiagnosed. It accounts for only 5% of all intestinal ischaemic events and is often severe, even fatal. The prevalence of asymptomatic mesenteric artery disease in the general population is not well established. In patients with atherosclerotic disease at other sites,

720 atherosclerosis in the mesenteric arteries may be relatively common: in patients with LEAD and renal artery disease, 27% of patients had \geq 50% stenosis in a mesenteric artery.¹⁴

Atherosclerosis occurs much less frequently in the arteries of the upper extremity compared with the lower extremity. The subclavian artery is often affected. In a study using data from four

- clavian artery is often affected. In a study using data from four cohorts in the USA, the prevalence of subclavian artery stenosis in the general population was 1.9%, with no significant difference between the sexes.¹⁵ Prevalence increased with age from 1.4% in those <50 years of age to 2.7% in those >70 years. Subclavian
- stenosis was defined in this study as an inter-arm pressure difference of \geq 15 mmHg, but, using angiography as the gold standard, the sensitivity of this definition has been shown to be only ~50% and specificity 90%. Thus the true prevalence of subclavian artery stenosis may be much higher than that observed in the cohorts. The majority of these cases are asymptomatic.
- Given the common aetiology of peripheral atherosclerosis occurring at different vascular sites, the presence of disease at one site increases the frequency of symptomatic and asymptomatic disease at another. The degree of concordance observed between
- 740 sites is, however, dependent on the methods of diagnosis and on the selected population. From a clinical perspective, such findings

indicate the need for a heightened awareness of the possibility of atherosclerotic disease occurring at sites other than the presenting one. This is especially so in the elderly in whom the degree of overlap of CAD, cerebrovascular disease, and LEAD is particularly 745 high.

3.2 Risk factors

Risk factors for PAD are similar to those important in the aetiology of CAD and are the typical risk factors for atherosclerotic disease. 750 These include the traditional risk factors: smoking, dyslipidaemia, diabetes mellitus, and hypertension. However, for some peripheral artery sites the evidence linking these factors to the development of disease is limited. Also, specific risk factors could be more important for the development of disease at certain sites, but 755 there are few comparative studies.

In LEAD, cigarette smoking has been shown consistently in several epidemiological studies to be an important risk factor and to be dose dependent.^{16,17} Smoking would appear to be a stronger risk factor for LEAD than for CAD and, in most 760 studies, patients with claudication have had a history of smoking at some point in their lives. Smoking cessation is associated with a rapid decline in the incidence of claudication, which equates to that in non-smokers after 1 year of stopping.⁷ Diabetes mellitus is the other risk factor especially important in the development 765 of LEAD. This is certainly true for severe disease, notably gangrene and ulceration, but for intermittent claudication the strength of the association with diabetes may be comparable with that for coronary heart disease. The association of diabetes with LEAD is inconsistent on multivariable analysis, which includes other risk 770 factors, but it appears that the duration and severity of diabetes affect the level of risk.^{16,17}

Most epidemiological studies show an association between hypertension and the presence of LEAD, although interpretation of such findings is difficult because blood pressure is a component 775 in the definition of disease [the ankle–brachial index (ABI)] and may also affect the degree of ischaemia and the occurrence of symptoms. However, no association has been found between increased blood pressure and claudication. In contrast, in the Limburg PAOD study, hypertension was associated with an 780 increased relative risk of 2.8 for LEAD¹⁸ and in the Rotterdam Study a low ABI (<0.90) was associated with both increased systolic and diastolic blood pressure.¹⁹

Most epidemiological studies have found that high total cholesterol and low high-density lipoprotein (HDL) cholesterol are independently related to an increased risk of LEAD. In the US Physicians Health Study, the ratio of total/HDL cholesterol was the lipid measure most strongly related to disease.²⁰

For other factors associated with CVD, such as obesity, alcohol consumption, and plasma homocysteine levels, the associations 790 with LEAD have been inconsistent. In recent years, particular interest in haemostatic, rheological, and inflammatory markers, such as plasma fibrinogen and C-reactive protein,²⁰ has led to studies that have shown independent associations with both the prevalence and incidence of LEAD, although whether such associ- 795 ations are primarily the cause or the effect is not clearly known. Currently genetic factors and many other novel biomarkers are being studied.

- In general, the risk factors for carotid stenosis are similar to those for LEAD, although smoking, while commonly associated with carotid disease, is not so dominant as with LEAD. Several population-based studies have found in both symptomatic and asymptomatic disease that the classic risk factors of smoking, high low-density lipoprotein (LDL) cholesterol, low HDL choles-
- 805 terol, hypertension, and diabetes mellitus are associated with higher risk in both men and women irrespective of age.^{9–11} The risk factors for carotid artery disease, however need to be distinguished from those for ischaemic stroke, which is not necessarily related to stenosis in the carotid arteries.
- Likewise, for atheromatous renal artery disease the pathogenesis is similar to that seen in other vascular sites and, although the evidence is limited, would appear to be associated with typical cardiovascular risk factors.²¹ These include pre-existing high blood pressure in which the hypertension is not necessarily a compli-
- 815 cation but may be a cause of the RAS and may partly explain why in many patients revascularization may not lead to a reduction in blood pressure.

In chronic mesenteric artery disease, the atheromatous lesions normally occur in the proximal segments of the splanchnic arteries.

- 820 The frequency of diffuse atherosclerosis has not been well described but would appear to occur mostly in patients with endstage renal disease (ESRD) or diabetes. The classic cardiovascular risk factors appear to be important, although hypocholesterolaemia (rather than hypercholesterolaemia) may be a presenting
- 825 finding due to a patient's chronic malnourished state. Significant associations were found between both increasing age and higher systolic blood pressure with the presence of upper extremity artery disease (UEAD).¹⁵ Compared with never smokers, the risks were increased in current and past smokers,
- and the odds ratio (OR) of 2.6 for current smokers was the highest of any risk factor, perhaps mirroring that found for LEAD. While a higher HDL cholesterol level appeared to be protective, surprisingly no association was found between total cholesterol and subclavian stenosis. Diabetes mellitus was also not related, although in another study the prevalence of UEAD
- was found to be slightly higher in diabetic compared with nondiabetic patients.²² Interestingly, in the four cohort study, LEAD, compared with CAD and cerebrovascular disease, was much more strongly related to UEAD.¹⁵
- 840

3.3 General diagnostic approach

3.3.1 History

History of risk factors and known co-morbidities is mandatory. Hypertension, dyslipidaemia, diabetes mellitus, smoking status, as well as history of CVD must be recorded. Medical history should include a review of the different vascular beds and their specific symptoms:

- Family history of CVD.
- Symptoms suggesting angina.
 - Any walking impairment, e.g. fatigue, aching, cramping, or pain with localization to the buttock, thigh, calf, or foot, particularly when symptoms are quickly relieved at rest.
- Any pain at rest localized to the lower leg or foot and its association with the upright or recumbent positions.

- Any poorly healing wounds of the extremities.
- Upper extremity exertional pain, particularly if associated with dizziness or vertigo.
- Any transient or permanent neurological symptom.
- History of hypertension or renal failure.
- Post-prandial abdominal pain and diarhoea, particularly if related to eating and associated with weight loss.
- Erectile dysfunction.

This cannot be an exhaustive list, and a review of symptoms should include all domains. It is important to emphasize that history is a cornerstone of the vascular evaluation.

One should remember that many patients, even with advanced disease, will remain asymptomatic or report atypical symptoms.

3.3.2 Physical examination

Although physical examination alone is of relatively poor sensitivity, specificity, and reproducibility, a systematic approach is mandatory. It must include at least:

- Measurement of blood pressure in both arms and notation of 875 inter-arm difference.
- Auscultation and palpation of the cervical and supraclavicular fossae areas.
- Palpation of the pulses at the upper extremities. The hands must be carefully inspected.
- Abdominal palpation and auscultation at different levels including the flanks, periumbilical region, and the iliac regions.
- Auscultation of the femoral arteries at the groin level.
- Palpation of the femoral, popliteal, dorsalis pedis, and posterior tibial sites.
- The feet must be inspected, and the colour, temperature, and integrity of the skin, and the presence of ulcerations recorded.
- Additional findings suggestive of LEAD, including calf hair loss and skin changes, should be noted.

Beyond their diagnostic importance, clinical signs could have a prognostic value. A meta-analysis published in 2008 emphasized the prognostic value of carotid bruit.²³ People with carotid bruits have twice the risk of myocardial infarction and cardiovascular death compared with those without. This predictive value can be extended to other clinical signs, such as femoral bruit, pulse abnormality in the lower extremity, or inter-arm blood pressure asymmetry. All of these abnormalities can be an expression of subclinical vascular disease.

3.3.3 Laboratory assessment

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The aim of the laboratory assessment is to detect major risk factors of CVD. The assessment should be performed according to the ESC Guidelines on Cardiovascular Disease Prevention²⁴ and the ESC/EAS Guidelines for the Management of Dyslipidaemias.²⁵ ₉₀₅

3.3.4 Ultrasound methods

3.3.4.1 Ankle-brachial index

The ABI is a strong marker of CVD and is predictive of cardiovascular events and mortality. Low ABI values (<0.90) are predictive 910 of atherosclerosis, such as CAD and carotid artery disease. A reduced ABI has been associated in several studies with an

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increased risk of cardiovascular morbidity and mortality.²⁶ Also a very high ABI (>1.40) in relation to stiffened arteries is associated

- with increased mortality.²⁷ Recently, the ABI has been shown to be a valid method of cardiovascular risk assessment in diverse ethnic groups, independent of traditional and novel risk factors, as well as other markers of atherosclerosis such as the coronary artery calcium score.²⁷ ABI is recommended as an office measurement
- in selected populations considered at high risk of CVDs. When 920 performed with a handheld Doppler device, the measurement remains inexpensive and minimally time consuming.

The use of ABI to diagnose LEAD is discussed in Section 4.5.2.1.

3.3.4.2 Duplex ultrasound 925

> Duplex ultrasound (DUS) is now widely available for the screening and diagnosis of vascular lesions. Initially, with continuous wave Doppler, severe stenoses were identified and quantified mainly by the peak systolic velocities. Nowadays, DUS includes B-mode

echography, pulsed-wave Doppler, colour Doppler, and power 930 Doppler in order to detect and localize vascular lesions and guantify their extent and severity.

By detecting subclinical artery disease, DUS provides relevant information regarding cardiovascular risk assessment. B-mode ultrasound is also a robust technique for the measurement of

- 935 the intima-media thickness (IMT), which has been studied (mostly in the carotid arteries) and validated in several epidemiological and interventional studies as a marker of atherosclerotic burden in individuals and a predictor of cardiovascular morbidity
- and mortality. Further, DUS allows a complete vascular evaluation 940 of the different beds and is often the first step in the clinical management. New techniques, such as B-flow imaging or live threedimensional (3D) echography, as well as the use of ultrasound contrast agents, will further improve the performance of DUS.

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3.3.5 Angiography

In the past, digital subtraction angiography (DSA) was the gold standard of vascular imaging. Given its invasive characteristics, this method has now been replaced by other effective non-invasive diagnostic methods and is used almost exclusively during endovascular procedures.

3.3.6 Computed tomography angiography

The introduction of multidetector computed tomography (MDCT) 955 has shortened the examination time and reduced motion and respiration artefacts while imaging the vessels and organs. The use of computed tomography angiography (CTA) is not recommended for screening purposes due to the high doses of radiation used, potential contrast nephrotoxicity, and the lack of data demonstrat-960 ing the effect of screening with CT.

When CTA is used for diagnostic purposes, nephrotoxicity can be limited by minimizing the volume of contrast agents and ensuring adequate hydration before and after imaging. The potential benefit of acetylcysteine to limit nephrotoxicity is uncertain.

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3.3.7 Magnetic resonance angiography

High-performance scanning is used during magnetic resonance angiography (MRA) with a high signal-noise ratio and rapid data

acquisition. Morphological and functional studies require at least 970 a 1.0 Tesla system. In order to increase the resolution, special phased-array surface coils are placed directly on the body, which provide a homogeneous magnetic field over a large area.

Absolute contraindications include cardiac pacemakers, implantable cardioverter defibrillators, neurostimulators, cochlear 975 implants, first-trimester pregnancy, and severe renal failure [glomerular filtration rate (GFR) < 30 mL/min per 1.73 m²]. Pacing systems suitable for magnetic resonance imaging (MRI) have been developed. Claustrophobia, metallic foreign objects, and second- or third-trimester pregnancy are regarded as relative 980 contraindications.

Time-of-flight angiography and phase-contrast angiography, without intravenous contrast, can be used to image the vascular bed. Development of the 'Angiosurf' and 'Bodysurf' techniques^{28,29} has been a breakthrough in imaging. Based on the 'Angiosurf' MRA 985 approach, a fairly comprehensive combined protocol can be used, which accomplishes the depiction of the head, thoracic, and all peripheral arteries from the carotids to the ankles.^{30,31}

Detailed descriptions of CTA and MRA are provided in Appendix 1 (available online at www.escardio.org/guidelines). 990

3.4 Treatment—general rules

Patient management should include lifestyle modification, focusing on smoking cessation, daily exercise (30 min/day), 995 normal body mass index ($\leq 25 \text{ kg/m}^2$), and a Mediterranean diet.²⁴ Pharmacological treatment can be added for blood pressure control and a lipid-lowering treatment to achieve LDL cholesterol <2.5 mmol/L (100 mg/dL) with an option of <1.8 mmol/L (<70 mg/dL) if feasible. In diabetic patients, 1000 glucose control should be obtained, with the target glycated haemoglobin (Hb_{A1c}) <7%. Site-dependent therapy and revascularization strategy are discussed in the respective sections. It must be emphasized that the management of patients with PAD should always be decided after multidisciplinary discussion, also including 1005 (depending on lesion site) specialists beyond the area of cardiovascular medicine, e.g. neurologists or nephrologists.

3.4.1 Smoking cessation

1010 Smoking is an important risk factor for PAD.³² In the general population smoking increased the risk of LEAD between twoand six-fold.¹⁶ Current smokers with LEAD also have an increased risk of amputation, and are at increased risk of postoperative complications and mortality.³³ Smokers should be 1015 advised to quit smoking and be offered smoking cessation programmes. Nicotine replacement therapy and/or bupropion or varenicline can facilitate cessation in patients with a high level of nicotine dependence, which can be estimated by the Fagerström's questionnaire or biomarkers such as exhaled carbon monoxide concentrations.³⁴ All three medications are safe to use in patients with CVD.³⁵

3.4.2 Lipid-lowering drugs

Statins reduce the risk of mortality, cardiovascular events, and 1025 stroke in patients with PAD with and without CAD. In the

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Heart Protection Study, 6748 participants had PAD; at 5-year follow-up, simvastatin caused a significant 19% relative reduction and a 6.3% absolute reduction in major cardiovascular events independently of age, gender, or serum lipid levels.³⁶ All patients with

PAD should have their serum LDL cholesterol reduced to <2.5 mmol/L (100 mg/dL), and optimally to <1.8 mmol/L (<70 mg/dL), or >50% LDL cholesterol reduction when the target level cannot be reached.24,25

The Antithrombotic Trialists' Collaboration meta-analysis com-

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3.4.3 Antiplatelet and antithrombotic drugs

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bined data from 42 randomized studies of 9706 patients with intermittent claudication and/or peripheral arterial bypass or angioplasty. The incidence of vascular death, non-fatal myocardial infarction, and non-fatal stroke at follow-up was significantly decreased, by 23%, by antiplatelet drugs.³⁷ Low-dose aspirin (75–150 mg daily) was at least as effective as higher daily doses. The efficacy of clopidogrel compared with aspirin was 1045 studied in the randomized Clopidogrel versus Aspirin in Patients at Risk for Ischaemic Events (CAPRIE) trial, including a subgroup of 6452 patients with LEAD.³⁸ At 1.9-year follow-up, the annual combined incidence of vascular death, non-fatal myocardial infarction, and non-fatal stroke in the LEAD group was 3.7% and 4.9%, 1050 respectively, in the clopidogrel and aspirin groups, with a significant 23.8% decrease with clopidogrel. These benefits appeared

higher than in patients enrolled for CAD or stroke. The small benefits of dual antiplatelet therapy do not justify its recommen-

dation in patients with LEAD due to an increased bleeding 1055 risk.^{39,40}

3.4.4 Antihypertensive drugs

Arterial hypertension in patients should be controlled adequately according to the current ESC/European Society of Hypertension 1060 guidelines.⁴¹ In general, target blood pressures of <140/ 90 mmHg are recommended, and \leq 130/80 mmHg in patients with diabetes or chronic kidney disease. However, the latter target has recently been contested.⁴²

- Treatment with angiotensin-converting enzyme (ACE) inhibitors 1065 has shown a beneficial effect beyond a blood pressure decrease in high-risk groups. In the Heart Outcomes Prevention Evaluation (HOPE) trial, ACE inhibitor treatment with ramipril significantly reduced cardiovascular events by 25% in patients with sympto-
- matic PAD without known low ejection fraction or heart 1070 failure.⁴³ The ONTARGET trial showed equivalence of telmisartan to ramipril in these patients.44

Importantly, B-blockers are not contraindicated in patients with LEAD. A meta-analysis of 11 randomized controlled studies found

- 1075 that β-blockers did not adversely affect walking capacity or symptoms of intermittent claudication in patients with mild to moderate LEAD.⁴⁵ At 32-month follow-up of 490 patients with LEAD and prior myocardial infarction, β-blockers caused a 53% significant independent relative decrease in new coronary events.⁴⁶ Consider-
- ing the cardioprotective effects of a low-dose, titrated β -blocker 1080 regimen in the perioperative setting, β -blockers are recommended in patients scheduled for vascular surgery according to the ESC guidelines.47

Recommendations in patients with PAD: general treatment

Recommendations	Class ^a	Level ^b	Ref ^c	
All patients with PAD who smoke should be advised to stop smoking.	I	В	48	1090
All patients with PAD should have their LDL cholesterol lowered to <2.5 mmol/L (100 mg/dL), and optimally to <1.8 mmol/L (70 mg/dL), or \geq 50% when the target level cannot be reached.	I	C₫	-	1095
All patients with PAD should have their blood pressure controlled to ≤140/90 mmHg.	I	A	41	1100
β-Blockers are not contraindicated in patients with LEAD, and should be considered in the case of concomitant coronary artery disease and/or heart failure.	lla	В	46, 47	1105
Antiplatelet therapy is recommended in patients with symptomatic PAD.	I	Cq	37	
In patients with PAD and diabetes, the HbA1c level should be kept at ≤6.5%.	I	Cq	-	1110
In patients with PAD, a multidisciplinary approach is recommended to establish a management strategy.	I	C₫	-	1115
ass of recommendation. vel of evidence.				
ferences. idence is not available for all sites ommendations specific for the vas tions.				112

Hb_{A1}c = glycated haemoglobin; LDL = low-density lipoprotein; LEAD = lower extremity artery disease; PAD = peripheral artery disease.

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4. Specific vascular areas

4.1 Extracranial carotid and vertebral artery disease

4.1.1 Carotid artery disease

4.1.1.1 Definition and clinical presentations

In the Western world, ischaemic stroke has a major public health impact as the first cause of long-term disability and the third leading cause of death. Stroke mortality ranges from 10% to 30%, and survivors remain at risk of recurrent neurological and 1135 cardiac ischaemic events. The risk of stroke and transient ischaemic attacks (TIAs), defined in most studies as transient neurological deficits usually lasting 1-2h and no longer than 24h, increases with age. Major risk factors for stroke include hypertension, hypercholesterolaemia, smoking, diabetes, cerebrovascular 1140 disease, atrial fibrillation, and other cardiac conditions that increase the risk for embolic complications. Large artery atherosclerosis, and specifically internal carotid artery stenosis, accounts for ${\sim}20\%$ of all ischaemic strokes.⁴⁹ Carotid artery stenosis is con-

- 1145 sidered symptomatic in the presence of TIA or stroke affecting the corresponding territory within the previous 6 months.^{50,51} In the vast majority of cases, carotid artery stenosis is caused by atherosclerosis. Rare aetiologies include radiation therapy, vasculitis, dissection, or fibromuscular dysplasia.
- 1150 For the purpose of these guidelines, the term carotid artery stenosis refers to a stenosis of the extracranial portion of the internal carotid artery, and the degree of stenosis is according to the NASCET criteria (see online Appendix 2).

In the North American Symptomatic Carotid Endarterectomy

- 1155 Trial (NASCET), the risk of recurrent ipsilateral stroke in patients with symptomatic carotid artery stenosis treated conservatively was 4.4% per year for 50–69% stenosis and 13% per year for >70% stenosis.⁵² In patients with asymptomatic carotid artery stenosis >60%, the risk of stroke is $\sim 1-2\%$ per year.^{53,54} However,
- 1160 the risk may increase to 3–4% per year in elderly patients or in the presence of contralateral carotid artery stenosis or occlusion, evidence of silent embolization on brain imaging, carotid plaque heterogeneity, poor collateral blood supply, generalized inflammatory state, and associated coronary or peripheral artery disease.^{1,52}
- ¹¹⁶⁵ Currently there are indications that the risk of stroke in patients with asymptomatic carotid artery disease is lower due to better medical treatment.^{55,56}

4.1.1.2 Diagnosis

1170 4.1.1.2.1 Clinical evaluation

The decision to revascularize patients with carotid artery stenosis is based on the presence of signs or symptoms related to the affected carotid artery, the degree of internal carotid artery stenosis, and on patient age, gender, co-morbidities, and life expectancy.

1175 Additional factors such as the presence of silent brain infarction in the corresponding territory, microembolization on intracranial Doppler, or the degree of stenosis progression may also be taken into account.

Neurological evaluation is essential to differentiate asymptomatic and symptomatic patients. All patients with neurological complaints should be seen as soon as possible by a neurologist since it may be challenging to determine whether symptoms are related to a carotid artery stenosis. Manifestations of carotid artery disease may be divided into hemispheric and/or ocular.

- Hemispheric (cortical) ischaemia usually consists of a combination of weakness, paralysis, numbness, or tingling (all affecting the same side of the body) and contralateral to the culprit carotid artery. Neuropsychological symptoms may also be present and may include aphasia if the dominant hemisphere (usually left) is affected,
- 1190 or neglect if the non-dominant hemisphere (usually the right, even in most left-handed individuals) is affected. Emboli to the retinal artery may cause temporary or permanent partial or total blindness in the ipsilateral eye. A temporary ocular deficit is called amaurosis fugax. While neurological symptoms of carotid disease
- 1195 are usually caused by distal embolization, they may seldom be due to cerebral hypoperfusion, either transient ('low-flow TIA') or permanent (haemodynamic stroke).

4.1.1.2.2 Imaging

Urgent imaging of the brain and supra-aortic vessels is mandatory in all patients presenting with TIA or stroke. While CT scan is 1200 widely available and allows for a differentiation between ischaemic and haemorrhagic stroke, MRI is more sensitive in the detection of brain ischaemia.

The risk of recurrent TIA or stroke in the first month is 10–30%.⁵⁷ In patients with carotid artery stenosis, imaging conveys 1205 important information such as the degree of carotid artery stenosis, carotid plaque morphology, the presence of intracranial disease, intracranial collateral circulation, asymptomatic embolic events, or other intracranial pathologies.

DUS is commonly used as the first step to detect extracranial 1210 carotid artery stenosis and to assess its severity. The peak systolic velocity measured in the internal carotid artery is the primary variable used for this purpose; secondary variables include the enddiastolic velocity in the internal carotid artery as well as the ratio of peak systolic velocity in the internal carotid artery as well as the ratio of peak systolic velocity in the internal carotid artery to that in 1215 the common carotid artery.⁵⁸ Although DUS evaluation may be hampered by severe plaque calcifications, tortuous vessels, tandem lesions, and slow turbulent flow in subtotal stenoses, this imaging modality allows for a reliable estimation of the degree of the stenosis as well as for the assessment of plaque morphology 1220 in the hands of an experienced investigator.

The advantages of CTA and MRA include the simultaneous imaging of the aortic arch, the common and internal carotid arteries in their totality, the intracranial circulation, as well as the brain parenchyma. MRA is more time-consuming than CTA but does not 1225 expose patients to radiation, and the used contrast agents are far less nephrotoxic. CTA offers excellent sensitivity and specificity for the detection of carotid artery stenosis; however, the presence of severe plague calcification may lead to overestimation of the degree of stenosis. In a systematic review and meta-analysis, no 1230 major difference was found between DUS, MRA, and CTA for the detection of a significant carotid artery stenosis.⁵⁹ In order to improve the accuracy of the diagnosis, the use of two imaging modalities prior to revascularization is suggested. DSA may be required for diagnostic purposes only in selected cases (e.g. discordant non- 1235 invasive imaging results, additional intracranial vascular disease). In patients with severe asymptomatic carotid artery stenosis, imaging of the brain to detect asymptomatic embolic events and a transcranial Doppler for emboli detection may be considered.

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Recommendation for evaluation of carotid artery stenosis

	Recommendations	Class ^a	Level ^b	Ref ^c		1245
	DUS, CTA, and/or MRA are indicated to evaluate carotid artery stenosis.	I	A	59		
^a Class of recommendation. ^b Level of evidence. ^c Reference. CTA = computed tomography angiography; DUS = duplex ultrasonography;						1250

MRA = magnetic resonance angiography.

4.1.1.3 Treatment modalities 1255

4.1.1.3.1 Medical therapy

The overall benefit of aspirin to prevent cardiovascular events in patients with atherosclerosis have been presented earlier (Section 3.4.3). Although, the use of antiplatelet agents has not been specifically addressed in patients with carotid artery disease

- 1260 (i.e. carotid plaques), low-dose aspirin (or clopidogrel in case of aspirin intolerance) should be administered to all patients with carotid artery disease irrespective of symptoms. The effectiveness of statins in patients with symptomatic cerebrovascular disease is
- well proven, irrespective of the initial cholesterol concentration. 1265 The Stroke Prevention by Aggressive Reduction in Cholesterol Levels (SPARCL) study evaluated the results of high-dose atorvastatin (80 mg/day) vs. placebo in 4731 patients with TIA or stroke. Patients allocated to atorvastatin had a significant 26% rela-
- tive risk reduction of the primary endpoint of fatal and non-fatal 1270 stroke at 5 years.⁶⁰ Among 1007 patients with carotid artery stenosis enrolled in the trial, the benefit of statin therapy was even more pronounced, with a 33% reduction of stroke, a 43% reduction of major coronary events, and a 56% reduction of carotid revascularization procedures at 5 years.⁶¹ 1275

4.1.1.3.2 Surgery

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The benefits of carotid endarterectomy (CEA) over medical management in randomized trials were conveyed by low perioperative complication rates [e.g. a stroke and death rate of 5.8% in NASCET⁵² and of 2.7% in the Asymptomatic Carotid Athero-

sclerosis Study (ACAS)⁵³] achieved by high-volume surgeons in low-risk patients.

Temporary interruption of cerebral blood flow during CEA can cause haemodynamic neurological deficits. This can potentially be

- 1285 avoided by using a shunt. Currently there is insufficient evidence to support or refute the use of routine or selective shunting as well as perioperative neurological monitoring during CEA. As suggested by a Cochrane review of seven trials, CEA using a patch (either prosthetic or vein based) may reduce the risk of restenosis and neurologi-
- 1290 cal events at follow-up compared with primary closure.⁶² A more recent randomized trial confirmed the lower restenosis rate associated with the patch, but could not find any difference in perioperative complications.⁶³ Usually, CEA is performed using a longitudinal arteriotomy. However, CEA with arterial eversion implies a transverse
- 1295 arteriotomy and reimplantation of the internal carotid artery on the common carotid artery. A Cochrane analysis on this subject suggested that CEA with eversion may be associated with a lower risk of (sub)acute occlusion and restenosis than conventional CEA. but no difference in clinical events was detected.64
- 1300 For decades it has been debated whether local anaesthesia is superior to general anaesthesia for CEA. The randomized General Anaesthesia versus Local Anaesthesia for Carotid Surgery (GALA) trial including 3526 patients showed no difference in terms of perioperative death, stroke, or myocardial infarction 1305 between general (4.8%) and local (4.5%) anaesthesia.⁶⁵

All patients undergoing CEA should receive perioperative medical management according to proper cardiovascular risk assessment. Low-dose aspirin is efficacious to reduce perioperative stroke.^{37,52,54,66} There is no clear benefit of dual therapy or high-

1310 dose antiplatelet therapy in patients undergoing CEA. Technical aspects of CEA are addressed in Appendix 2.

4.1.1.3.3 Endovascular techniques

Carotid artery stenting (CAS) is a revascularization option less invasive than CEA. It is performed under local anesthaesia, avoids neck dissection with the consequent risk of peripheral nerve damage, and 1315 is less painful. Although patients at high risk for surgery are not well defined, CAS is frequently advocated for patients at increased cardiopulmonary risk or with unfavourable neck anatomy, restenosis after CEA, prior neck dissection or radiation therapy, as well as in the presence of carotid artery stenosis difficult to access (i.e. high 1320 internal carotid or low common carotid artery lesions).

The optimal anticoagulation regimen for CAS remains unknown. Periprocedure unfractionated heparin is commonly used. Dual antiplatelet therapy with aspirin and clopidogrel (or ticlopidine) is recommended. Two small, randomized trials comparing aspirin 1325 alone with double antiplatelet therapy for CAS were terminated prematurely due to high rates of stent thrombosis and neurological events in the aspirin-alone group.67,68

In patients with proven intolerance to dual antiplatelet therapy, CEA should be preferred to CAS. Newer antiplatelet agents such 1330 as prasugrel or ticagrelor have not yet been adequately tested in CAS.

4.1.1.3.4 Operator experience and outcomes of carotid artery stenting While comparing the results of CAS and CEA, it should be acknowledged that CAS gained maturity more recently than CEA, and that 1335 the endovascular technique is evolving rapidly. Overall, available evidence supports the notion that experience does play a major role in CAS outcomes. The benefit is probably conveyed by optimal procedure management and appropriate patient selection. In this respect, several CAS vs. CEA trials have been criticized for the insuf-1340 ficient endovascular experience required and for the possibility of treating patients with CAS under proctoring conditions.⁶⁹

More detailed information on the importance of operator experience in CAS is provided in Appendix 2.

4.1.1.3.5 Embolic protection devices

The use of embolic protection devices (EPDs) during CAS remains controversial. At present, only two very small, randomized studies have evaluated CAS with vs. without EPDs, and failed to prove an improved clinical outcome with the use of the devices.^{70,71}

Opposing these results, two systematic reviews showed a 1350 reduction in neurological events associated with protected CAS.^{72,73} A benefit from EPDs was also suggested from a large-scale prospective registry documenting an in-hospital death or stroke rate of 2.1% among 666 patients undergoing CAS with 1355 adjunctive EPD and of 4.9% in the group of patients (n = 789) treated without EPDs (P = 0.004).⁷⁴ In the same study, the use of EPDs was identified in multivariable analysis as an independent protective factor for this endpoint (adjusted OR 0.45, P =0.026). Importantly, the complication rate associated with the 1360 use of EPD appears to be low (<1%).⁷⁵

In contrast, secondary analyses from two randomized CAS vs. CEA trials reported a lack of benefit from EPD use during CAS. In the SPACE trial, the rate of 30-day ipsilateral stroke or death after CAS was 8.3% among 145 patients treated with EPDs and 1365 6.5% in 418 patients treated without EPDs (P = 0.40).⁷⁶ In a substudy of the ICSS trial, new diffusion-weighted MRI lesions after CAS were observed in 38 (68%) of 56 patients who had stenting with EPDs and in 24 (35%) of 68 patients who had unprotected

stenting [OR 3.28, 95% confidence interval (CI) 1.50-7.20; P = 0.003].⁷⁷ Importantly, the use of EPDs in both trials was left to 1370 the discretion of the operator. The best results for CAS so far in randomized trials-for both symptomatic and asymptomatic patients-have been obtained in studies that mandated embolic

protection with a single device and in which operators were trained in the use of the specific device [Stenting and Angioplasty 1375 with Protection in Patients at High Risk for Endarterectomy (SAP-PHIRE)⁷⁸ and CREST,⁷⁹ as detailed below]. Finally, recent registry data suggest that proximal occlusion systems may be useful in embolic protection.⁸⁰

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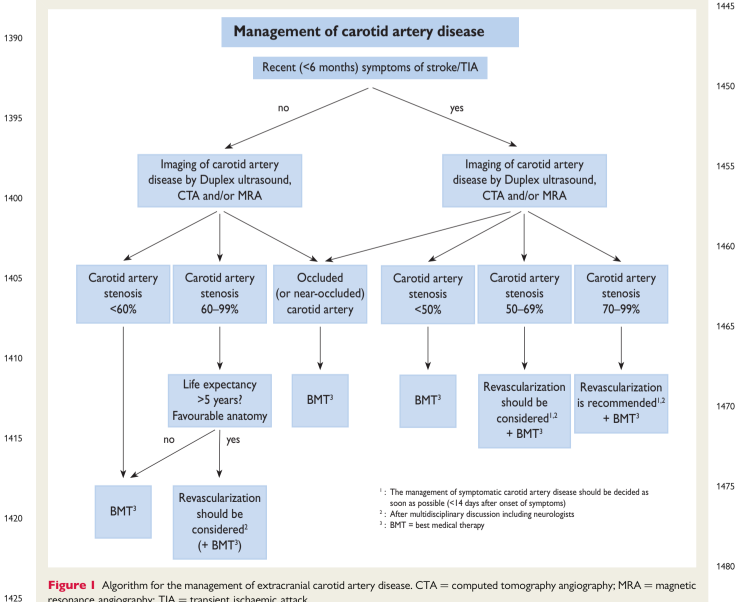
4.1.1.4 Management of carotid artery disease

The management of carotid artery disease is summarized in Figure 1.

Recommendations for embolic protection in patients undergoing CAS

Recommendations	Class ^a	Level ^b	R ef ^c		1430
Dual antiplatelet therapy with aspirin and clopidogrel is recommended for patients undergoing CAS.	I	В	67, 68		
The use of EPDs may be considered in patients undergoing CAS.	ШЬ	В	73		1435
^a Class of recommendation.				-	
² Level of evidence. References.					1440

CAS = carotid artery stenting; EPD = embolic protection device.



resonance angiography; TIA = transient ischaemic attack.

4.1.1.4.1 Asymptomatic carotid artery disease 4.1.1.4.1.1 Surgery

- 1485 A total of 5233 patients with asymptomatic carotid artery disease were enrolled in randomized multicentre trials comparing CEA with medical management.^{53,54,66,81} After 4657 patientyears of follow-up, the randomized Asymptomatic Carotid Atherosclerosis Study (ACAS) estimated the 30-month risk of
- ¹⁴⁹⁰ ipsilateral stroke in the case of carotid artery stenosis >60%at 5.1% for patients who underwent CEA in addition to best medical therapy (at that time) vs. 11.0% for those with best medical therapy alone.⁵³ The Asymptomatic Carotid Surgery Trial (ACST) randomized 3120 asymptomatic patients to either
- immediate CEA or indefinite deferral of CEA.⁵⁴ The 5-year risks were 6.4% vs. 11.8% for all strokes (absolute risk reduction 5.4%, P = 0.0001), 3.5% vs. 6.1% for fatal or disabling stroke (absolute risk reduction 2.6%, P = 0.004), and 2.1% vs. 4.2% for fatal strokes (absolute risk reduction 2.1%, P = 0.006), respect-
- 1500 ively. Combining perioperative events and strokes, net risks were 6.9% vs. 10.9% at 5 years (gain 4.1%, 2.0–6.2) and 13.4% vs. 17.9% at 10 years (gain 4.6%, 1.2–7.9).⁶⁶ Medication was similar in both groups; throughout the study, most patients were on antithrombotic and antihypertensive therapy. Net benefits
- 1505 were significant irrespective of the use of lipid-lowering therapy, for men and women under the age of 75 years at entry. In the three trials, the benefit was greater in men than in women, but the number of women enrolled was low.
- It can be concluded that CEA is beneficial in asymptomatic patients (especially men) between 40 and 75 years of age with >60% stenosis, if their life expectancy is >5 years and operative mortality <3%.^{66,70–77,79,81} However, the absolute benefit of revascularization in terms of stroke prevention is small (1–2%

per year), and those trials were performed prior to extensive 1540 use of statins. Therefore, the benefit of revascularization on top of optimal medical management should be reassessed.

4.1.1.4.1.2 Endovascular therapy

The results of eight CAS registries enrolling >1000 patients have been published recently (*Table 3*).⁸² The registries included $>20\ 000\ patients at high surgical risk, mainly asymptomatic. Pre$ and post-procedure neurological assessment and blinded event adjudication were required in most studies. Overall, the studies demonstrated that death and stroke rates with CAS are in the rangeexpected in current recommendations for CEA even in patients athigh surgical risk, and that CAS results tend to improve over time.1545

So far, the randomized evidence for CAS in asymptomatic patients is limited. While no study has compared endovascular treatment with medical therapy, two trials (SAPPHIRE and CREST) comparing CAS vs. CEA have also enrolled asymptomatic patients (for details see Section 4.1.1.4.2.2).

4.1.1.4.2 Symptomatic carotid artery disease

It should be emphasized that neurological assessment and appropriate treatment should be proposed as soon as possible after the index event. At a very minimum patients need to be seen and treated within 2 weeks, with important benefit of instituting medical treatment⁸⁸ and performing revascularization as soon as possible after the onset of symptoms.^{89,90}

4.1.1.4.2.1 Surgery

Pooled data from the NASCET, the European Carotid Surgery Trial (ECST), and the Veterans Affairs Trial included >35 000 patient-years of follow-up in patients (28% women) with symptomatic disease.^{50,51,91,92} CEA increased the 5-year risk of ipsilateral ischaemic stroke over medical therapy alone in patients with ¹⁵⁷⁰

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Table

le 3	Thirty-day	v event rates in	carotid artery	stenting regist	ries enrolling	>1000 patients
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Name	Year	N	Industry sponsored	Surgical high-risk	EPD	Sympt patients	Neurologistª	CEC	D/S	D/S/MI	D/S sympt	D/S asympt
CAPTURE ⁸³	2007	3500	Yes	Yes	Mandatory	14%	Yes	Yes	5.7%	6.3%	10.6%	4.9%
CASES-PMS ⁸⁴	2007	1493	Yes	Yes	Mandatory	22%	Yes	Yes	4.5%	5.0%	NA	NA
PRO-CAS ⁸⁵	2008	5341	No	No	75%	55%	70%	No	3.6% ⁵	NA	4.3% ⁵	2.7% [♭]
SAPPHIRE-W78	2009	2001	Yes	Yes	Mandatory	28%	No ^c	Yes	4.0%	4.4%	NA	NA
Society for Vascular Surgery ⁸⁶	2009	1450	No	No	95%	45%	No	No	NA	5.7%	NA	NA
EXACT ⁸⁷	2009	2145	Yes	Yes	Mandatory	10%	Yes	Yes	4.1%	NA	7.0%	3.7%
CAPTURE-287	2009	4175	Yes	Yes	Mandatory	13%	Yes	Yes	3.4%	NA	6.2%	3.0%
Stabile et al. ⁸⁰	2010	1300	No	No	Mandatory	28%	Yes	No	1.4%	NA	3.0%	0.8%

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^aIndependent pre- and post-procedural assessment by a neurologist.

^bIn-hospital events.

^cNeurological assessment performed by stroke-scale-certified staff member.

CAPTURE = Carotid ACCULINK/ACCUNET Post Approval Trial to Uncover Rare Events; CASES-PMS = Carotid Artery Stenting with Emboli Protection Surveillance Study; CEC = clinical event committee adjudication; D = death; EPD = embolic protection device; EXACT = Emboshield and Xact Post Approval Carotid Stent Trial; MI = myocardial infarction; N = number of patients; PRO-CAS = Predictors of Death and Stroke in CAS; S = stroke; SAPPHIRE = Stenting and Angioplasty with Protection in Patients at High Risk for Endarterectomy.

Reproduced with permission from Roffi et al.⁸²

<30% stenosis (n = 1746, absolute risk increase 2.2%, P = 0.05). CEA had no effect in patients with 30–49% stenosis (n = 1429, absolute risk reduction 3.2%, P = 0.06) and had a small benefit

- in patients with 50–69% stenosis (n = 1549, absolute risk reduction 4.6%, P = 0.04). CEA was highly beneficial in patients with >70% stenosis but with no near occlusion (n = 1095, absolute risk reduction 16.0%, P < 0.001; the number needed to treat to prevent one ipsilateral stroke in 5 years was 6). In contrast, in
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grade flow ('string-flow') in the internal carotid artery, CEA did not show any advantage over medical treatment.

A pooled analysis of the ESCT and NASCET trials (5893 patients with 33 000 patient-years of follow-up) convincingly demonstrated

patients with a 99% stenosis (near occlusion) and sluggish ante-

- that carotid revascularization should be performed rapidly in symptomatic patients with TIA or mild stroke. The number needed to treat to prevent one ipsilateral stroke in 5 years was 5 for those randomized within 2 weeks after the last ischaemic event vs. 125 for patients randomized after 12 weeks.⁹³
- 1615 In symptomatic patients, the benefit of surgery is clearly established for patients with stenosis >70%, but no near occlusion, and to a lesser degree in patients with stenosis 50–69%. It should be underscored that medical therapy in these old trials did not include the use of statins.
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4.1.1.4.2.2 Endovascular therapy versus surgery

A total of six large-scale (i.e. enrolling >300 patients) clinical trials comparing CEA and CAS have been published. The CAVATAS,⁹⁴ EVA-3S,⁹⁵ ICSS,⁹⁶ and SPACE⁹⁷ trials enrolled exclusively symptomatic patients. The SAPPHIRE^{98,99} and CREST⁷⁹ trials included both symptomatic and asymptomatic patients at high and conventional risk for surgery, respectively.

In the CAVATAS study (504 symptomatic patients), performed prior to the introduction of EPDs, most patients allocated to endovascular therapy were treated with angioplasty alone. Only 26% received a stent. There was no statistical difference in terms of any stroke or death at 30 days between CEA and angioplasty (9.9% vs. 10%).⁹⁴ Despite higher restenosis rates in the endovascular arm, no difference in the rates of non-periprocedural ipsilateral stroke was reported at 8-year follow-up.¹⁰⁰

The SAPPHIRE study randomized symptomatic and asymptomatic patients at high risk for surgery.⁹⁸ All endovascular patients were systematically treated with the same stent and a protection device. The trial was designed to prove non-inferiority of CAS and was termi-

- 1640 nated prematurely because of slow enrolment. The primary endpoint of the trial was the cumulative incidence of death, stroke, or myocardial infarction within 30 days after the procedure or ipsilateral stroke occurring between 31 days and 1 year. Among the 334 randomized patients (29% symptomatic), the primary endpoint occurred in
- 1645 12.2% in the CAS group and in 20.1% in the CEA group (P = 0.053). The difference was driven mainly by the rate of myocardial infarction (2.4% in the CAS group vs. 6.1% in the CEA group; P = 0.10). No cranial nerve injury was observed in the CAS group, compared with 5.3% in the CEA group. The durability of CAS was docu-
- 1650 mented by a comparable cumulative percentage of major (1.3% for CAS vs. 3.3% for CEA) and minor (6.1% for CAS vs. 3.0% for CEA) ipsilateral strokes at 3 years and a low rate of repeat revascularization during the same period (3.0% for CAS vs. 7.1% for CEA).⁹⁹

The SPACE study randomized 1200 symptomatic patients.¹⁰¹ Left at the discretion of the treating physician, EPDs were used in 27% of 1655 the cases. The trial was prematurely stopped because of slow enrolment and lack of funding. The incidence of ipsilateral stroke or death at 30 days was the primary endpoint of the study and did not differ between the groups. With an insufficient sample size, SPACE failed to prove the non-inferiority of CAS with the pre-specified absolute 1660 difference of 2.5% (P = 0.09). Follow-up analysis showed no difference in the 2-year rate of adverse events between groups (8.8% for CEA and 9.5% for CAS; P = 0.62).¹⁰²

The EVA-3S trial randomized 527 symptomatic patients with a stenosis $\geq 60\%$ to CAS or CEA.⁹⁵ The primary endpoint was the cumulative incidence of any stroke or death within 30 days after treatment. Although not mandated, CAS without EPD protection was rapidly halted because of excessive risk of stroke compared with those with an EPD (OR 3.9, 95% CI 0.9–16.7).¹⁰³ The trial was stopped prematurely because of significant increased event rates in the CAS 1670 arm (death or stroke 9.6% vs. 3.9% in the CEA arm; P = 0.01). Beyond 30 days, no difference in death or stroke rate was observed, but at 4-year follow-up, the results of CEA were still more favourable than those of CAS, driven by the periprocedural events.¹⁰⁴

The ICSS study randomized 1710 symptomatic patients to CEA 1675 or CAS (EPD use was not mandatory and protected CAS was performed in 72% of patients). The primary endpoint was the 3-year rate of fatal or disabling stroke. While follow-up is ongoing, an interim safety analysis of events between randomization and 120 days reported an incidence of death, stroke, or periprocedural 1680 myocardial infarction in favour of CEA, with an incidence of 8.5% in the CAS group and 5.2% in the CEA group [hazard ratio (HR) 1.69, 95% CI 1.16–2.45; P = 0.004].⁹⁶ The difference was driven mainly by a lower rate of non-disabling strokes in the CEA arm.

The CREST study was a multicentre, randomized controlled trial 1685 (RCT) with the primary endpoint of periprocedural stroke, myocardial infarction, or death, plus ipsilateral stroke up to 4 years. The study was characterized by strict requirements in terms of endovascular credentialing and a lead-in phase that included the treatment of 1541 patients with CAS that preceded the randomized enrolment. 1690 Owing to slow enrolment, this study—initially designed for symptomatic patients—was then extended to include asymptomatic individuals.⁷⁹ The primary endpoint occurred in 7.2% of the CAS group and in 6.8% of the CEA group (HR 1.11, 95% CI 0.81-1.51; P = 0.51). With respect to periprocedural death, stroke, or myocardial 1695 infarction, no difference was observed, with an event rate of 5.2% in the CAS group and 4.5% in the CEA group (P = 0.38). Patients randomized to CAS had more periprocedural strokes (HR 1.79, 95% Cl 1.14–2.82; P = 0.01), but they had fewer myocardial infarctions (1.1% vs. 2.3%; 95% CI 0.26-0.94; P = 0.03) compared with 1700 those receiving CEA. The incidence of major periprocedural strokes was low and not different between the two groups (0.9% vs. 0.6%; P = 0.52). Cranial nerve palsy occurred in 0.3% of patients randomized to CAS and in 4.7% of those treated with CEA (HR 0.07, 95% CI 0.02-0.18; P < 0.0001). At 4 years, no difference in 1705 rates of ipsilateral stroke after the periprocedural period was observed (HR 0.94, 95% CI 0.50-1.76; P = 0.85).

A meta-analysis of 13 randomized trials and including those mentioned above involved 7484 patients, of which 80% had symptomatic disease. Compared with CEA, CAS was associated with 1710 1.40; 95% CI 0.85-2.33).¹⁰⁵

Recommendations for management of asymptomatic carotid artery disease

increased risk of any stroke (RR 1.45; 95% CI 1.06-1.99),

decreased risk of periprocedural myocardial infarction (RR 0.43;

95% CI 0.26-0.71), and non-significant increase in mortality (RR

1720	Recommendations	Class ^a	Level ^b	Ref ^c
	All patients with asymptomatic carotid artery stenosis should be treated with long-term antiplatelet therapy.	I	В	52, 54, 66
1725	All patients with asymptomatic carotid artery stenosis should be treated with long-term statin therapy.	I	С	-
1730 1735	In asymptomatic patients with carotid artery stenosis ≥60%, CEA should be considered as long as the perioperative stroke and death rate for procedures performed by the surgical team is <3% and the patient's life expectancy exceeds 5 years.	lla	A	52, 54, 66
1740	In asymptomatic patients with an indication for carotid revascularization, CAS may be considered as an alternative to CEA in high-volume centres with documented death or stroke rate <3%.	Шь	В	79,99

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^cReferences.

 $\mathsf{CAS}=\mathsf{carotid}$ artery stenting; $\mathsf{CEA}=\mathsf{carotid}$ endarterectomy.

4.1.2 Vertebral artery disease

4.1.2.1 Definition and natural history

The prevalence of vertebral artery (VA) disease due to atherosclerotic disease in the general population is unknown as this condition often remains undiagnosed, because it is either asymptomatic or due to neglected symptoms of vertebrobasilar ischaemia.¹⁰⁶ Approximately 20% of all ischaemic strokes are estimated to involve the vertebrobasilar territory.^{107,108} Vertebrobasilar stroke is primarily the result of an embolic process—most frequently artery-to-artery embolism from the VA origin or cardioembolism. On occasion, dissection, thrombotic, and low-flow haemodynamic mechanisms may be involved.¹⁰⁹ A significant ste-

nosis of the extracranial VA—mostly located at its origin—may account for up to 20% of all vertebrobasilar strokes or TIAs.¹¹⁰

4.1.2.2 Imaging

1765 Data on the accuracy of non-invasive imaging for the detection of extracranial VA are limited and none of the studies has compared different imaging modalities against contrast angiography. A recent

Recommendations for management of symptomatic carotid artery disease

Recommendations	Class ^a	Level ^b	Ref ^c	
All patients with symptomatic carotid stenosis should receive long-term antiplatelet therapy.	I	А	37	1775
All patients with symptomatic carotid stenosis should receive long-term statin therapy.	I.	В	60, 61	
In patients with symptomatic 70-99% stenosis of the internal carotid artery, CEA is recommended for the prevention of recurrent stroke.	I	A	50,51,91, 92	1780
In patients with symptomatic 50-69% stenosis of the internal carotid artery, CEA should be considered for recurrent stroke prevention, depending on patient-specific factors.	lla	A	50, 51, 91, 92	1785
In symptomatic patients with indications for revascularization, the procedure should be performed as soon as possible, optimally within 2 weeks of the onset of symptoms.	I	В	93	1790
In symptomatic patients at high surgical risk requiring revascularization, CAS should be considered as an alternative to CEA.	lla	В	79, 99, 102	1800
In symptomatic patients requiring carotid revascularization, CAS may be considered as an alternative to CEA in high-volume centres with documented death or stroke rate <6%.	Шь	В	79, 99, 102	1800

^cReferences.

CAS = carotid artery stenting; CEA = carotid endarterectomy.

systematic review suggested that MRA offers better sensitivity and specificity than DUS for extracranial VA stenosis.¹¹¹ While CTA is increasingly used for assessment of VA disease, this technique still needs validation.¹¹¹ Both MRA and CTA may be inadequate for ostial VA lesions, especially in the presence of severe angulation or tortuosity of the VA take-off. Despite those limitations, contrast angiography is rarely used merely for diagnostic purposes.

4.1.2.3 Management of vertebral artery disease

The overall benefits of antiplatelet and statin therapy have been presented earlier in these guidelines (Section 3.4.3). Although there are no prospective studies evaluating different therapeutic

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- 1825 strategies in patients with VA disease, aspirin (or if not tolerated clopidogrel) and statins should be administered in all patients, irrespective of symptoms. Asymptomatic VA disease does not require intervention. In general, the need to intervene is tempered by the fact that the posterior circulation is supplied by the confluence of
- 1830 the two VAs, and a large proportion of patients remain asymptomatic despite an occlusion of one VA. However, in patients with recurrent ischaemic events under antiplatelet therapy or refractory vertebrobasilar hypoperfusion, revascularization may be considered.
- 1835 Although surgery of extracranial VA stenosis has been performed with low rates of stroke and mortality by surgeons with extensive experience,¹¹² in most centres the surgical approach has been replaced by endovascular techniques. However, data for VA revascularization are limited to retrospective and mainly
- 1840 single-centre studies.

More information is provided in the online Appendix 2.

Recommendations for revascularization in patients with VA stenosis

	Recommendations	Class ^a	Level ^b
1850	In patients with symptomatic extracranial VA stenosis, endovascular treatment may be considered for lesions ≥50% in the case of recurrent ischaemic events despite optimal medical management.	Шь	С
1855	Revascularization of an asymptomatic VA stenosis is not indicated, irrespective of the degree of severity.	III	С
	 lass of recommendation.		

^bLevel of evidence. VA = vertebral artery.

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4.2 Upper extremity artery disease

4.2.1 Definition and clinical presentation

The subclavian artery and brachiocephalic trunk are the most common locations for atherosclerotic lesions in the upper extremities. However, UEAD can be caused by a number of conditions, involving different levels of the upper extremity arterial system (see online Appendix 3). The most common manifestation for subclavian arterial occlusive disease is unequal arm pressures. A differ-

- 1870 ence of \geq 15 mmHg is highly suspicious for subclavian stenosis. It is not uncommon to detect this occlusive disease in asymptomatic patients. Nevertheless, when the subclavian or brachiocephalic trunk becomes symptomatic, the clinical scenario can be diverse. Subclavian steal syndrome due to flow reversal in the VA, which
- 1875 is worsened by exercising the arm, can evoke symptoms of vertebrobasilar insufficiency (dizziness, vertigo, blurred vision, alternating hemiparesis, dysphasia, dysarthria, confusion, and loss of consciousness, drop attacks, ataxia or other postural disturbances including sensory and visual changes). Patients with coronary
- 1880 bypass with an internal mammary artery can develop symptoms of myocardial ischaemia as the manifestation of subclavian steal

syndrome. Brachiocephalic occlusive disease can also lead to stroke related to the carotid and vertebral territories. Ischaemic arm symptoms are characterized by crampy pain on exercise also referred to as arm claudication. In more severe cases— 1885 especially in more distal disease—rest pain and digital ischaemia with gangrene can develop.

4.2.2 Natural history

Little is known about the natural history of subclavian stenosis, but 1890 the prognosis appears relatively benign. Only subclavian steal with myocardial ischaemia in patients revascularized using the internal mammary artery as well as symptomatic brachiocephalic atherosclerosis with stroke episodes can be considered as life-threatening clinical conditions. However, any symptomatic subclavian occlusive disease should be investigated and treated. Vertebrobasilar insufficiency related to subclavian artery stenosis can be recurrent even after revascularization procedures. It can be explained by numerous other conditions such as cardiac arrhythmias, or intracerebral small vessel disease that can mimic symptoms of vertebrobasilar insufficiency. The combination of proximal and distal arm occlusive disease can present a clinical challenge, with poor prognosis for the extremity.

4.2.3 Clinical examination

Clinical diagnosis of upper limb ischaemia is based on history and physical examination including bilateral blood pressure measurement and assessment of the axillary, brachial, radial, and ulnar artery pulses. Auscultation is an important part of upper extremity examination and should begin in the supraclavicular fossa. Signs and 1910 symptoms, such as pulse deficit, arm pain, pallor, paraesthesia, coldness, and unequal arm pressures, warrant further investigation for occlusive artery disease of the upper limb. The Allen test should be performed in patients in whom the radial artery is instrumented or harvested for coronary revascularization. Adequate col-1915 lateral flow via the ulnar artery is to be confirmed by this test.

4.2.4 Diagnostic methods

4.2.4.1 Duplex ultrasonography

The proximal location of subclavian arterial occlusive disease 1920 makes DUS challenging. However, duplex scanning is of particular value in differentiating occlusion from stenosis, in determining the direction of the vertebral blood flow, and in screening for concurrent carotid artery stenosis. Subclavian steal can be present in the absence of retrograde vertebral flow at rest. Dynamic examination with cuff compression of the upper arm and consecutive hyperaemia after decompression can change the vertebral flow direction.

4.2.4.2 Computed tomography angiography

Upper limb atherosclerosis can be imaged in excellent detail using 1930 CTA. To avoid misinterpretations, it is important to detect congenital abnormalities, in order to define precisely the four vessels perfusing the head. CTA should be analysed interactively, based on a combination of axial images and post-processed views.

4.2.4.3 Magnetic resonance angiography

The use of MRI and contrast-enhanced MRA should also be considered because it enables acquisition of both functional and

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- morphological information. This information can be used to distinguish antegrade from retrograde perfusion. MRA can be combined with special sequences to detect vessel wall oedema and contrast enhancement after administration of intravenous contrast. MRA can detect dilatation and stenosis of the supra-aortic vessels that may be associated with both arteritis and atherosclerosis.
- 1945 Assessment of antegrade and retrograde flow is particularly helpful when steal syndrome is suspected. MRA is particularly useful for follow-up studies.

4.2.4.4 Digital subtraction angiography

1950 DSA is the gold standard in imaging. However, it is increasingly being replaced by other imaging modalities, such as CTA and MRA.

4.2.5 Treatment

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Control of the risk factors for atherosclerosis should be offered to all patients with UEAD, including asymptomatic subjects, because they are at increased risk of death.¹¹³

Revascularization is sometimes indicated in asymptomatic patients, such as CAD patients with planned use of the internal mammary artery for the coronary bypass grafting, or patients with bilateral upper limb lesions to enable blood pressure measurement.

In symptomatic patients endovascular and surgical treatment options are available.

- Neither acute results nor long-term patency rates have been compared in randomized studies for the two techniques. The risk of severe complications is low with both approaches, and in particular the risk of vertebrobasilar stroke is rarely reported. Atherosclerotic lesions of the upper extremities, mostly subclavian lesions, are nowadays treated primarily by endovascular tech-
- 1970 niques. The primary technical success rate is very high and similar to that for surgical treatment. The less invasive nature of endovascular treatment outweighs supposedly better long-term results of surgical interventions.¹¹⁴

Ostial lesions should preferably be treated with balloon-expandable stents because they can be placed more precisely than self-expanding stents. Furthermore, the ostial lesions are more likely to be highly calcified, and in this situation the higher radial force of balloon-expandable stents might be beneficial.

Sixt et al.¹¹⁴ reported a primary success rate of 100% for treatment of stenoses and 87% for occlusions. They also compared stenting procedures with balloon angioplasty and found a trend for an improved 1-year primary patency rate after stent-supported angioplasty (89% vs. 79%). For occlusions, the primary patency rate was 83%.

De Vries et al.¹¹⁵ reported an initial technical success rate of 100% for stenosis and 65% for occlusions. However devices and the experience of the interventionists have since improved and are associated with better results, including for treatment of occlusions. The long-term clinical results in that study were favourable, with a 5-year primary patency rate of 89%.

For subclavian artery occlusions, surgical reimplantation demonstrated long durability with low operative mortality and morbidity rates. Carotid-subclavian bypass with a prosthetic graft is a good surgical alternative. $^{\rm 116}$

Other extra-anatomical bypass modalities, such as axilloaxillary and subclavian–subclavian, are considered the third surgical choice for this pathology. The transthoracic approach is generally 2000 reserved for patients with multivessel aortic and supraortic trunk disease, which may preclude an extra-anatomical repair. The latter surgical option is related to higher mortality and morbidity when compared with transpositions or extra-anatomical reconstructions.¹¹⁷ 2005

Some clinical or anatomical circumstances, such as old age, high surgical risk, previous sternotomy, or calcified ascending aorta, can preclude the transthoracic surgical approach. In these cases, an extra-anatomical or endovascular approach can be applied.¹¹⁸ Nevertheless, no randomized trials have been performed to 2010 compare different therapeutic options. Other therapies, including prostanoid infusion and thoracocervical sympathectomy, may be considered when revascularization is not possible.¹¹⁹

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extremity artery disease Level^b Recommendations Class^a Revascularization is indicated in symptomatic patients. When revascularization is indicated, an endovascular-first strategy is recommended I in patients with atherosclerotic lesions of the upper extremities. Surgery should be considered after failed lla endovascular treatment in low-surgical-risk patients. Revascularization may be considered in

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Recommendations for the management of upper

^aClass of recommendation. ^bLevel of evidence.

occlusions.

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4.3 Mesenteric artery disease

asymptomatic patients with former or future

mammary-coronary bypass or to monitor

blood pressure in bilateral upper limb

4.3.1 Definition

Patients with mesenteric artery disease may be asymptomatic.¹²⁰ Symptomatic mesenteric artery disease is an uncommon, potentially underdiagnosed condition caused by fixed stenoses or occlusion of at least two visceral arteries. Stenosis of one and even two visceral vessels is usually well tolerated because of the abundant collateral circulation between the coeliac trunk, the superior mesenteric artery, and the inferior mesenteric artery—the latter

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sclerosis is the leading cause of mesenteric artery disease (95%). Typically, patients affected by mesenteric artery disease have diffuse atherosclerotic disease including CAD.^{120,121} Nonatherosclerotic causes of mesenteric artery disease such as fibromuscular disease, Dunbar syndrome (compression of the coeliac trunk by the arcuate ligament), and vasculitis will not be discussed.

being connected to branches of the internal iliac arteries. Athero-

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4.3.2 Clinical presentation

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Patients with mesenteric artery disease usually present with abdominal angina, a clinical syndrome characterized by painful abdominal cramps and colic occurring typically in the post-prandial phase.¹²¹ Patients may suffer from ischaemic gastropathy, a condition characterized by the fear of food, nausea, vomiting, diarrhoea, malabsorption, and unintended progressive weight loss.^{122,123} Acute mesenteric ischaemia may also be caused by mesenteric artery thrombosis, with a grim prognosis.

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4.3.3 Prevalence and natural history

The incidence of mesenteric artery disease in the general population is \sim 1 per 100 000 per year.¹²⁴ In patients with known ather-2075 osclerotic disease, the prevalence of mesenteric artery disease may range from 8% to 70%, and a >50% stenosis of more than one splanchnic artery may be detected in up to 15% of cases.¹²⁵⁻¹²⁸ In patients with abdominal aortic aneurysm, aortoiliac occlusive disease, and infrainguinal LEAD, a significant stenosis of at least 2080

one of the three visceral arteries may be found in 40, 29, and 25% of cases, respectively.¹²⁰ Predisposing conditions for the development of mesenteric artery disease include arterial hypertension, diabetes mellitus, smoking, and hypercholesterolaemia. Untreated symptomatic mesenteric artery disease may lead to 2085 starvation, bowel infarction, and death.

4.3.4 Diagnostic strategy

DUS has become the imaging method of choice for mesenteric artery disease.¹²⁹⁻¹³³ The diagnostic performance may be 2090 improved by a post-prandial test, revealing increased velocity and turbulences, which may seem trivial in a fasting patient. CTA and gadolinium-enhanced MRA are useful initial tests for supporting the clinical diagnosis of symptomatic mesenteric artery disease if the results of DUS are inconclusive.^{134–137} Recently, 24 h gastro-2095 intestinal tonometry has been validated as a diagnostic test to detect splanchnic ischaemia and to guide treatment.¹³⁸ Basically, gastrointestinal tonometry measures gut intraluminal CO₂. Intra-

based on the concept that in situations where gastrointestinal per-2100 fusion is reduced oxygen delivery falls below a critical level, resulting in anaerobic cellular metabolism that leads to local lactic acidosis and generation of CO_2 .

Ischaemic colitis is frequently diagnosed by histology following biopsy during bowel endoscopy. DSA is still considered the diag-

luminal gut CO₂ is elevated when local perfusion is compromised

2105 nostic gold standard, but its use is now limited to periinterventional imaging.^{139,140}

Recommendations for diagnosis of symptomatic chronic mesenteric ischaemia

Recommendations	Class ^a	Level ^b	Ref ^c	
DUS is indicated as the first- line diagnostic test in patients suspected of mesenteric artery disease.	I	A	29- 33, 38	2115
When DUS is inconclusive, CTA or gadolinium-enhanced MRA are indicated.	I	В	135-137, 139, 141	2120
Catheter-based angiography is indicated exclusively during the endovascular therapy procedure.	I	С	-	2425
				2125

^aClass of recommendation.

^bLevel of evidence.

^cReferences.

CTA = computed tomography angiography; DUS = duplex ultrasonography; MRA = magnetic resonance angiography.

4.3.5 Prognostic stratification

Five-year mortality in asymptomatic patients with mesenteric artery disease is estimated at 40%, and up to 86% if all three main visceral arteries are affected.¹²⁰ Diffuse mesenteric artery 2135 disease in asymptomatic subjects should be considered as a marker of increased cardiovascular mortality, justifying aggressive management of cardiovascular risk factors.

4.3.6 Treatment

Recent reports have suggested that endovascular therapy, with or without stenting, may have a lower perioperative mortality rate than open surgery for revascularization of mesenteric artery disease. Retrospective data from a US nationwide inpatient sample analysis (1988-2006) including >22 000 patients suggested a lower mortality rate ₂₁₄₅ after endovascular therapy compared with surgical bypass (3.7% vs. 13%, P < 0.01).¹⁴² In addition, bowel resection was less frequent in the endovascular group than in the surgical group (3% vs. 7%, P <0.01). Bowel resection was, in general, associated with a high in-hospital mortality rate [percutaneous transluminal angioplasty 2150 (PTA)/stenting 25% and surgery 54%, respectively]. The lower in-hospital mortality rates reported after angioplasty with or without stenting indicate that this strategy should be proposed when possible. Longitudinal data are needed to determine the durability of this benefit. So far no randomized controlled data are available. 2155

Symptom relief following revascularization is reported in up to 100% of cases, although restenosis after endovascular therapy may be frequent (29-40%). Although no controlled data support the strategy, dual antiplatelet therapy for 4 weeks postprocedure, followed by long-term aspirin treatment, has become 2160 the standard of care. DUS follow-up every 6-12 months is recommended. The use of drug-eluting stents, flared stent devices, or drug-eluting balloons in conjunction with bare-metal stents has not yet been evaluated in larger studies.

Recommendations for the management of mesenteric artery disease

2170	Recommendations	Class ^a	Level ^b	Ref ^c
2175	Mesenteric revascularization should be considered in patients with symptomatic mesenteric artery disease.	lla	В	120, 143-150
	In the case of revascularization, endovascular treatment should be considered as the first-line strategy.	lla	с	-

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^aClass of recommendation. ^bLevel of evidence. ^cReferences.

4.4 Renal artery disease 2185

Renal artery disease is increasingly related to atherosclerosis with advancing age and prevalent hypertension, diabetes mellitus, renal disease, aortoiliac occlusive disease, and CAD.¹⁵¹ In the elderly population, atherosclerosis accounts for \sim 90% of cases and usually involves the ostium and proximal third of the main renal artery and the perirenal aorta. Less frequent causes are fibromuscular dysplasia and arteritis. Screening angiography in potential kidney donors indicates that RAS can be asymptomatic and may be present in up to 3-6% of normotensive individuals.¹⁵²

2195 4.4.1 Clinical presentation

Major clinical signs of RAS include refractory hypertension, unexplained renal failure, and flash pulmonary oedema (Table 4). RAS may cause or deteriorate arterial hypertension and/or renal failure. Hypoperfusion of the kidney activates the renin-angiotensin-aldosterone system (RAAS), causing classic renovascular hypertension, primarily in young patients with fibromuscular dysplasia.^{151,153} However, in patients with atherosclerosis, RAS may induce an acute or subacute acceleration of a pre-existing essential hypertension including flash pulmonary oedema usually in bilateral kidney disease.¹⁵¹ The association between RAS severity and ischaemic nephropathy^{154,155} has recently been challenged.¹⁵⁶ The loss of filtration capacity of the kidney in RAS may be due

not only to hypoperfusion, but also to recurrent microembolism. Renal failure may occur with severe bilateral RAS or unilateral 2210 stenosis in a single functional kidney.

Kidney disease and renovascular disease promote CVD and hypertension. Increased risk of CVD in atherosclerotic RAS patients may result from activation of the RAAS and sympathetic nervous systems, decreased GFR, or concomitant atherosclerosis

2215 in other vascular beds.^{157–159} The prevalence of left ventricular hypertrophy with RAS is 79% vs. 46% in patients with essential hypertension, with a substantial impact on morbidity and mortality.^{160–162}

2220 4.4.2 Natural history

Data on progression of atherosclerotic RAS are inconsistent. More recent studies show significant disease progression to high-grade stenosis or occlusion in only 1.3-11.1% of patients, whereas

Table 4 Clinical situations where the diagnosis of RAS 2225 should be considered

	_	
Clinical presentation		
• Onset of hypertension before the age of 30 years and after 55 years		
 Hypertension with hypokalemia, in particular when receiving thiazide diuretics 		2230
• Hypertension and abdominal bruit		
 Accelerated hypertension (sudden and persistent worsening of previously controlled hypertension) 		2235
• Resistant hypertension (failure of blood-pressure control despite full doses of an appropriate three-drug regimen including a diuretic)		
 Malignant hypertension (hypertension with coexistent end-organ damage, i.e. acute renal failure, flash pulmonary oedema, hypertensive left ventricular failure, aortic dissection, new visual or neurological disturbance, and/or advanced retinopathy) 		2240
• New azotemia or worsening renal function after the administration of an angiotensin-converting enzyme inhibitor or an angiotensin II receptor blocker		
Unexplained hypotrophic kidney		2245
Unexplained renal failure		
RAS = renal artery stenosis.	-	

older studies documented occlusion rates up to 18% over 5 years.¹⁶³⁻¹⁶⁶ After 2 years, 3, 18, and 55% of the kidneys had lost their function in the case of unilateral stenosis, bilateral stenosis, and contralateral occlusion, respectively.¹⁶⁷

4.4.3 Diagnostic strategy

R

Baseline diagnostic evaluation includes physical examination, exclusion of other potential causes of secondary hypertension, and ambulatory blood pressure measurement. In clinical situations in which RAS is suspected, such as those listed in Table 4, renal 2260 artery imaging should be considered.

DUS is the first-line screening modality for atherosclerotic RAS. It can be applied serially to assess the degree of stenosis and physiological patterns, such as flow velocities and vascular resistance. Increased peak systolic velocity in the main renal artery associated 2265 with post-stenotic turbulence is most frequently used to determine relevant RAS, and corresponds to >60% angiographic RAS with a sensitivity and specificity of 71-98% and 62-98%, respectively.^{168–170} Several duplex criteria should be used to identify significant (>60%) stenosis. These include imaging of intrarenal 2270 interlobar or segmental arteries, including calculation of the sidedifference of the intrarenal resistance index, missing early systolic peak, retarded acceleration, and increased acceleration time, which are less specific and should be used to support the diagnosis based on peak systolic velocity.¹⁷¹⁻¹⁷³ 2275

Common pitfalls of DUS include failure to visualize the entire renal artery and missing the highest peak systolic velocity during spectral Doppler tracing. Accessory renal arteries are generally not adequately examined or identified. The accuracy of DUS is operator dependent.

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Both 3D MRA and multidetector CTA have demonstrated equally high sensitivities (>90%) for detection of haemodynamically significant stenoses, with excellent interobserver and intermodality agreement. 174

Gadolinium-enhanced MRA provides excellent characterization of the renal arteries, surrounding vessels, renal mass, and occasion-

ally renal function. It is less useful in patients with renal artery

2285 Currently CTA provides higher spatial resolution than MRA and may be more readily available; however, the requirement to use iodinated contrast makes it an unattractive modality in patients with impaired renal function.

2290

stents because of artefacts. In addition, MRA tends to overestimate the degree of luminal narrowing. A recent concern in the use of gadolinium-enhanced MRI is nephrogenic systemic fibrosis, with an incidence ranging from 1% to 6% for dialysis patients, and a

GFR < 30 mL/min was designated as a contraindication.¹⁷⁵ In recent years measuring the translessional pressure gradient with a dedicated pressure wire was proposed to identify a significant RAS. A distal-to-the-lession to aortic pressure ratio at rest of <0.9 was linked to an upregulation of renin production.¹⁵¹ This

2300 <0.9 was linked to an upregulation of renin production.¹⁵¹ This ratio correlates to a papaverine-induced hyperaemic systolic

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Recommendations for diagnostic strategies for RAS

2305				
	Recommendations	Class ^a	Level ^b	Ref ^c
2310	DUS is recommended as the first-line imaging test to establish the diagnosis of R		В	171, 172
2310	CTA (in patients with creatir clearance >60 mL/min) is recommended to establish th diagnosis of RAS.		В	151, 174
2315	MRA (in patients with creatii clearance >30 mL/min) is recommended to establish th diagnosis of RAS.	1.1	В	174
2320	When the clinical index of suspicion is high and the results of non-invasive tests are inconclusive, DSA is recommended as a diagnostic test (prepared for intervention) to establish th diagnosis of RAS.		с	-
2325 2330	Captopril renal scintigraphy selective renal vein renin measurements, plasma renin activity, and the captopril test are not recommended as useful screening tests to establish the diagnosis of R	п Ш	В	151, 178

^aClass of recommendation.

^bLevel of evidence.

^cReferences.

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CTA = computed tomography angiography; DSA = digital subtraction angiography; DUS = duplex ultrasonography; MRA = magnetic resonance angiography; RAS = renal artery stenosis. pressure gradient of >21 mmHg.¹⁷⁶ A dopamine-induced mean pressure gradient of >20 mmHg predicted a beneficial blood pressure response to renal stenting.¹⁷⁷

DSA is generally limited to pre-angioplasty visualization and quantification of the stenosis. It may also be considered in patients with high clinical suspicion of RAS already scheduled for another angiographic examination (e.g. coronary angiography) or in the case of inconclusive non-invasive imaging. 2345

4.4.4 Prognostic stratification

Among patients with ESRD, the life expectancy of those with RAS is the poorest.¹⁷⁹ However, life expectancy is also significantly reduced in patients with RAS without ESRD.¹⁷⁹ Two-year mor- 2350 tality in patients with baseline serum creatinine concentrations before revascularization of <1.2 mg/dL, 1.2–2.5 mg/dL, and >2.5 mg/dL were 5, 11, and 70%, respectively.¹⁸⁰ More than 80% of patients die due to cardiovascular events.

4.4.5 Treatment

Beyond secondary prevention of atherosclerosis, the treatment of renal artery disease should be aimed at control of blood pressure and preservation of renal function.

4.4.5.1 Medical treatment

ACE inhibitors and calcium channel blockers are effective in the treatment of hypertension in the presence of RAS and may lead to slowing of the progression of renal disease.¹⁸¹ Most patients with haemodynamically significant RAS tolerate RAAS blockade 2365 without difficulty. However, ACE inhibitors can reduce glomerular capillary hydrostatic pressure enough to cause a transient decrease in GFR and raise serum creatinine, warranting caution and close follow-up. A significant (\geq 30%) fall in GFR (or a >0.5 mg/dL rise in serum creatinine) may be an indication to consider renal 2370 revascularization. ACE inhibitors are contraindicated in the case of bilateral RAS and when this lesion affects a single functional kidney.

There is evidence that thiazides, hydralazine, angiotensin II receptor blockers, and β -blockers are also effective in achieving 2375 target blood pressures in individuals with RAS.^{182–184}

All patients with atherosclerotic RAS should be treated according to the European Guidelines on Cardiovascular Disease Prevention. 24

4.4.5.2 Revascularization

The decision regarding the potential revascularization strategy should be based on the patient's individual characteristics, such as life expectancy, co-morbidities, quality of blood pressure control, and renal function.

Evidence supporting the benefit of aggressive diagnosis and timing of renal revascularization remains unclear. Among patients receiving medical therapy alone, there is the risk for deterioration of kidney function with worsening morbidity and mortality. Renal artery revascularization can provide immediate improvement in 2390 kidney function and blood pressure; however, as with all invasive interventions, it may result in mortality or substantial morbidity in a small percentage of patients. This is particularly the case for renovascular lesions that pose no immediate hazard or risk of

2395 progression. There is general consensus that renal revascularization should be performed in patients with anatomically and functionally significant RAS who present with particular clinical scenarios such as sudden onset or 'flash' pulmonary oedema or congestive heart failure with preserved left ventricular function 2400 and acute oligo-/anuric renal failure with kidney ischaemia.

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4.4.5.2.1 Impact of revascularization on blood pressure control

Twenty-one uncontrolled series of stenting/angioplasty published before 2007 in 3368 patients gave no unifying pattern regarding mortality rates. Cure, improvement, or worsening of arterial hypertension was documented to range from 4% to 18%, from 35% to 79%, and from 0% to 13%, respectively. Two studies reported a statistically significant reduction in the New York Heart Association functional class after stent placement in patients with either bilateral disease or stenosis to a solitary functioning

when childe blatteral disease of scenesis to a solidary functioning kidney (global ischaemia). For these patients with congestive heart failure and repeated admissions for pulmonary oedema not associated with CAD, improved volume management, restored sensitivity to diuretics, and lowered rehospitalization rates suggest that some individualized patient categories benefit substantially from renal revascularization.^{185–188}

Three RCTs compared endovascular therapy with medical treatment with ≥ 6 months of follow-up.^{166,183,189} Notably, these trials were small and had no adequate power for clinical outcomes. Stents were rarely used and medical therapies varied both between and within studies. In a randomized study including 49 patients, the investigators concluded that endovascular therapy in unilateral atherosclerotic RAS enables reduction of the number of antihypertensive drugs,¹⁸⁹ but that previous uncontrolled studies overestimated the potential for lowering blood pressure. In the

- ²⁴²⁵ Dutch Renal Artery Stenosis Intervention Cooperative (DRASTIC) study involving 106 patients, ¹⁶⁶ there were no significant differences between the angioplasty and drug therapy groups in terms of systolic and diastolic blood pressures or renal function, whereas daily drug doses were reduced in the angioplasty group. However, a significant
- ²⁴⁵⁰ improvement in systolic and diastolic blood pressures was reported after angioplasty in a meta-analysis of these three studies.¹⁹⁰ Two recent randomized trials comparing stent angioplasty combined with medical therapy with medical therapy alone [Angioplasty and Stenting for Renal Artery Lesions trial (ASTRAL) and the Stent Pla-
- ²⁴³⁵ cement in Patients With Atherosclerotic Renal Artery Stenosis and Impaired Renal Function (STAR)] failed to demonstrate any significant difference in blood pressure.^{191,192} However, in the ASTRAL trial, the daily drug dosage was reduced.¹⁹¹

2440 4.4.5.2.2 Impact of revascularization on renal function The ASTRAL trial is so far the largest RCT to determine whether percutaneous revascularization combined with medical therapy compared with medical therapy alone improves renal function.¹⁹¹ Eight-hundred and six patients with atherosclerotic RAS in whom

- the need for revascularization was uncertain were enrolled. Fiftynine per cent of patients were reported to have RAS >70%, and 60% had a serum creatinine of \geq 150 µmol/L. At a mean follow-up of 33.6 months (range 1–4 years), differences in renal function and kidney and cardiovascular events were all similarly unimpressive,
- 2450 even in the highest risk groups, which included patients with global ischaemia or impaired or rapidly decreasing kidney function. The

primary study endpoint-the decline in renal function over timecalculated as the mean slope of the reciprocal of the serum creatinine concentration over time, was slightly slower in the revascularization group, but the difference was not statistically significant. 2455 The STAR multicentre trial enrolled 140 patients to detect a \geq 20% decrease in creatinine clearance.¹⁹² At 2 years, the primary endpoint was reached in 16% of patients in the stented group and in 22% of patients in the medical treatment group. The difference was not statistically significant and was inconclusive, given the wide 2460 confidence intervals around the estimate of effect. It was noteworthy that >50% of the patients randomized to stenting had a <70%diameter stenosis and 28% of patients did not receive a stent (19%) because of no RAS > 50%. This largely underpowered trial showed that deterioration of renal function may progress despite 2465 successful revascularization, underscoring the complex cause of ischaemic nephropathy, with an important parenchymal component affected by risk factors for atherosclerosis. It also showed that if technical skills are insufficient, a considerable number of stent-related complications can occur (two procedure-related 2470 deaths, one death secondary to an infected haematoma, and one case of deterioration of renal function resulting in dialysis).

4.4.5.2.3 Impact of revascularization on survival

In the ASTRAL and STAR trials no difference was seen in the secondary endpoints—cardiovascular morbidity and death. A recent analysis of two consecutive registries comparing conservative treatment with revascularization showed a 45% reduction in mortality for the revascularization cohort.¹⁹³ To date, no major differences in survival are evident between patients undergoing either surgical or endovascular procedures, although only a few studies addressed this issue directly.

Several factors may argue against renal revascularization or predict poorer outcomes, including the presence of proteinuria >1 g/24 h, renal atrophy, severe renal parenchymal disease, and severe diffuse intrarenal arteriolar disease. Moreover, adverse consequences of renal atheroembolization at the time of surgical revascularization have been documented.¹⁹⁴ Similarly, atheroembolization may be provoked by percutaneous revascularization.^{192,195,196}

The potential physiological benefits of renal stent placement 2490 include reperfusion of the ischaemic kidney(s), resulting in a reduction in the stimulus to renin production, which decreases angiotensin and aldosterone production, thereby decreasing peripheral arterial vasoconstriction and preventing hypervolaemia. Improvement in renal perfusion enhances glomerular filtration 2495 and therefore promotes natriuresis. Moreover, reduction of humoral activation may result in reduction of left ventricular mass and improvement of diastolic dysfunction.^{197–199}

The ASTRAL study did not provide information on how to treat patients with a clinical need for revascularization. This question is 2500 being addressed by two ongoing RCTs. The Cardiovascular Outcomes in Renal Atherosclerotic Lesions (CORAL) trial tests the hypothesis that stenting atherosclerotic RAS >60% (systolic pressure gradient >20 mmHg) in patients with systolic hypertension reduces the incidence of cardiovascular and renal events. 2505 The Randomized, Multicentre, Prospective Study Comparing Best Medical Treatment Versus Best Medical Treatment Plus Renal Artery Stenting in Patients With Haemodynamically Relevant

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Atherosclerotic Renal Artery Stenosis (RADAR) investigates the impact of renal stenting on the change in renal function in 300 patients.²⁰⁰

4.4.5.2.4 Technical outcomes of endovascular revascularization Balloon angioplasty with bailout stent placement if necessary is recommended for fibromuscular dysplasia lesions.^{201–204} In

Recommendations: treatment strategies for RAS

520	Recommendations	Class ^a	Level ^b	Ref ^c
	Medical therapy			
525	ACE inhibitors, angiotensin II receptor blockers, and calcium channel blockers are effective medications for treatment of hypertension associated with unilateral RAS.	I.	В	166, 182, 183, 189, 192, 219
530	ACE inhibitors and angiotensin II receptor blockers are contraindicated in bilateral severe RAS and in the case of RAS in a single functional kidney.	ш	В	151, 166, 182, 183, 189, 192
	Endovascular therapy			
535	Angioplasty, preferably with stenting, may be considered in the case of >60% symptomatic RAS secondary to atherosclerosis.	llb	A	151, 201-204
540	In the case of indication for angioplasty, stenting is recommended in ostial atherosclerotic RAS.	I	В	205, 220
545	Endovascular treatment of RAS may be considered in patients with impaired renal function.	ΠР	В	193, 206, 221-223
545	Treatment of RAS, by balloon angioplasty with or without stenting, may be considered for patients with RAS and unexplained recurrent congestive heart failure or sudden pulmonary oedema and preserved systolic left ventricular function.	Ш	c	-
	Surgical therapy			
555	Surgical revascularization may be considered for patients undergoing surgical repair of the aorta, patients with complex anatomy of the renal arteries, or after a failed endovascular procedure.	ШЬ	с	-

^bLevel of evidence.

^cReferences.

ACE = angiotensin-converting enzyme; RAS = renal artery stenosis.

atherosclerotic RAS, stent placement has consistently proven superior to balloon angioplasty in the treatment of renal artery atherosclerotic lesions.²⁰⁵ Restenosis rates range from 3.5% to ~20%^{206,207}; drug-eluting stents have not yet been shown to achieve a significantly better outcome.^{208,209} The appropriate treat-2570 ment modality of in-stent RAS has not yet been defined. Balloon angioplasty, bare-metal stent, covered stent, and drug-eluting stent placement are still under investigation.^{210–213} The role of distal protection devices is still a matter of debate. Following several promising single-centre reports, results from a small, randomized trial¹⁹⁶ showed no significantly improved renal function outcome for distal filter protection during stent revascularization except when an adjunctive glycoprotein Ilb/Illa receptor antagonist was used.

4.4.5.2.5 Role of surgical revascularization

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Renal artery surgery offers major benefits for patients undergoing surgical repair of the aorta, and for patients with complex disease of the renal arteries, e.g. aneurysms or failed endovascular procedures. Thirty-day mortality rates range from 3.7% to 9.4%. After a follow-up of up to 5 years, the need for reoperation has been reported in 5–15% and survival in 65–81% of patients.^{214–218} ²⁵⁸⁵ Major arguments against surgical revascularization include higher mortality linked to surgery in patients with co-morbidities and similar benefits of endovascular repair.

The list of pivotal published and ongoing trials in patients with RAS is provided in Appendix 4. 2590

4.5 Lower extremity artery disease

4.5.1 Clinical presentation

LEAD has several different presentations, categorized according to 2595 the Fontaine or Rutherford classifications (*Table 5*). Importantly, even with a similar extent and level of disease progression, symptoms and their severity may vary from one patient to another.

4.5.1.1 Symptoms

Many patients are asymptomatic. In this situation, LEAD is diagnosed by clinical examination (absent pulses) or by the ABI. Importantly, asymptomatic patients are also at high risk for cardiovascular events.²

The most typical presentation of LEAD is intermittent claudication, characterized by pain in the calves, increasing with walking; the pain 2605 typically disappears quickly at rest (Fontaine stage II; Rutherford grade I). In the case of a more proximal level of arterial obstruction (i.e. the aortoiliac segment), patients may complain of pain extension into the thighs and buttocks. Isolated buttock claudication is rare and due to bilateral hypogastric severe disease. The pain should be distin- 2610 guished from that related to venous disease (usually at rest, increasing in the evening, often disappearing with some muscle activity), hip or knee arthritis (pain on walking but not disappearing at rest), and peripheral neuropathy (characterized more by instability while walking, pain not relieved by resting). Typical intermittent claudication can 2615 also be caused by lumbar spinal stenosis. The Edinburgh Claudication Questionnaire²²⁴ is a standardized method to screen and diagnose intermittent claudication, with a 80-90% sensitivity and >95% specificity (available online at http://www.ncbi.nlm.nih.gov/pmc/articles/ PMC2560464/?page=1). More recently, several studies highlighted 2620 that a substantial proportion of patients with symptomatic LEAD present with atypical symptoms.²²⁵

In more severe cases pain is present at rest, in the supine position (Fontaine stage III; Rutherford grade II). Rest pain is localized more

- 2625 often in the foot and should be distinguished from muscle cramping or arthritis. Patients often complain of permanent coldness in the feet. Ulcers and gangrene (Fontaine stage IV; Rutherford grade III) indicate severe ischaemia and begin mostly at the level of toes and the distal part of the limb. Arterial ulcers are, in most cases, extre-
- 2630 mely painful; they are frequently secondary to local trauma, even minor, and should be distinguished from venous ulcers. When pain is absent, peripheral neuropathy should be considered. Ulcers are often complicated by local infection and inflammation.

Critical limb ischaemia is the most severe clinical manifestation 2635 of LEAD, defined as the presence of ischaemic rest pain, and

2640	Fontaine classification			Ruth	erford cla	ssification	
		Stage	Symptoms	\leftrightarrow	Grade	Category	Symptoms
		I	Asymptomatic	\leftrightarrow	0	0	Asymptomatic
2645					I	I	Mild claudication
		П	Intermittent claudication	\leftrightarrow	I	2	Moderate claudication
2650					I	3	Severe claudication
		ш	lschaemic rest pain	\leftrightarrow	II	4	lschaemic rest pain
2655		IV	Ulceration or		ш	5	Minor tissue loss
		IV	gangrene		ш	6	Major tissue loss

Table 5Clinical staging of LEAD

LEAD = lower extremity artery di 2660

ischaemic lesions or gangrene objectively attributable to arterial 2680 occlusive disease.

4.5.1.2 Clinical examination

Clinical examination can be quite informative both for screening and for diagnosis. Patients should be relaxed and acclimatized to the room temperature. Inspection may show pallor in more severe cases, sometimes at leg elevation. Pulse palpation is very informative for screening purposes and should be done systematically. Pulse abolition is a specific rather than a sensitive clinical sign. Auscultation of bruits over the femoral artery at the groin and more distally is also suggestive, but poorly sensitive. The value of the clinical findings in patients with LEAD can be strongly improved by measuring the ABI. The blue toe syndrome is characterized by a sudden cyanotic discolouration of one or more toes; it is usually due to embolic atherosclerotic debris from the proximal arteries. 2690 2690 2690 2690

4.5.2 Diagnostic tests

4.5.2.1 Ankle-brachial index

The primary non-invasive test for the diagnosis of LEAD is the ABI. $_{2700}$ In healthy persons, the ABI is >1.0. Usually an ABI <0.90 is used to define LEAD. The actual sensitivity and specificity have been estimated, respectively, at 79% and 96%.²²⁶ For diagnosis in primary care, an ABI <0.8 or the mean of three ABIs <0.90 had a positive predictive value of \geq 95%; an ABI >1.10 or the mean of three ABIs >1.00 had a negative predictive value of \geq 99%.²²⁷ The level of ABI also correlates with LEAD severity, with high risk of amputation when the ABI is <0.50. An ABI change >0.15 is generally required to consider worsening of limb perfusion over time, or improving after revascularization.²²⁸

For its measurement (*Figure* 2), a 10–12 cm sphygmomanometer cuff placed just above the ankle and a (handheld) Doppler instrument (5–10 MHz) to measure the pressure of the posterior and anterior tibial arteries of each foot are required. Usually the highest ankle systolic pressure is divided by the highest brachial systolic pressure, resulting in an ABI per leg. Recently some papers reported higher sensitivity to detect LEAD if the ABI numerator is the lowest pressure in the arteries of both ankles.²²⁹

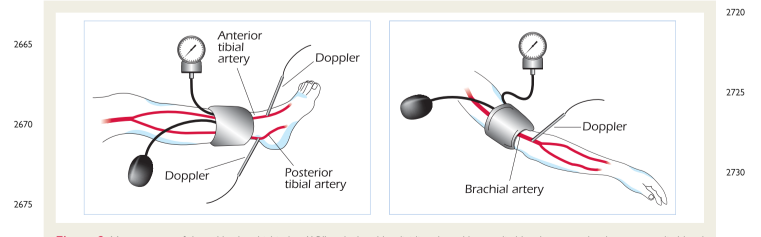


Figure 2 Measurement of the ankle-brachial index (ABI), calculated by dividing the ankle systolic blood pressure by the arm systolic blood pressure.

Measuring ABI after exercise enables the detection of additional subjects with LEAD, who have normal or borderline ABI at rest. The patient is asked to walk (commonly on a treadmill at 3.2 km/

h at a 10-20% slope) until claudication pain occurs and impedes 2740 walking. An ABI drop after exercise seems especially useful when resting ABI is normal but there is clinical suspicion of LEAD.²³⁰

Some patients have an ABI >1.40, related to stiff (calcified) arteries, a condition often observed in the case of diabetes,

ESRD, and in the very elderly. Importantly, a substantial proportion 2745 of patients with an elevated ABI actually do have occlusive artery disease.²³¹ Alternative tests such as measurement of toe systolic pressures and Doppler waveform analysis are useful to unmask LEAD.²³¹ A toe-brachial index < 0.70 is usually considered diagnostic of LEAD. 2750

Recommendations for ABI measurement

2755	Recommendations	Class ^a	Level ^b	Ref ^c
2733	Measurement of the ABI is indicated as a first-line non- invasive test for screening and diagnosis of LEAD.	I	В	226
2760	In the case of incompressible ankle arteries or ABI >1.40, alternative methods such as the toe-brachial index, Doppler waveform analysis or pulse volume recording should be used.	I	В	231

^aClass of recommendation. ^bLevel of evidence.

^cReferences.

ABI = ankle-brachial index; LEAD = lower extremity artery disease.

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4.5.2.2 Treadmill test

The treadmill test is an excellent tool for obtaining objective functional information, mainly on symptom onset distance and maximum walking distance. It is useful in patients with borderline ABI at rest with symptoms suggestive of LEAD. It can 2775 also help to differentiate vascular claudication (with leg pressure drop after exercise) from neurogenic claudication (leg pressure remains stable or increases). The standardized treadmill test is also proposed to assess treatment efficacy (exercise rehabilitation, drug therapies, and/or revascularization) during follow-up. 2780

- Usually the test is performed on a treadmill walking at 3.2 km/h with a 10% slope. However, there are several technical variations,²³² such as introducing a steady increase in elevation of the treadmill every 3 min while keeping the speed constant.
- The test should be supervised to observe all symptoms occur-2785 ring during the test. It should be avoided in the case of severe CAD, decompensated heart failure, or major gait disturbances. It is usually associated with ABI measurement before and after exercise. A pressure drop >20% immediately after exercise confirms the arterial origin of symptoms.²³³ For patients 2790
- unable to perform treadmill exercise, alternative tests such as repeated pedal flexions can be used, with excellent correlation with the treadmill test.

Recommendations for treadmill testing in patients with 2795 LEAD

R	ecommendations	Class ^a	Level ^b	Ref ^c		
cc as to	ne treadmill test should be onsidered for the objective sessment of treatment improve symptoms in audicants.	lla	A	234, 235		2800
aty of sh dia fo	the case of typical or ypical symptoms suggestive LEAD, the treadmill test ould be considered for agnostic confirmation and/or r baseline quantification of nctional severity.	lla	В	234		2805
Level	of recommendation. of evidence. ences.				<u> </u>	2810

LEAD = lower extremity artery disease.

4.5.2.3 Ultrasound methods

DUS provides extensive information on both arterial anatomy and blood flow. Compared with DSA, several concordant meta-analyses estimated DUS sensitivity to detect >50% diameter angiographic stenosis at 85–90%, with a specificity >95%.^{236–238} No significant 2820 differences were found between the above- and below-knee lesions.^{236,238} DUS can also visualize run-off vessels, especially when using the colour mode. DUS depends greatly on the examiner's experience, and adequate qualification and training are mandatory. Combined with the ABI, DUS provides all the information 2825 necessary for management decisions in the majority of patients with LEAD, confirms the diagnosis, and provides information on lesion location and severity. The lesions are located by twodimensional (2D) ultrasonography and colour-Doppler mapping, while the degree of stenosis is estimated mostly by Doppler wave- 2830 form analysis and peak systolic velocities and ratios. The interobserver reproducibility of the DUS to detect >50% stenosis in lower extremity arteries is good, except for pedal arteries.^{239,240}

DUS is also highly useful for the follow-up after angioplasty or to monitor bypass grafts.^{241,242} Excellent tolerance and lack of radiation 2835 exposure make DUS the method of choice for routine follow-up.

Pitfalls of DUS are related mainly to difficulties in assessing the lumen in highly calcified arteries. Insonation in the area of open ulcers or excessive scarring may not be possible. Also in some cases (e.g. obesity, gas interpositions), the iliac arteries are more 2840 difficult to visualize and alternative methods should be considered when the imaging is suboptimal. The major disadvantage of DUS compared with other imaging techniques (DSA, CTA, or MRA) is that it does not provide full arterial imaging as a clear roadmap, as do the other techniques. However, in contrast to 2845 other imaging technique (DSA, CTA, and MRA), DUS provides important information on haemodynamics. Complete DUS scanning of the entire arterial network can be time-consuming. Although aggregate images or schemas can be provided, another imaging technique is usually required, especially when bypass is 2850 considered.²⁴³ However, even in this situation, DUS can be an important aid in determining the most appropriate site of anastomosis by identification of the least calcified portion of the vessel.²⁴⁴ Intravascular ultrasound has been proposed for plaque charac-

terization and after angioplasty, but its routine role in the clinical 2855 setting requires further investigation.

4.5.2.4 Computed tomography angiography

- CTA using MDCT technology allows imaging with high resolution. Compared with DSA, the sensitivity and specificity for occlusions 2860 reported using the single-detector techniques already reached a high degree of accuracy. In a recent meta-analysis, the reported sensitivity and specificity of CTA to detect aortoiliac stenoses >50% were 96% and 98%, respectively.²⁴⁵ The same study showed similar sensitivity (97%) and specificity (94%) for the 2865
- femoropopliteal region, comparable with those reported for the below-knee arteries (sensitivity 95%, specificity 91%).²⁴⁵

The great advantage of CTA remains the visualization of calcifications, clips, stents, and bypasses. However, some artefacts may be present due to the 'blooming effect'.

4.5.2.5 Magnetic resonance angiography

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- MRA can non-invasively visualize the lower limb arteries even in the most distal parts. The resolution of MRA using gadolinium-enhanced contrast techniques reaches that of DSA. In 2875 comparison with DSA, MRA has an excellent sensitivity (93-100%) and specificity (93-100%).^{237,246-250} Owing to different techniques (2D and 3D, with or without gadolinium), the results are not as uniform as for CTA, and studies comparing MRA with
- CTA are not available. In direct comparison, MRA has the greatest 2880 ability to replace diagnostic DSA in symptomatic patients to assist decision making, especially in the case of major allergies. There are also limitations for the use of MRA in the presence of pacemakers or metal implants (including stents), or in patients with claustro-2885
- phobia. Gadolinium contrast agents cannot be used in the case of severe renal failure (GFR < 30 mL/min per 1.73 m²). Of note, MRA cannot visualize arterial calcifications, which may be a limitation for the selection of the anastomotic site for a surgical bypass.

4.5.2.6 Digital subtraction angiography 2890

For the aorta and peripheral arteries, retrograde transfemoral catheterization is usually used. Cross-over techniques allow the direct antegrade flow imaging from one side to the other. If the femoral access is not possible, transradial or transbrachial approaches and

direct antegrade catheterization are needed. Considered as the 2895 gold standard for decades, DSA is now reserved for patients undergoing interventions, especially concomitant to endovascular procedures. Indeed, the non-invasive techniques provide satisfying imaging in almost all cases, with less radiation, and avoiding complications inherent to the arterial puncture, reported in <1% of cases. 2900

4.5.2.7 Other tests

Several other non-invasive tests can be used routinely, either to localize the lesions or to evaluate their effect on limb perfusion: segmental pressure measurements and pulse volume record-2905 ings,²⁵¹ (laser) Doppler flowmetry, transcutaneous oxygen pressure assessment (TCPO₂), and venous occlusion plethysmography before and during reactive hyperaemia.²⁵²

Recommendations for diagnostic tests in patients with I FAD

Recommendations	Class ^a	Level ^b	Ref ^c	
Non-invasive assessment methods such as segmental systolic pressure measurement and pulse volume recording, plethysmography, Doppler flowmetry, and DUS are indicated as first-line methods to confirm and localize LEAD lesions.	I	В	251,252	2915
DUS and/or CTA and/or MRA are indicated to localize LEAD lesions and consider revascularization options.	I	A	237, 238, 241–250	
The data from anatomical imaging tests should always be analysed in conjunction with haemodynamic tests prior to therapeutic decision.	I	С	-	2925
ass of recommendation. vel of evidence. ferences. A = computed tomography angio	graphy; DUS :	= duplex ultra	asonography;	2930

LEAD = lower extremity artery disease; MRA = magnetic resonance angiography.

4.5.3 Therapeutic strategies

All patients with LEAD are at increased risk of further CVD events, and general secondary prevention is mandatory to improve prognosis. Patients with asymptomatic LEAD have no indication for 2940 prophylactic revascularization. The following paragraphs focus on the treatment of symptomatic LEAD.

4.5.3.1 Conservative treatment

The aim of conservative treatment in patients with intermittent 2945 claudication is to improve symptoms, i.e. increase walking distance and comfort. To increase walking distance, two strategies are currently used: exercise therapy and pharmacotherapy.

4.5.3.1.1 Exercise therapy

In patients with LEAD, training therapy is effective in improving 2950 symptoms and increasing exercise capacity. In a meta-analysis²⁵³ including data from 1200 participants with stable leg pain, compared with usual care or placebo, exercise significantly improved maximal walking time, with an overall improvement in walking ability of \sim 50–200%. Walking distances were also significantly 2955 improved. Improvements were seen for up to 2 years. Best evidence comes from studies with a short period of regular and intensive training under supervised conditions.²⁵⁴ In a meta-analysis of eight trials collecting data from only 319 patients, supervised exercise therapy showed statistically significant and clinically relevant 2960 differences in improvement of maximal treadmill walking distance compared with non-supervised exercise therapy regimens (+150 m on average)²⁵⁵ In general, the training programme lasts for 3 months, with three sessions per week. The training intensity on the treadmill increases over time, with a session duration of

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30-60 min.²⁵⁶ Of note, in a small randomized trial²⁵⁷ comparing supervised exercise therapy with usual care, while no significant changes in peak cardiovascular measurements were noted after 12 weeks of exercise, patients under supervised exercise therapy were more efficient in meeting the circulation and ventilation demands of exercise.

Individuals with LEAD should undertake exercise as a form of treatment. Any type of regular exercise should be continued after completion of the intensive training programme. Daily walking, or repeated series of heel raising or knee bending, are realistic possibilities.²⁵⁸ Other training programmes have been

2975 listic possibilities.²⁵⁸ Other training programmes have been suggested, but their effectiveness is less well documented. In a pilot trial, dynamic arm exercise training was followed by similar improvement (pain-free and maximal walking distance) to that seen with treadmill walking exercise training.²⁵⁹

2980 There are obvious limitations to training therapy. Muscular, articular, or neurological diseases may be limiting factors. General cardiac and/or pulmonary diseases can decrease capacity to achieve a level of training that is sufficient to obtain positive results. In conjunction with practical aspects, such as difficulties

- 2985 in attending the sessions or neglecting continuous training, the actual results in the clinical setting have often been poorer than in trials. Patients with Fontaine class IV should not be submitted to regular exercise training.
- 2990 4.5.3.1.2 Pharmacotherapy

Several pharmacological approaches were claimed to increase walking distance in patients with intermittent claudication. However, objective documentation of such an effect is often lacking or limited. In terms of walking distance improvement, the

2995 benefits, if any, are generally mild to moderate, with wide confidence of intervals. Also, mechanisms of action are diversified and often unclear. The drugs with best proof of efficacy are discussed briefly below. Among them, the best-documented drugs are cilostazol and naftidrofuryl.

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4.5.3.1.2.1 Cilostazol

Cilostazol is a phosphodiesterase-3 inhibitor. In a pooled analysis of nine trials (1258 patients) comparing cilostazol with placebo,²⁶⁰ this drug was associated with an absolute improvement of +42.1 m vs. placebo (P < 0.001) over a mean follow-up of 20 weeks. In another meta-analysis,²⁶¹ maximal walking distance increased on average by 36 m with cilostazol 50 mg/day, and almost twice (70 m) with the 100 mg dose. Improvement in quality of life is also reported in claudicants.²⁶² Owing to its pharmacological properties, it should be avoided in the case of heart failure. The most frequent side effects are headache, diar-

4.5.3.1.2.2 Naftidrofuryl

rhoea, dizziness, and palpitations.

Naftidrofuryl has been available in Europe for many years. It is a 5-hydroxytryptamine type 2 antagonist that reduces erythrocyte and platelet aggregation. The efficacy of naftidrofuryl was examined in a meta-analysis of five studies including 888 patients: pain-free walking distance was significantly increased by 26% vs.
placebo.²⁶³ This positive effect on intermittent claudication was confirmed by a recent Cochrane analysis.²⁶⁴ Quality of life was

also improved with naftidrofuryl treatment.²⁶⁵ Mild gastrointestinal disorders are the most frequently observed side effect.

4.5.3.1.2.3 Pentoxifylline

This phosphodiesterase inhibitor was among the first drugs to show improvement in red and white cell deformability, and, as a consequence, decrease blood viscosity. In a recent meta-analysis²⁶¹ of six studies including 788 patients, a significant increase in maximal walking distance was found with pentoxifylline (+59 m). 3030

4.5.3.1.2.4 Carnitine and propionyl-L-carnitine

These drugs are likely to have an effect on ischaemic muscle metabolism. In two multicentre trials,^{266,267} propionyl-L-carnitine improved walking distance and quality of life better than placebo. Additional trials are expected to evaluate their efficacy in large ³⁰³⁵ groups of patients.

4.5.3.1.2.4 Buflomedil

Buflomedil may cause inhibition of platelet aggregation and improve red blood cell deformability. It also has α -1 and α -2 adrenolytic effects. In a recent placebo-controlled study in 2078 patients,²⁶⁸ significant symptomatic improvement was shown. However, in a recent meta-analysis,²⁶⁹ these results were quoted as 'moderately' positive, with some degree of publication bias. The therapeutic dose range is narrow, with a risk of seizures.²⁷⁰ Buflomedil has been recently withdrawn from the market in some European countries for potential major side effects and uncertain benefits.

4.5.3.1.2.5 Antihypertensive drugs

In a recent review, antihypertensive drugs did not differ in respect 3050 of their effect on intermittent claudication. 271 According to a recent meta-analysis of four studies, the benefits of ACE inhibitors on walking distance are uncertain, and the main expectation of prescribing this drug class is in the general prognostic improvement of these patients (see Section 3.4.4). 272 Notably, β -blockers do not 3055 exert a negative effect on claudication. 273,274

4.5.3.1.2.6 Lipid-lowering agents

Beyond the evidence that statins improve the cardiovascular prognosis of patients with LEAD, several studies reported preliminary positive effects of statins on intermittent claudication.²⁶¹ The increase in maximal walking distance reported varied, on average, from 50 to 100 m. In one meta-analysis, the pooled effect estimate was in favour of lipid-lowering agents, with a relevant increase in maximal walking distance of 163 m.²⁶¹ 3065

4.5.3.1.2.7 Antiplatelet agents

The use of antiplatelet drugs is indicated in patients with LEAD to improve event-free survival (see Section 3.4.3). In contrast, data on the potential benefits of antiplatelet drugs to improve clinical symptoms are scarce. In a recent meta-analysis,²⁶¹ data from ³⁰⁷⁰ studies assessing five drugs (ticlopidine, cloricromene, mesoglycan, indobufen, and defibrotide) were pooled, with a significant increase in maximal walking distance of 59 m. Available data are too disparate to formulate any conclusions.

4.5.3.1.2.8 Other therapies

Other pharmacological agents assessed are inositol, proteoglycans, and prostaglandins. Although positive, the results require further

- confirmation. A recent meta-analysis showed no significant improvement in walking distance with gingko biloba.²⁷⁵ 3080
 - Intermittent pneumatic compression may be a relevant treatment for symptomatic LEAD. In a review,²⁷⁶ concordant data are reported in several studies showing increased flow (13-240%) in the popliteal or infragenicular arteries. Rest pain and walking
- distance were also improved. In a recent small, randomized trial 3085 comparing a portable intermittent pneumatic compression device with best medical therapy, maximal walking distance improved by 50% (90 m).²⁷⁷
- 4.5.3.2 Endovascular treatment of lower extremity artery disease 3090 Endovascular revascularization for the treatment of patients with LEAD has developed rapidly during the past decade, and a great number of patients can now be offered the less invasive treatment option. An increasing number of centres favour an endovascular-
- first approach due to reduced morbidity and mortality-compared 3095 with vascular surgery—while preserving the surgical option in case of failure.

The optimal treatment strategy concerning endovascular vs. surgical intervention is often debated due to the paucity of randomized

- studies; furthermore, most of these studies are underpowered. 3100 Moreover, owing to the rapid development, a thorough evaluation of new endovascular treatment options within adequately designed clinical studies is difficult. Another problem is the lack of uniform endpoint definitions, making a direct comparison among studies dif-
- ficult.²⁷⁸ It is important to report results including clinical, morpho-3105 logical, and haemodynamic outcomes.

The selection of the most appropriate revascularization strategy has to be determined on a case-by-case basis in a specialized vascular centre in close cooperation with an endovascular specialist

and a vascular surgeon. The main issues to be considered are 3110 the anatomical suitability (Table 6), co-morbidities, local availability and expertise, and the patient's preference.

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While revascularization is obligatory in patients with CLI, the evidence of any long-term benefit of endovascular treatment over supervised exercise and best medical treatment is inconclusive,

- especially in patients with mild to moderate claudication.²⁷⁹ However, advances in the endovascular treatment of LEAD have prompted many physicians to consider more liberal indications for percutaneous intervention. Endovascular revascularization is also
- indicated in patients with lifestyle-limiting claudication when clinical 3120 features suggest a reasonable likelihood of symptomatic improvement and there has been an inadequate response to conservative therapy. In aortoiliac lesions, endovascular revascularization can be considered without initial extensive conservative treatment.
- The major drawback of endovascular interventions-compared 3125 with surgery—is the lower long-term patency. The primary patency after angioplasty is greatest for lesions in the common iliac artery and decreases distally, and with increasing length, multiple and diffuse lesions, poor-quality run-off, diabetes, and renal
- 3130 failure. Currently there is no established method—besides stent implantation-to improve at least the mid-term patency of angioplasty. The use of drug-eluting balloons seems promising; however, the current limited data do not justify a general recommendation.

In general, endovascular interventions are not indicated as pro-

phylactic therapy in an asymptomatic patient. Patients undergoing 3135

Table 6 Lesion classification according to the TransAtlantic Inter-Society Consensus for the Management of Peripheral Arterial Disease (TASC II)

	Aorto-iliac lesions	31
Lesion type	Description	
Туре А	 Unilateral or bilateral stenosis of CIA Unilateral or bilateral single short (≤3 cm) stenosis of EIA 	31
Туре В	 Short (≤3 cm) stenosis of infrarenal aorta Unilateral CIA occlusion Single or multiple stenosis totaling 3-10 cm involving the EIA not extending into the CFA Unilateral EIA occlusion not involving the origins of internal iliac or CFA 	31
Туре С	 Bilateral CIA occlusions Bilateral EIA stenoses 3-10 cm long not extending into the CFA Unilateral EIA stenosis extending into the CFA Unilateral EIA occlusion that involves the origins of internal iliac and/or CFA Heavily calcified unilateral EIA occlusion with or without involvement of origins of internal iliac and/ or CFA 	31
Туре D	 Infra-renal aorto-iliac occlusion Diffuse disease involving the aorta and both iliac arteries requiring treatment Diffuse multiple stenosis involving the unilateral CIA, EIA and CFA Unilateral occlusions of both CIA and EIA Bilateral occlusions of EIA Iliac stenosis in patients with AAA requiring treatment and not amenable to endograft placement or other laesions requiring open aortic or iliac surgery 	31
	Femoral-popliteal lesions	
Lesion type	Description	31
Туре А	 Single stenosis ≤10 cm in length Single occlusion ≤5 cm in length 	
Туре В	 Multiple lesions (stenoses or occlusions), each ≤5 cm Single stenosis or occlusion ≤15 cm not involving the infra geniculate popliteal artery Single or multiple lesions in the absence of continuous tibial vessels to improve inflow for a distal bypass Heavily calcified occlusion ≤5 cm in length 	31
	 Single popliteal stenosis Multiple stenoses or occlusions totaling >15 cm 	
Туре С	 with or without heavy calcifications Recurrent stenoses or occlusions that need treatment after two endovascular interventions 	31
Туре D	 Chronic total occlusion of CFA or SFA (>20 cm, involving the popliteal artery) Chronic total occlusion of popliteal artery and 	- 1

AAA = abdominal a ortic aneurysm; CFA = common femoral artery;CIA = common iliac artery; EIA = external iliac artery; SFA = superficial femoral artery.

After Norgren et al.⁶ with permission.

endovascular revascularization for claudication or CLI should be entered into a clinical surveillance programme.

The primary goals of stent implantation are: (i) to improve an 3195 insufficient primary result-residual stenosis, extensive recoil, flow-limiting dissection; and (ii) to improve long-term patency. The placement of stents should generally be avoided in bending areas (hip and knee joints), although special stents have been developed recently. Stent implantation should also be avoided in 3200 a segment suitable as a landing zone for a potential bypass.

4.5.3.2.1 Aortoiliac segment

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Obstructive atherosclerotic disease of the distal aorta and iliac arteries is preferentially treated with endovascular techniques, and an endovascular-first strategy can be recommended for all TransAtlantic Inter-Society Consensus (TASC) A-C lesions. Low morbidity and mortality as well as a >90% technical success rate justify the endovascular-first approach. In experienced centres, TASC D lesions are also primarily treated percutaneously. The main limitation in recommending the endovascular-first strategy for almost all aortoiliac lesions is the lack of published data from randomized trials.

The only randomized trial comparing primary stent implantation with provisional stenting in the case of a persistent pressure gradient after angioplasty alone did not demonstrate any benefit of 3215 primary stent implantation.²⁸⁰ Based on an older meta-analysis, stenting can be recommended as the primary therapy for common and external iliac stenosis and occlusions.²⁸¹ The patency rates with stenting of iliac arteries compare favourably with those of surgical revascularization.²⁸²

The choice of balloon vs. self-expandable stents is determined

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mainly by operator preference. The main advantages of balloonexpandable stents are the higher radial stiffness and the more accurate placement, which is especially important in bifurcation lesions.²⁸³ In the external iliac artery, a primary stenting strategy 3225 using self-expandable stents compared with provisional stenting is

preferred mainly due to a lower risk of dissection and elastic recoil. In the case of doubt about the haemodynamic significance of morphologically borderline iliac lesions, pressure gradients at rest and with induced hyperaemia should be measured.²⁸⁴

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Recommendations for revascularization in patients with aortoiliac lesions

A primary endovas 3240 patients with seven	cular approach may be	с
3240 considered in aorto patients with sever	· · · · · · · · · · · · · · · · · · ·	
an experienced tea	e comorbidities, if done by	с
	antation rather than g may be considered for IIb	с

^bLevel of evidence.

TASC = TransAtlantic Inter-Society Consensus.

4.5.3.2.2 Femoropopliteal segment

One of the main problems with endovascular therapy in this segment is the high prevalence of diffuse disease. Furthermore, different mechanical forces act on the superficial femoral artery. This artery is deformed repetitively in multiple directions by leg movements. A high technical success rate, due to technical devel- 3255 opments and increasing operator experience, in combination with low risk, make endovascular therapy the preferred choice also in patients with long and complex femoropopliteal lesions.

The landscape of endovascular treatment of femoropopliteal disease has changed decisively with the development of self- 3260 expandable nitinol stents. The previous strategy was to use stents as the treatment option only in the case of initial PTA failure or late recurrence. However, according to an increasing number of randomized studies, primary nitinol stenting can now be recommended as the first-line treatment for intermediate 3265 length superficial femoral artery lesions due to improvement of at least mid-term patency.^{285,286} The restenosis rate after 1-2years is 20-30% lower after primary stenting compared with angioplasty.

The decision to stent the superficial femoral artery is based 3270 mainly on the clinical indication for revascularization and on the lesion length and complexity. In the case of CLI, stenting can be applied more liberally for limb salvage and ulcer healing.

In the past, there was much concern about stent fractures. Several risk factors have been identified for stent fractures: 3275 number and length of implanted stents, overlapping stents, amount of calcification, and deployment technique.²⁸⁷ The higher fracture resistance of the latest generation of stents in combination with the production of long nitinol stents (up to 20 cm in length) broadens the possibilities of endovascular therapies in the case of 3280 more difficult and complex lesions.

In-stent restenosis is the major drawback of stent implantation. To date there is no proof of any impact of stent design on restenosis rates. Isolated balloon angioplasty of restenosis lesions has a very high failure rate. Other treatment modalities have been inves- 3285 tigated, but there is no single randomized trial in patients with in-stent restenosis demonstrating the superiority of one technique over the other. Drug-eluting stents have been investigated in a few studies in the superficial femoral artery, and until now no advantage has been shown compared with bare-metal nitinol stents.²⁸⁸ 3290 Early studies with drug-eluting balloons in the femoropopliteal arteries showed improved short-term patency rates compared with plain balloon angioplasty.²⁸⁹

Covered stents (stent grafts) appear to be a viable option for the treatment of complex superficial femoral artery lesions, with out- 3295 comes comparable with prosthetic above-knee femoropopliteal bypass surgery.²⁹⁰

Despite its widespread use, research data regarding subintimal angioplasty are sparse. There are no data comparing patency rates between intraluminal and subintimal angioplasty. However, in 3300 many interventions an unintentional subintimal passage is unavoidable. Regarding atherectomy, different devices are used with unclear long-term benefits. Currently there are niche indications in severely calcified lesions and non-stent areas (e.g. the common femoral and popliteal artery). However, there are some concerns 3305 regarding the risk of distal embolization with these devices.

	_				
3310		Recommendations	Class ^a	Level ^b	R ef ^c
3315		When revascularization is indicated, an endovascular-first strategy is recommended in all femoropopliteal TASC A–C lesions.	I	С	-
		Primary stent implantation should be considered in femoropopliteal TASC B lesions.	lla	A	285, 286, 291
3320 3325		A primary endovascular approach may also be considered in TASC D lesions in patients with severe comorbidities and the availability of an experienced interventionist.	Шь	С	-
	٥L	lass of recommendation. evel of evidence. eferences.			

Recommendations for revascularization in patients with femoropopliteal lesions

4.5.3.2.3 Infrapopliteal arteries

TASC = TransAtlantic Inter-Society Consensus.

Most patients with CLI have multisegmental disease involving the infrapopliteal arteries. Therefore, limb salvage is the primary indi-3335 cation for endovascular treatment of infrapopliteal lesions, while angioplasty of these arteries is usually not indicated in patients with intermittent claudication. There is increasing evidence to support a recommendation for angioplasty in patients with CLI where straight-line flow to the foot in at least one lower leg 3340 artery can be re-established according to the pre-interventional

angiogram and in the case of important co-morbidities.²⁹²

Primary PTA remains the standard of care, as it provides an

acceptable clinical outcome at a low procedural cost.²⁹³ The

limb salvage rate is definitely higher than the angiographic

patency rate after initially successful intervention below the knee.

Therefore, long-term patency is not obligatory in CLI patients

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Recommendations for revascularization in patients

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with infrapopliteal lesions

Recommendations	Class ^a	Level ^b
When revascularization in the infrapopliteal segment is indicated, the endovascular-first strategy should be considered.	lla	с
For infrapopliteal lesions, angioplasty is the preferred technique, and stent implantation should be considered only in the case of insufficient PTA.	lla	с

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PTA = percutaneous transluminal angioplasty.

with persistent clinical improvement. Stent implantation in infrapopliteal vessels is generally reserved for cases with a suboptimal 3365 outcome after PTA. The use of drug-eluting stents is associated with a favourable restenosis rate²⁹⁴; the balloon-expandable sirolimus-eluting stent is approved in Europe for this indication.

4.5.3.3 Surgery

Vascular surgery offers different revascularization techniques for lower limb ischaemia. Bypass surgery presents the most common surgical approach for diffuse occlusive disease and creates new conduits following anatomical or extra-anatomical routes. In some circumstances, local endarterectomy with or without patching can 3375 restore blood perfusion. Different graft materials can be applied. Autologous vein or artery grafts are the best options, but are not always available or applicable. In such cases, prosthetic grafts are considered. Homografts represent the third option for vascular substitution, especially in the case of infective complications. 3380

Patients with extensive necrosis or infectious gangrene and those who are non-ambulatory may best be served with primary amputation. Amputation remains the last surgical step to solve irreversible limb ischaemia, allowing patient recovery with rehabilitation and prosthesis. For a moribund patient, adequate analgesia and other 3385 supportive measures may also be the best option. Other adjuvant surgical options can be considered. Skin reconstruction is useful to cover large areas of lost tissue. The use of lumbar sympathectomy is controversial and is not supported by evidence.

4.5.3.3.1 Aortoiliac disease

Aorto-biiliac or -bifemoral bypass is usually recommended for diffuse aortoiliac disease. In some situations, when an abdominal approach is perilous, a modified retroperitoneal approach or a unilateral bypass with a femoro-femoral cross-over may be considered. Other extra-anatomical surgical alternatives are axillo(bi)femoral or thoracic(bi)femoral bypasses. The surgical strategy depends on the lesion location and technical possibilities. Compared with the aortofemoral bypass, extra-anatomical bypasses present poorer patency rates and higher risk of complications. The 10-year primary patency rates of aortobifemoral bypass range from 80% to 90%.²⁹⁵

4.5.3.3.2 Infrainguinal disease

When infrainguinal disease is the cause of claudication, the appropriateness of intervention is more debated than for aortoiliac disease, depending on the level of symptoms, quality of femoral 3405 profundis artery and its collaterals, and local haemodynamic status. In contrast, in the case of CLI, any patent proximal vessel, including the iliac, common, or superficial femoral arteries, femoral profundis, and popliteal arteries, may serve as the inflow vessel for distal arterial reconstruction. Autologous vein grafts (in 3410 situ or reversed vein graft, or using the contralateral saphenous vein) provide the best patency results.²⁹⁶ Prosthetic grafts may be used if the autogenous vein is not available. Conflicting results are reported on the usefulness of vein cuffs to improve graft patency.^{297,298} In a recent meta-analysis²⁹⁹ involving data from 3415 seven contemporary trials (1521 patients) comparing Dacron with polytetrafluoroethylene femoropopliteal bypasses, the cumulative primary patency rates were similar at 3 years (60.2% vs. 53.8%, respectively) and at 5 years (49.2% vs. 38.4%). Pooling the three studies that included exclusively above-knee femoropopliteal 3420

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bypasses revealed lower risk for primary occlusion with Dacron grafts (HR 0.71 vs. polytetrafluoroethylene, P = 0.003), but long-term results are awaited. The pooled weighted data for 1-, 3-, and 5-year primary patency rates for femorodistal (tibial or

- 3425 pedal) bypasses are, respectively, reported at 85, 80, and 70% for venous bypass and 70, 35, and 25% with a prosthetic graft.⁶ In one trial with above-knee grafting, the 4-year primary and secondary patency rates were significantly better with the use of the saphenous vein (73% and 90%, respectively) compared with
- 3430 polytetrafluoroethylene (47% and 47%, both P < 0.05) and Dacron (54% and 60%, both P < 0.01). Two trials comparing *in situ* and reversed saphenous vein grafts to the above- and belowknee popliteal artery revealed no differences in primary and secondary patency as well as survival with an intact limb. Three
- 3435 trials comparing polytetrafluoroethylene with human umbilical vein showed significantly higher secondary patency rates with the latter.³⁰⁰ Comparison of polytetrafluoroethylene grafts with and without a vein cuff found no difference in above-knee grafts. However, primary patency for below-knee bypass was higher 3440 with a polytetrafluoroethylene prosthesis with vein cuff bypass at
- 2 years.^{296,301}

Only one randomized trial has compared angioplasty with infrainguinal bypass. In the Bypass versus Angioplasty in Severe Ischaemia of the Leg (BASIL) trial, 452 patients with severe limb ischaemia due to infrainguinal disease were randomized to angioplasty or infrainguinal bypass. The primary endpoint was amputation-free survival. Secondary endpoints included all-cause mortality, morbidity, reintervention, quality of life, and hospital costs.³⁰² The 30-day mortality was similar in both groups (5%

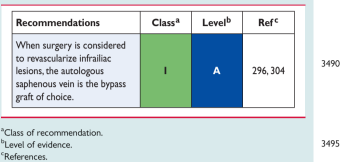
- for surgery and 3% for angioplasty). However, surgery was associated with a higher morbidity (57% vs. 41%), mainly due to myocardial infarction and wound infection. Moreover, surgery was more expensive during the first year, due to the longer hospital stay. The 6-month amputation-free survival was similar in both strat-
- egies. Angioplasty patients presented higher failure rates (20% vs. 3% at 1 year), resulting in higher reintervention rates (27% vs. 17%). These results suggest that surgical revascularization is superior to angioplasty in patients with good quality veins for bypass. Recently additional data with a longer follow-up period
- 3460 (>3 years) have been published^{211,303}: overall, there was no significant difference in amputation-free or overall survival between the two strategies. However, for patients who survived for at least 2 years after randomization, the surgery-first revascularization strategy was associated with a significant increase in subsequent overall

3465 survival and a trend towards improved amputation-free survival. One small, randomized trial comparing stenting with femoral-to-above-knee prosthetic bypass found no difference in primary and secondary patency rates at 12 months.²⁹⁰ Further trials are required comparing infrainguinal stenting with surgery.

- 3470 Another infrainguinal surgical reconstruction is the profundoplasty, the correction of a stenosis at the origin of the deep femoral artery. It may be considered as an inflow procedure, instead of a distal bypass, in the presence of an excellent proximal inflow, >50% stenosis of the proximal third of the profunda femoris artery, and the 3475 presence of excellent collateral flow to the tibial vessels.
 - Secondary amputation should be performed when revascularization has failed and reintervention is no longer possible or when the

limb continues to deteriorate because of infection or necrosis despite a patent graft. The goals of secondary amputation are: ischaemic pain relief, complete removal of diseased, necrotic, or 3480 infected tissue, and construction of a stump suitable for ambulation with prosthesis.

Recommendation for surgical revascularization in patients with LEAD



LEAD = lower extremity artery disease.

4.5.3.3.3 Surveillance

Clinical surveillance including clinical assessment and ankle ³⁵⁰⁰ pressure follow-up should be performed following any revascularization procedure. Although there is no consensual protocol of surveillance, regular monitoring of revascularized limbs can permit a prompt prophylactic intervention (e.g. repair of an arterial bypass at high risk of occlusion according to DUS criteria) and improve long-term patency.³⁰⁵ However, in a multicentre randomized trial including 594 patients with vein grafts, a systematic duplex surveillance programme was not found to be beneficial in terms of graft patency and limb survival rates, and was less cost-effective than clinical surveillance.³⁰⁶ DUS could be useful to select high-risk prosthetic grafts, which may require long-term anticoagulation to reduce the risk of graft thrombosis,³⁰⁷ but these data are based on observational series and require confirmation in trials.

4.5.3.3.4 Antiplatelet and anticoagulant therapy after revascularization 3515 Beyond potential benefits of antiplatelet agents to reduce fatal or non-fatal CVD events in patients with LEAD, these drugs are also specifically proposed after revascularization to improve patency rates. In a meta-analysis of 16 studies, the effect of antiplatelet therapy administered post-operatively was evaluated in 3520 patients receiving infrainguinal bypasses.³⁰⁸ Antiplatelet treatment with aspirin or a combination of aspirin and dipyridamole had an overall positive effect on primary patency 12 months after the procedure (OR 0.59, 95% CI 0.45-0.79). Subgroup analysis indicated that patients receiving a prosthetic graft were more likely to 3525 benefit from administration of platelet inhibitors than patients treated with venous grafts.³⁰⁸ The multicentre, prospective Dutch Bypass Oral Anticoagulants or Aspirin (BOA) trial³⁰⁹ randomized 2690 lower extremity bypass patients into two groups: anticoagulation (with the international normalized ratio targeted 3530 within the 3.0-4.5 interval) vs. antiplatelet therapy (aspirin 80 mg/day). Overall patency rates did not differ, but the results of a subgroup analysis suggested that oral anticoagulation improved vein graft patency compared with aspirin. Conversely, aspirin

- ³⁵³⁵ improved prosthetic graft patency vs. anticoagulation. Notably, the risk of major bleeding was two-fold higher in the anticoagulation group. In another trial,³¹⁰ 665 patients undergoing femoropopliteal bypass were randomized to aspirin (325 mg/day) plus warfarin (goal international normalized ratio 1.4–2.8) vs. aspirin (325 mg/
- day) alone. This trial failed to demonstrate any improvement in terms of graft patency with dual therapy. However, the results were in favour of combination therapy for patients with prosthetic bypasses. The haemorrhagic risk doubled when warfarin was added to aspirin. In another randomized study,³¹¹ warfarin (inter-
- national normalized ratio 2.0–3.0) plus aspirin (325 mg/day) was compared with aspirin (325 mg/day) alone in 56 patients with highrisk vein grafts (defined as poor arterial run-off, suboptimal vein conduit, and repeat interventions). At 3 years, patency and limb salvage rates were significantly higher in those receiving warfarin
- and aspirin, with in turn higher bleeding rates with this combination. Recently, the Clopidogrel and Acetylsalicylic Acid in Bypass Surgery for Peripheral ARterial disease (CASPAR) randomized double-blind trial assessed the efficacy of aspirin plus clopidogrel vs. aspirin alone to increase primary patency, limb salvage, and
- 3555 survival in patients receiving a below-knee bypass graft.³¹² Among the 851 patients enrolled, almost 70% had a venous graft and 30% a prosthetic graft. After a mean follow-up of 1 year, no overall difference was found regarding the combined primary outcome between the two groups. Subgroup analysis was in favour of a ben-
- eficial effect of clopidogrel in association with aspirin in prosthetic grafts. The number needed to treat using the dual antiplatelet therapy to save one limb after below-knee surgery was dramatically low, estimated at 10.2 patients.
- The role of anticoagulation after infrainguinal balloon PTA and stenting has been assessed in three prospective randomized trials.³¹³ None of these trials showed any significant improvement in arterial patency with the use of anticoagulation therapy, while bleeding complications increased.³¹³ Yet, anticoagulation therapy cannot be recommended routinely after lower extremity PTA or stenting.

4.5.3.4 Stem cell and gene therapy for revascularization

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The development of novel therapies to stimulate neovascularization, known as therapeutic angiogenesis, is based on the use of angiogenic factors or stem cells to promote revascularization and remodelling of collaterals with the aim of ameliorating symptoms and preventing amputation.

While several trials reported relief of ischaemic symptoms, functional improvement, and prevention of amputation,^{314–317} others failed to confirm this early promise of efficacy.^{318–320}

For autologous cell transplantation in humans, bone marrow and peripheral blood are rich sources of stem and progenitor cells. Bone marrow is currently the most frequent source of cells used for clinical repair trials, because it is easy to obtain and no

- 3585 complex purification steps are required. Another advantage is that it contains a variety of stem and progenitor cells with suggested superiority over one selected type of progenitor cell. With the many different cell types that can be used for stem cell therapy, it is not yet clear which ones are the most promising.³²¹
- 3590 In a recent meta-analysis of 37 trials, autologous cell therapy was effective in improving surrogate indexes of ischaemia, subjective

Recommendations for antiplatelet and anticoagulant therapy after revascularization

Recommendations	Class ^a	Level ^b	Ref ^c	
Antiplatelet therapy with aspirin is recommended in all patients with angioplasty for LEAD to reduce the risk of systemic vascular events.	I	с		3600
Dual antiplatelet therapy with aspirin and a thienopyridine for at least one month is recommended after infrainguinal bare-metal-stent implantation.	I	с		3605
Antiplatelet treatment with aspirin or a combination of aspirin and dipyridamole is recommended after infrainguinal bypass surgery.	I	A	308	3610
Antithrombotic treatment with vitamin K antagonists may be considered after autogenous vein infrainguinal bypass.	llb	В	309	3615
Dual antiplatelet therapy combining aspirin and clopidogrel may be considered in the case of below-knee bypass with a prosthetic graft.	Шь	В	312	3620

^aClass of recommendation. ^bLevel of evidence.

^cReferences.

LEAD = lower extremity artery disease.

symptoms, and hard endpoints (ulcer healing and amputation). Patients with thromboangiitis obliterans showed larger benefits than patients with atherosclerotic LEAD. The TAMARIS study is the largest randomized placebo-controlled trial of gene therapy in CLI, including >520 patients from 30 countries with CLI and skin lesions, unsuitable for standard revascularization. This study found no statistical difference between the two groups regarding the primary efficacy endpoint of death or first major amputation on the treated leg, whichever came first (37.0% vs. 33.2%, P = 0.48).³²² At present angiogenic gene and stem cell therapy are still being investigated and it is too early to give firm recommendations.

4.5.4 Management of intermittent claudication

The management of intermittent claudication consists of optimal 3640 risk factor control in order to improve the vital prognosis (see Section 3.4) and the symptoms. Therapeutic options to relieve symptoms are non-invasive (mostly exercise therapy and drug therapy) or invasive (revascularization). An algorithm for the management of intermittent claudication is proposed in *Figure 3*. With 3645 the increasing use of endovascular therapy to improve walking distance, there is an apparent need to compare it with 'supervised exercise training'. In a study of 51 patients with intermittent

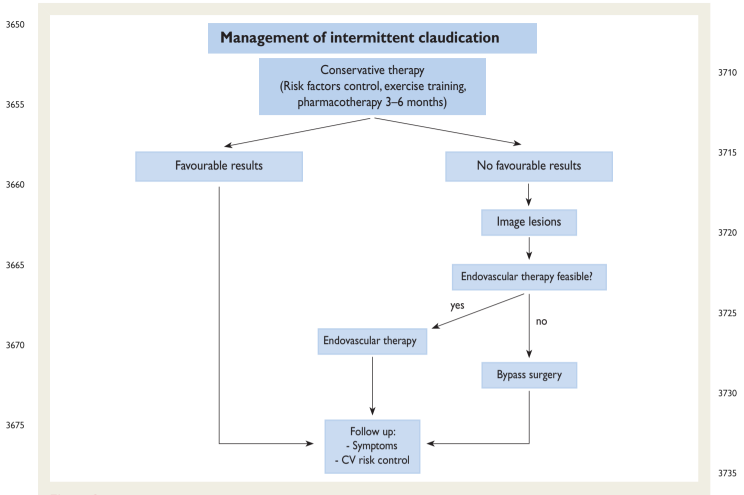


Figure 3 Algorithm for the management of intermittent claudication. CV = cardiovascular. 3680

claudication, there was no significant difference in walking distance or quality of life 2 years after treatment.³²³ More recently, a randomized controlled study initiated in 151 patients with intermittent claudication confirmed no difference in quality of life 12

- months after intervention. However, this study showed a higher cost for the endovascular intervention group.²⁷⁹ The adjuvant benefit of endovascular therapy to 'supervised exercise training' associated with best medical therapy has been assessed in patients
- 3690 with mild to moderate intermittent claudication.³²⁴ Although no difference in quality of life was reported in this study, at 24 months the improvement in walking distance in the angioplasty group was 38% greater than that in the control group in the case of femoropopliteal lesions, and 78% in the case of aortoiliac
- 3695 lesions. The ongoing Claudication: Exercise Versus Endoluminal Revascularization (CLEVER) trial will provide important insights into the indications of these therapeutic options in the management of patients with intermittent claudication.³²⁵

4.5.4.1 Medical treatment 3700

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In patients with intermittent claudication, the primary goal is to reduce the risk of CVD morbidity and mortality. This risk is present in all patients with LEAD, including those with mild, atypical, and even no symptoms.^{2,326} Therefore, the management and

3705 control of risk factors is necessary in every patient with LEAD, to reach the goals of secondary prevention. Among them, smoking cessation also provides the most noticeable improvement 3740 in walking distance when combined with regular exercise training, especially when lesions are located below the femoral arteries.

Symptoms can be improved by exercise training (preferably supervised) and drug therapy. Walking tests on the treadmill should be performed regularly to assess the evolution objectively. 3745 Patients should also be advised to keep a logbook to follow their home training and the evolution of their walking distance and symptoms. The logbook can help the patient adhere to medical advice. In the case of typical claudication, drug therapy to improve walking distance can be initiated.

In many patients with mild to moderate symptoms, these first steps will lead to a significant improvement in claudication and in quality of life. In this case, training (and eventually drug therapy) should be continued and the patients should be evaluated at regular intervals. ABI should be controlled periodically, although 3755 substantial functional improvement may not necessarily follow significant ABI change. The risk factor profile should be checked regularly and treatment adapted accordingly.

4.5.4.2 Interventional therapy

In severe cases with disabling claudication, medical therapy including 'supervised exercise training' is often insufficient to improve

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symptoms, and imaging of the lesions should be performed to define the exact location and characteristics of the lesions. This will help to decide whether interventional treatment is indicated

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and/or possible. Evidence for any long-term benefit of revascularization over supervised exercise and best medical therapy is inconclusive, especially in patients with mild to moderate claudication.³²⁴

- However, the expansion of endovascular therapy has prompted many physicians to consider more liberal indications for percutaneous intervention. The indications for endovascular revascularization also depend on the level of daily disability related to claudication, when clinical and imaging features suggest a reason-
- 3775 able likelihood of symptomatic improvement and there is insufficient response to exercise or pharmacological therapy. Owing to the limited probability of improvement in symptoms with exercise therapy in the case of aortoiliac lesions, revascularization should be considered without initial conservative treatment. Surgery is
- 3780 limited to extensive lesions without the possibility for endovascular treatment. The management of patients with intermittent claudication is summarized in *Figure 3*.

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Recommendations for patients with intermittent claudication

	Recommendations	Class ^a	Level ^b	Ref ^c
3790	Supervised exercise therapy is indicated.	I	A	255
3795	Non-supervised exercise therapy is indicated when supervised exercise therapy is not feasible or available.	I	С	-
	In patients with intermittent claudication with symptoms affecting daily life activity, drug therapy may be considered.	ШЬ	A	260-265, 269
3800	In the case of intermittent claudication with poor improvement after conservative therapy, revascularization should be considered.	lla	С	-
3805 3810	In patients with disabling intermittent claudication that impacts their activities of daily living, with culprit lesions located at the aorta/ iliac arteries, revascularization (endovascular or surgical)	lla	с	-
	should be considered as first- choice therapeutic option, along with the risk factor management.			
3815	Stem cell/gene therapy is not indicated.	ш	С	-

^aClass of recommendation. ^bLevel of evidence. ^cReferences.

4.5.5 Critical limb ischaemia (CLI)

4.5.5.1 Definition and clinical presentation

CLI is the most severe clinical manifestation of LEAD, defined as the presence of ischaemic rest pain, and ischaemic lesions or gangrene objectively attributable to arterial occlusive disease. It implies a chronic condition, to be distinguished from acute limb 3825 ischaemia (ALI) (see Section 4.5.6). An ankle pressure <50 mmHg is usually recommended as a diagnostic criterion because it includes most patients for whom rest pain or ischaemic lesions do not improve spontaneously without intervention. Since healing needs additional perfusion above that required for supporting intact skin, the ankle and toe pressure levels needed for healing are higher than the pressures found in ischaemic rest pain. For patients with ischaemic lesions or gangrene, CLI is suggested by an ankle pressure of <70 mmHg. Toe pressure <30 mmHg replaces the ankle pressure criteria in case of medial calcinosis.⁶ 3835 The investigation of the microcirculation (i.e. transcutaneous oxygen pressure) is also helpful in some cases, not only for diagnostic and prognostic purpose, but also sometimes to determine the level of amputation (Table 7).

Primary amputation rates range from 5% to 20%, mainly in 3840 patients unsuitable for revascularization, who are neurologically impaired or non-ambulatory.^{6,327} CLI is also a marker for generalized, severe atherosclerosis, with a three-fold risk excess of future myocardial infarction, stroke, and vascular death compared with patients with intermittent claudication.⁶ 3845

4.5.5.2 Therapeutic options

Comprehensive management requires multidisciplinary care to control atherosclerotic risk factors, provide revascularization as far as possible, optimize wound care, adapt shoe wear, treat infection, and initiate rehabilitation therapy (*Figure 4*).

The cornerstone of the management is arterial reconstruction and limb salvage.³²⁸ Revascularization should be attempted without delay in all patients presenting CLI, whenever technically possible. The screening or assessment of coronary or cerebrovascular diseases should not delay the management of patients with CLI if clinically stable. Medical baseline therapy including at least platelet inhibitors and statins should be initiated.^{329,330}

All patients with CLI should be referred to a vascular specialist early in the course of their disease, to plan revascularization. The most sig- 3860 nificant change in the treatment of CLI is the increasing tendency to shift from bypass surgery to less invasive endovascular procedures as an accepted first-choice revascularization strategy including tibial arteries, with bypass surgery reserved as a back-up option if necessary.⁶ The main advantages of endovascular revascularization are the 3865 low complication rates, ranging from 0.5% to 4.0%, high technical success rates (even in long occlusions) approaching 90%, and an acceptable short-term clinical outcome. The BASIL trial demonstrated that rates of amputation-free survival are similar for surgery and balloon angioplasty for at least 2 years after the procedure.^{302,331} 3870 The endovascular approach, including liberal use of stents above the knee level, is justified as long as low rates of complications are encountered and the surgical landing zone for the distal anastomosis of a potential secondary bypass remains unaffected by the interventional procedure. In patients with extensive foot gangrene or sepsis, open 3875 procedures possibly deliver more immediate pulsatile flow to the limb; however, the higher morbidity of surgery and the risk of graft

Assessment	Feature	Presentation to define CLI	Remarks
History	Duration of symptoms and clinical signs of CLI	>2 weeks	Needs morphine analgesics to be controlled
Symptoms	Rest pain	Toe, forefoot	Especially with elevation of limb (i.e. during night sleep). Calf pain/cramps do not constitute clinical presentation of CLI
	Ischaemic lesions	Periungual, toes, heel, over-bone prominences	
	Infection		Secondary complication: inflammation and infection
	Probe-to-bone test		Positive test identifies osteomyelitis with high specificity and sensitivity
Haemodynamics	Absolute ankle pressure	<50 mmHg or <70 mmHg	Plus rest pain Plus ischaemic lesion(s)
	Absolute great toe pressure	<30 mmHg	To be measured in the presence of medial calcinosis (incompressible or falsely elevated ankle pressure, ABI >1.40)
	Transcutaneous partial oxygen pressure	<30 mmHg	Estimation of wound healing, considerable variability

ABI = ankle-brachial index; CLI = critical limb ischaemia.

infection must be kept in mind.³³² Very distal venous bypass grafts to the pedal arteries are feasible and are characterized by an excellent patency rate of 88% at 4 years.^{333,334}

There are large discrepancies between the reported results of arterial reconstruction,³³⁵ mostly because of the inappropriate inclusion of patients with non-critical limbs in studies on CLI. Of

- note, there is a lower risk group consisting of patients with rest pain, and a higher risk group consisting of patients with true limb ischaemia with major tissue loss. At 1 year, 73% of patients in the low-risk group lost their leg or died if treated conservatively. For those patients fitting the high-risk criteria, 95% of those treated con-
- 3915 servatively required amputation within a year. In comparison, for those high-risk patients who received reconstruction, only 25% required major amputation.³³⁶ The primary efficacy endpoint of therapy is vascular reconstruction patency and limb salvage, whereas the patient-related main successful outcome includes main-
- 3920 tenance of ambulation and independence. Despite acceptable patency and limb salvage rates, reinterventions within 3 months and readmission to the hospital within 6 months occur in over a half of patients. Independent predictors of failure include impaired ambulatory status at presentation (HR 6.44), presence of infrainguinal disease
- (HR 3.93), ESRD (HR 2.48), and presence of gangrene (OR 2.40).³³⁷ In patients with CLI unsuitable for revascularization, the only drugs with some positive results within randomized studies are prostanoids.^{338,339} However, due to some divergent results in other studies, there is no conclusive evidence on effectiveness.³⁴⁰
- ³⁹³⁰ The safety and efficacy of various forms of therapeutic angiogenesis (gene or stem cell therapy) are promising, but robust evidence from RCTs is needed. The benefits of spinal cord stimulation are still debated, but a Cochrane review published in 2005 suggests some efficacy.³⁴¹

The management of patients with CLI is summarized in Figure 4.

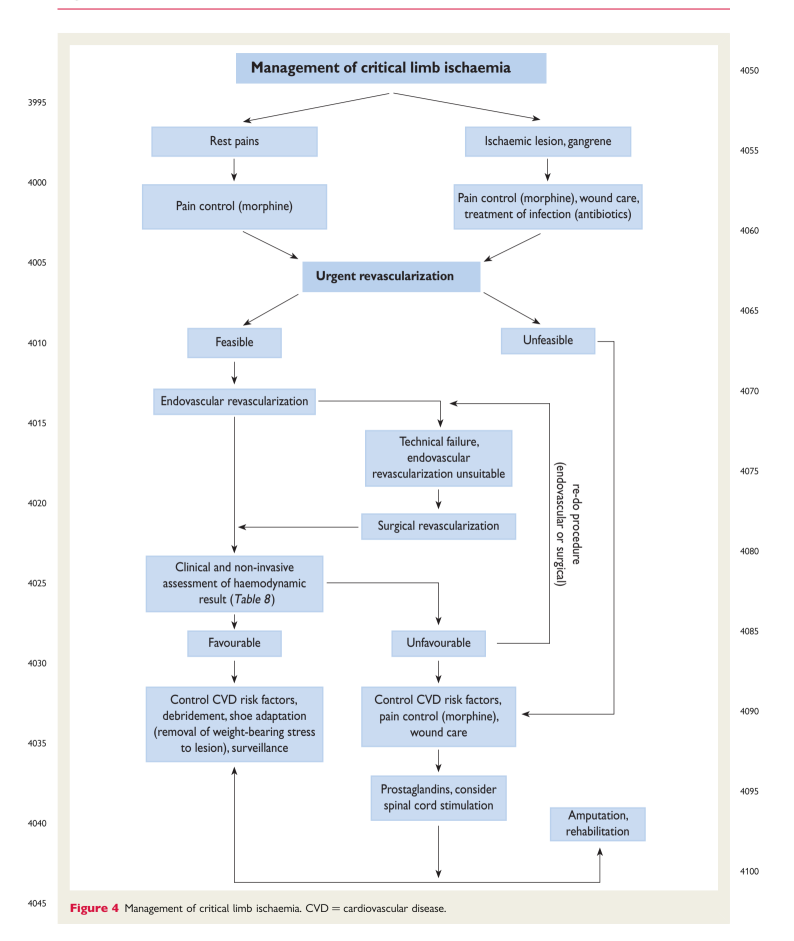
Recommendations for the management of critical limb ischaemia

Recommendations	Class ^a	Level ^b	Ref ^c	
For limb salvage, revascularization is indicated whenever technically feasible.	I	А	302, 331, 336	39
When technically feasible, endovascular therapy may be considered as the first-line option.	ШЬ	В	302, 331	37
If revascularization is impossible, prostanoids may be considered.	ШЬ	В	338, 339	39

4.5.6 Acute limb ischaemia (ALI)

ALI is related to a sudden decrease in arterial perfusion in the limb. Thrombotic or embolic causes can be involved. Artery disease 3985 progression, cardiac embolization, aortic dissection or embolization, graft thrombosis, thrombosis of a popliteal aneurysm, entrapment or cyst, trauma, phlegmasia cerulea, ergotism, hypercoagulable states, and iatrogenic complications related to cardiac catheterization, endovascular procedures, intra-aortic 3990 balloon pump, extra-corporeal cardiac assistance, as well as

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vessel closure devices are the potential causes of ALI. The viability 4105 of the limb is mostly threatened in this context. Quick and proper management is needed for limb salvage.

Once the clinical diagnosis is established, treatment with unfractionated heparin should be given.^{6,342} Analgesic treatment is often

- necessary. The level of emergency and the choice of therapeutic 4110 strategy depend on the clinical presentation, mainly the presence of neurological deficiencies, and the thrombotic vs. embolic cause. The clinical categories are presented in Table 8.
- An irreversible or unsalvageable extremity may require amputation before deterioration of the patient's clinical condition, although 4115 attempts are usually made to save the limb, or at least to limit the level of amputation. A viable limb mandates urgent imaging as well as the assessment of major co-morbidities. In the case of severely deteriorated renal function, detailed DUS imaging may replace
- angiography. In some cases, a clear cardiac embolization in poten-4120 tially normal arteries can be treated by surgical embolectomy without previous angiographic imaging. Otherwise, given the emergency level of care, angiography can be performed with no previous vascular ultrasound to avoid therapeutic delays.
- Different revascularization modalities can be applied (Figure 5). The 4125 options for guick revascularization consist of percutaneous catheterdirected thrombolytic therapy, percutaneous mechanical thrombus extraction or thromboaspiration (with or without thrombolytic therapy), and surgical thrombectomy, bypass, and/or arterial repair.
- The therapeutic strategy will depend on the type of occlusion (throm-4130 bus or embolus) and its location, duration of ischaemia, co-morbidities, type of conduit (artery or graft), and therapy-related risks and outcomes. Owing to reduced morbidity and mortality compared with open surgery, endovascular therapy is the initial treatment
- of choice, especially in patients with severe co-morbidities, if the 4135 degree of severity allows time for revascularization, and pending local availability of an emergency interventional team. Treatment results are best with an ALI duration < 14 days.⁶ Intra-arterial thrombolysis is the classic endovascular technique for thrombus removal. Various techniques and different thrombolytic agents are currently

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Table 8 Clinical categories of acute limb ischaemia

Grade	Category	Sensory loss	Motor deficit	Prognosis
I	Viable	None	None	No immediate threat
IIA	Marginally threatened	None or minimal (toes)	None	Salvageable if promptly treated
IIB	Immediately threatened	More than toes	Mild/ moderate	Salvageable if promptly revascularized
111	Irreversible	Profound, anaesthetic	Profound, paralysis (rigor)	Major tissue loss Amputation. Permanent nerve damage inevitable

Adapted from Rutherford et al., with permission.³²⁸

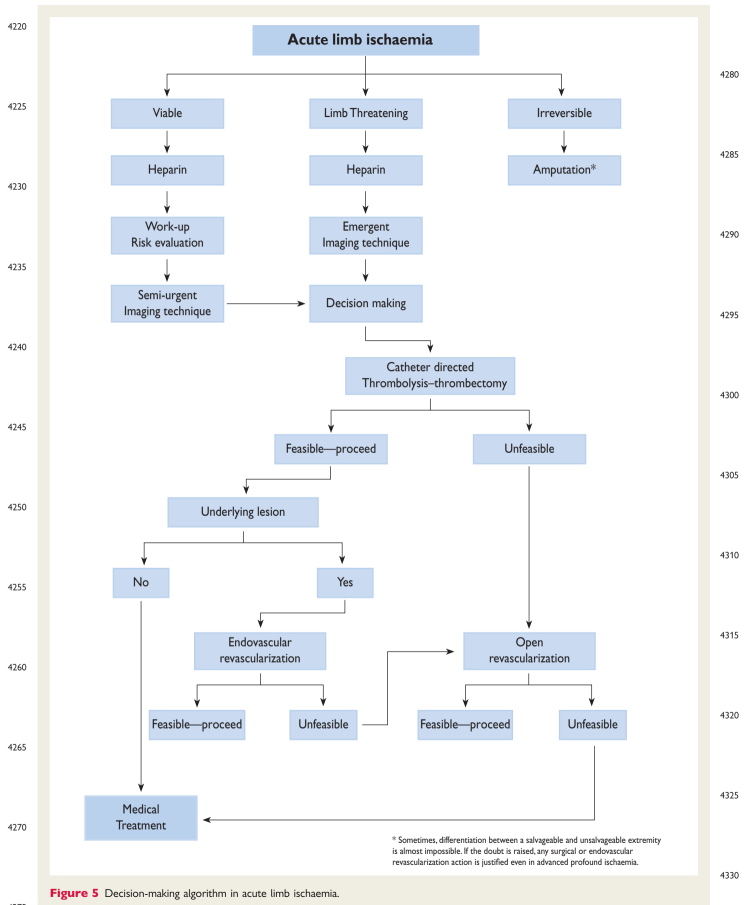
used.⁶ Intrathrombotic delivery of the thrombolytic agent is more effective than non-selective catheter-directed infusion. Different devices aiming at mechanical removal of the clot have been developed and are commonly used alone or in combination with thrombolysis, 4165 with the main advantage of decreasing delay to reperfusion. The modern concept of the combination of intra-arterial thrombolysis and catheter-based clot removal is associated with 6-month amputation rates <10%⁶ Systemic thrombolysis has no role in the treatment of patients with ALI. 4170

Based on the results of old randomized trials,³⁴³⁻³⁴⁵ there is no clear superiority of thrombolysis vs. surgery on 30-day mortality or limb salvage. Thrombolysis offers better results when applied within the first 14 days after the onset of symptoms. Thrombectomy devices have been proposed to treat ALI, but the benefits are not 4175 well documented. After thrombus removal, the pre-existing arterial lesion should be treated by endovascular methods or open surgery. Based on clinical presentation and availability of an emergency centre, surgical revascularization should be preferred when limb ischaemia is highly threatening and catheter-based treatment 4180 attempts may delay revascularization. Lower extremity fourcompartment fasciotomies are sometimes performed to prevent a post-reperfusion compartment syndrome, especially in the setting of class IIb and III ischaemia with surgical revascularization. In cases of viable limb, open or endovascular revascularization may not be 4185 possible, especially in the case of absent distal arteries, even after primary in situ thrombolysis; the only option then is to stabilize the ischaemic status with medical therapy (anticoagulation, prostanoids).

Recommendations	Class ^a	Level ^b	R ef ^c	
Urgent revascularization is indicated for ALI with threatened viability (stage II).	I	А	6, 342	
In the case of urgent endovascular therapy, catheter-based thrombolysis in combination with mechanical clot removal is indicated to decrease the time to reperfusion.	I	В	6, 304	
Surgery is indicated in ALI patients with motor or severe sensory deficit (stage IIB).	I.	В	304	
In all patients with ALI, heparin treatment should be instituted as soon as possible.	I	с	-	
Endovascular therapy should be considered for ALI patients with symptom onset <14 days without motor deficit (stage IIA).	lla	A	6, 304	

^aClass of recommendation ^bLevel of evidence. ^cReferences. ALI = acute limb ischaemia.

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4.6 Multisite artery disease

4.6.1 Definition

⁴³³⁵ Multisite artery disease is defined as the simultaneous presence of clinically relevant atherosclerotic lesions in at least two major vascular territories. Although patients with multisite artery disease are encountered regularly in clinical practice, no randomized trials have been designed to compare different treatment strategies,
 ⁴³⁴⁰ and the available data originate only from subgroup analyses or consecutive patient series.

The recent ESC/European Association for Cardio-Thoracic Surgery guidelines on myocardial revascularization offer for the first time specific recommendations for the management of

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patients suffering from CAD associated with carotid artery disease, renal artery disease, or LEAD.³⁴⁶

When dealing with a patient with multisite artery disease, one must focus attention not only on lesion sites and inherent technical difficulties related to specific treatment options, but also on the

- ⁴³⁵⁰ overall clinical status of the patient, taking into account the presence of cardiovascular risk factors and co-morbidities. Consequently, the treatment strategy should be chosen individually, based more on clinical rather than technical issues. A multidisciplinary team approach is required.
- ⁴³⁵⁵ The present guidelines address the impact of multisite artery disease on prognosis, as well as the screening for and management of multisite artery disease, taking into account the combinations most relevant for clinical practice.

4360 4.6.2 Impact of multisite artery disease on prognosis

- In patients with atherosclerotic disease in one vascular site, the presence of co-existing disease in a different vascular bed is associated with a higher risk of recurrent symptoms and complications in the first site. In fact, among 828 patients enrolled in the Framingham Study who had a myocardial infarction, those with a history of
- stroke or symptomatic LEAD had a two-fold increase in the risk of recurrent myocardial infarction.³⁴⁷ The REACH Registry enrolled 68 236 patients with either established atherosclerotic arterial disease (CAD, LEAD, cerebrovascular disease; n = 55 814) or
- three or more risk factors for atherothrombosis (n = 12 422).³⁴⁸ The incidence of cardiovascular death, myocardial infarction, stroke, or hospitalization for atherothrombotic events at 1 year increased with the number of symptomatic sites, ranging from 5.3% for patients with risk factors only to 12.6, 21.1, and 26.3% for patients with one,
- $_{\rm 4375}$ two, and three symptomatic sites, respectively (P < 0.001 for trend).¹ At 3 years, the rates of myocardial infarction/stroke/vascular death/rehospitalization were 25.5% for patients with symptomatic vascular disease in one vascular site vs. 40.5% for patients symptomatic in multiple vascular sites (P < 0.001).³⁴⁸ In a survey on 7783 out-
- 4380 patients who had experienced an atherothrombotic event, the rate of a first recurrent event at 1 year was almost doubled for patients with multisite disease vs. single disease location.³⁴⁹

4.6.3 Screening for and management of multisite artery disease

4385 4.6.3.1 Peripheral artery disease in patients presenting with coronary artery disease

Screening for and management of carotid artery disease, renal artery disease, and LEAD in patients presenting with CAD are addressed below.

4.6.3.1.1 Carotid artery disease in patients presenting with coronary 4390 artery disease

4.6.3.1.1.1 Carotid artery stenosis in patients not scheduled for coronary artery bypass grafting

In patients with CAD, the prevalence of severe carotid stenosis increases concurrently with the severity of CAD and is a known predictor of worse cardiovascular prognosis. Furthermore, a complex morphology of carotid plaque, such as echolucent plaque, is associated with heterogeneous coronary plaques and clinically unstable CAD. In a general review of cohorts with consecutive CAD patients enrolled without exclusion criteria,³⁵⁰ an average prevalence of >50, >60, >70, and >80% carotid stenosis was reported in 14.5, 8.7, 5.0, and 4.5% of patients, respectively. Thus, although the association between carotid artery stenosis and CAD is evident, the prevalence of significant carotid stenosis over the entire cohort is relatively low. Therefore, systematic carotid duplex screening is of limited value.

4.6.3.1.1.2 Carotid artery stenosis in patients scheduled for coronary artery bypass grafting

The question of prophylactic carotid revascularization in patients 4410 needing coronary artery bypass grafting (CABG) who also have a severe carotid artery stenosis arises from the higher risk of stroke in this population (*Table 9*).

4415 Table 9 Risk of stroke related to CABG **Patient category** Stroke risk (%) No carotid stenosis 1.4-3.8 4420 Unilateral >50% carotid stenosis 3.0 Bilateral >50% carotid stenosis 5.0 7.0 Carotid occlusion Previous stroke or TIA 85 4425

CABG = coronary artery bypass grafting; TIA = transient ischaemic attack. Modified from Blacker *et al.*³⁵¹

4.6.3.1.1.2.1 Screening for carotid stenosis in patients undergoing coron- 4430 ary artery bypass grafting

The prevalence of carotid stenosis in patients undergoing CABG varies in the literature, because of patient specificities, selection bias, DUS diagnostic criteria, and the extent of stenosis considered. Several studies attempted to identify clinical risk factors for the 4435 presence of significant carotid artery stenosis among patients undergoing planned CABG.³⁵² The most frequent risk factors are increasing age, history of cerebrovascular disease, or the co-existence of LEAD. Other risk factors mostly reported are female sex, multivessel CAD, and smoking. These risk factors are 4440 taken into consideration in the ESC/EACTS guidelines on myocardial revascularization.³⁴⁶ The criteria for screening carotid artery disease in patients undergoing CABG differ slightly from their expert-based recommendation, based on data from a study which assessed the efficacy of a clinical score to propose carotid 4445 DUS scanning in patients undergoing CABG.³⁵² The authors

identified four independent risk factors for carotid stenosis in candidates for CABG: age >70 years, neck bruit, history of cerebrovascular disease, and presence of clinical or subclinical LEAD. In a prospective assessment, they found that performing DUS scanning only in patients with at least one of these risk factors detected 100% of those with a carotid stenosis >70%, and decreased the number of useless scans by 40%. This approach does, however, need validation in a multicentre study.

Recommendations for screening for carotid artery

stenosis in patients undergoing CABG

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4460	Recommendations	Class ^a	Level ^b	Ref ^c
4465	In patients undergoing CABG, DUS scanning is recommended in patients with a history of cerebrovascular disease, carotid bruit, age ≥70 years, multivessel CAD, or LEAD.	I	В	352
4470	Screening for carotid stenosis is not indicated in patients with unstable CAD requiring emergent CABG with no recent stroke/TIA.	Ш	В	352

^aClass of recommendation. ^bLevel of evidence.

^cReferences

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CABG = coronary artery bypass grafting; CAD = coronary artery disease; DUS = duplex ultrasonography; LEAD = lower extremity artery disease; TIA = transient ischaemic attack.

4480 4.6.3.1.1.2.2 Management of carotid artery disease in patients undergoing coronary artery bypass grafting

It is unclear whether the benefits expected from CEA in the case of asymptomatic carotid artery stenosis are similar in those with concomitant CAD, and no specific randomized trial has been conducted in CAD patients with asymptomatic carotid stenosis. The

- ⁴⁴⁸⁵ Asymptomatic Carotid Atherosclerosis Study (ACAS) trial⁵³ found no interaction between perioperative outcomes after CEA and a history of myocardial infarction. A subgroup analysis of the ACST⁵⁴ observed long-term benefits with carotid surgery similar
- to those for the overall sample in the subset of 830 patients with CAD. However, the occurrence of stroke after CABG is multifactorial. In patients with carotid stenosis who undergo CABG without intervention on the carotid arteries, only 40% of postoperative strokes are ipsilateral to the carotid lesion. Besides,
- only a quarter of the strokes in patients with combined carotid and coronary surgery are exclusively ipsilateral to the stenotic carotid artery.³⁵³ In fact, the most common single cause of stroke after CABG is embolization with atherothrombotic debris from the aortic arch, while atrial fibrillation, low cardiac output, and hypercoagulation states resulting from tissue injury also con-
- tribute to the risk of stroke. Thus, the presence of carotid stenosis appears more as a marker of high risk of stroke after CABG rather

than the causal factor. Only those patients who have symptomatic carotid artery disease and those with asymptomatic bilateral 4505 carotid artery stenosis or unilateral carotid occlusion are definitely at higher risk of stroke during cardiac surgery, compared with patients without carotid artery stenosis.^{351,354}

Owing to the multitude of causes of stroke during CABG, prophylactic carotid revascularization before coronary surgery offers a 4510 partial solution for stroke risk reduction, at the expense of the risk related to the carotid revascularization itself, including the risk of myocardial infarction if carotid surgery is considered before coronary surgery in patients who often have severe CAD. Irrespective of whether the patient will undergo prophylactic carotid revasculari-4515 zation, the risk of stroke in these patients is overall higher than in the absence of CAD. The 30-day rate of stroke/death after combined (either synchronous or staged) CABG + CEA $^{353,355-363}$ or $CABG + CAS^{363-368}$ is >9% in most reports (ranging from 4.0%) to 19.2%). On the other hand, a recent study reported a 5-year 4520 rate of death/stroke or myocardial infarction as low as 8% after isolated CABG in low-risk patients with asymptomatic carotid stenosis >70%³⁶⁹ Thus, in the absence of clear proof that CEA or CAS is beneficial in patients undergoing CABG, all patients should be assessed on an individual basis, by a multidisciplinary team including 4525 a neurologist. Based on trials in patients with symptomatic carotid disease, it is reasonable to propose carotid revascularization (see Section 4.1.1.3.2) in patients scheduled for non-emergency CABG with recent (<6 months) TIA/stroke and symptomatic carotid stenosis, although those trials do not address the specific 4530 issue of patients undergoing coronary bypass.

Management of asymptomatic carotid stenosis should be delayed in cases of acute coronary events, because of increased rates of unstable carotid plaques concomitant to unstable CAD, with high perioperative risk of stroke in the case of carotid intervention.³⁵⁰ Selected 4535 patients with high-grade, asymptomatic carotid stenosis, particularly in the case of bilateral stenosis, may benefit from prophylactic carotid revascularization. The preoperative evaluation of such patients should include a detailed neurological examination, history aimed at unreported TIA symptoms, and a brain CT or MRI study 4540 to assess the presence of 'silent' ipsilateral infarcts.

Choice of carotid revascularization method in patients scheduled for coronary artery bypass grafting

Timaran et al. compared the in-hospital outcome of patients 4545 who underwent CAS before CABG with those who were treated by combined CEA and CABG between 2000 and 2004.³⁶³ During this 5-year period, 27 084 concurrent carotid revascularizations and CABGs were done. Of these, 96.7% underwent CEA-CABG, whereas only 3.3% (887 patients) had CAS-4550 CABG. Patients undergoing CAS-CABG had significantly lower rates of post-operative stroke (2.4% vs. 3.9%; P <0.001) and tended to have lower rates of combined stroke and death (6.9% vs. 8.6%; P = 0.1) compared with patients undergoing CEA-CABG, although in-hospital death rates were similar (5.2% vs. 4555 5.4%, respectively). After risk stratification, CEA-CABG patients had a 65% increased risk of post-operative stroke compared with patients undergoing CAS-CABG (OR 1.65, 95% CI 1.1-2.6; P = 0.02). However, no differences in the risk of combined

stroke and death were observed (OR 1.26, 95% Cl 0.9-1.6; P = not significant).

The most recent meta-analysis on the management of concomitant coronary and carotid artery disease was published by Naylor et al., in 2009.³⁷⁰ The results of different strategies (timing, revas-

- 4565 et al., in 2009.³⁷⁰ The results of different strategies (timing, revascularization modalities) are presented in *Table 10*. Of note, these results are stratified neither according to the coronary and neurological symptoms nor according to the severity of coronary and carotid artery disease.
- 4570 An overview of these results indicates no strong benefit of one strategy over another, although some need further studies to narrow their confidence intervals. Interestingly, the presence of carotid artery stenosis may lead to reconsideration of the technique of surgical coronary revascularization. Indeed, the
- 4575 co-existence of severe carotid disease in patients with CAD indicates widespread atherosclerosis with high risk for the presence of atherothrombotic lesions of the aortic arch, a risk factor for stroke. The avoidance of cross-clamping of the aorta during off-pump surgery may explain the lower rates of perioperative
- 4580 stroke when combined with CEA, although the number of patients subject to this strategy (n = 324) is too low to draw firm

Table 10Meta-analysis of cumulative results ofrevascularization strategies, with an indication forCABG and concomitant carotid revascularization

)	Strategy	Operative mortality (%)	Death ± any stroke/ TIA (%)	Death ± any stroke/ TIA ± MI (%)
	Synchronous CEA+CA	BG		÷
	CEA prebypass (n = 5386)	4.5 (3.9–5.2)	8.2 (7.1–9.3)	.5 (0. - 3.)
	CEA performed on bypass (n = 844)	4.7 (3.1–6.4)	8.1 (5.8–10.3)	9.5 (5.9–13.1)
	CEA+off-pump CABG (n = 324)	1.5 (0.3–2.8)	2.2 (0.7–3.7)	3.6 (1.6–5.5)
)	Staged CEA-CABG	·		÷
	CEA then CABG (n = 917)	3.9 (1.1–6.7)	6.1 (2.9–9.3)	10.2 (7.4–13.1)
;	CABG then CEA (n = 302)	2.0 (0.0–6.1)	7.3 (1.7–12.9)	5.0 (0.0–10.6)
	Staged CAS-CABG	·		·
	Staged CAS+CABG (n = 760)	5.5 (3.4–7.6)	9.1 (6.2–12.0)	9.4 (7.0–11.8)
)		1	1	1

 $\label{eq:CABG} CABG = \text{coronary artery bypass grafting; } CAS = \text{carotid artery stenting; } CEA = \text{carotid endarterectomy; } MI = \text{myocardial infarction; } TIA = \text{transient ischaemic attack.}$

Two other recent meta-analyses on CAS + CABG 371,372 provided similar results. Adapted from Naylor et al. 370

conclusions. Similarly, the higher risk of lesions of the aortic arch, a risk factor for stroke during catheterization of the carotid arteries, may explain why—although apparently less invasive— 4620 CAS does not present superior results to CEA in this situation. As expected, the staged approaches provide different myocardial and neurological protection, depending on the timing of the two interventions. This is probably the key issue when the staged approach is considered, and the neurological or myocardial risk 4625 may be prioritized according to the patient's clinical presentation as well as the level of severity of carotid and CADs.

Of note, in both the SAPPHIRE and CREST trials of CEA vs. CAS, the 30-day rate of myocardial infarction after carotid revascularization was significantly lower with CAS vs. CEA.^{79,98} Moreover, 4630 in a recent meta-analysis evaluating 2973 patients enrolled in randomized CAS vs. CEA trials, Wiesmann et al. reported a myocardial infarction rate of 2.3% with CEA vs. 0.9% with CAS (P = 0.03; OR 0.37).³⁷³ However, although CAS appears to be associated with a lower risk of periprocedural myocardial infarction com- 4635 pared with CEA, the overall data including death and stroke reported in Table 10 do not clearly favour one revascularization strategy over another. If CAS is performed before elective CABG, the need for double antiplatelet therapy usually delays cardiac surgery for ~ 5 weeks. Such deferral of CABG may 4640 expose the patient to the risk of myocardial infarction between CAS and CABG procedures (0-1.9%), and presents a major drawback of this treatment strategy.^{364,366,368} Recently, a few studies described the results of synchronous CAS + CABG, with CAS performed immediately before cardiac surgery.^{367,374} Such a strat- 4645 egy yielded a more favourable 4.0% 30-day rate of death or stroke.³⁷⁴ However, the bleeding risk during CABG, a factor predictive of long-term mortality, has not been considered extensively when comparing CAS with CEA concomitant (or before) to CABG. 4650

More details on the management of carotid stenosis in patients with CAD are given in Appendix 5.

4655

Recommendations for the management of carotid stenosis in patients undergoing CABG

	Recommendations	Class ^a	Level ^b		4660
	The indication for carotid revascularization should be individualized after discussion by a multidisciplinary team including a neurologist.	I	с		
	If carotid revascularization is indicated, the timing of the carotid and coronary interventions should be decided according to the clinical presentation, level of emergency, and severity of carotid disease and CAD.	I	С		4665
_			-	_	
C	lass of recommendation.				4670

^bLevel of evidence.

CABG = coronary artery by pass grafting; CAD = coronary artery disease.

4615

4585

Recommendations for carotid artery revascularization in patients undergoing CABG

	-			
		Recommendations	Class ^a	Level ^b
4680		In patients undergoing CABG, with a <6-mo TIA/stroke and corresponding carotid arter		y of
		Carotid revascularization is recommended in 70–99% carotid stenosis.	I	С
4685		Carotid revascularization may be considered in 50–69% carotid stenosis, depending on patient-specific factors and clinical presentation.	ШЬ	с
		Carotid revascularization is not recommended if the carotid stenosis is <50%.	ш	С
4690		In patients undergoing CABG with no histo stroke within 6 months	ry of TIA/	
4695		Carotid revascularization may be considered in men with bilateral 70–99% carotid stenosis or 70–99% carotid stenosis and a contralateral occlusion.	ШЬ	с
		Carotid revascularization may be considered in men with 70–99% carotid stenosis and ipsilateral previous silent cerebral infarction.	llb	с
4700	-			

4730

CABG = coronary artery bypass grafting; TIA = transient ischaemic attack.

4.6.3.1.2 Renal artery disease in patients presenting with coronary 4705 artery disease

In patients with CAD, RAS >50% is found in 10–20% of cases, mostly using renal angiography concomitant to cardiac catheterization, with almost a guarter being bilateral.^{13,375-380} These studies

- are concordant in reporting even higher rates in patients with 4710 triple-vessel CAD, as well as in those with hypertension or renal failure, although the use of contrast agents should be limited in patients with renal failure. Other situations where renal artery disease should be considered are recurrent episodes of heart failure and/or refractory angina, pulmonary oedema, and renal 4715
- function deterioration after the introduction of ACE inhibitors or angiotensin receptor antagonists.

In CAD patients with a suspicion of renal artery disease, as for any other patient, DUS should be used as the first-line non-invasive imaging test (see Section 4.4.3),^{171,172} even in the case of planned 4720 cardiac catheterization, in order to limit the use of ionized contrast agents and irradiation, and for cost issues. While CTA or MRA are usual second-line imaging tests, in the case of planned coronary angiography with a suspicion of renal artery disease after DUS (or poor quality imaging) in the absence of renal failure, renal 4725

angiography during the same procedure can be considered.

Although the co-existence of significant renal artery disease in patients with CAD is not negligible, a systematic screening for RAS does not appear reasonable because the management of these patients is barely affected. The use of systematic renal angioplasty has been challenged recently by the results of the ASTRAL trial¹⁹¹ (see Section 4.4.5.2), and there are no specific data for

patients who also suffer from CAD. Similarly, the presence of renal artery disease does not affect the management of patients with CAD, with the exception of renal failure after the use of ACE inhibitors or angiotensin II receptor antagonists. Yet, the indi- 4735 cations for screening renal artery disease in patients with CAD are similar to those for any patient.

4740 Screening for RAS in patients planned for coronary angiography

	Recommendations	Class ^a	Level ^b				
	DUS should be considered first in the case of clinical suspicion of renal artery disease in patients planned for coronary angiography.	lla	С		4745		
	Renal angiography concomitant to cardiac catheterization may only be considered in the case of persisting suspicion after DUS.						
^a C	lass of recommendation.	-	-		4750		

^bLevel of evidence.

DUS = duplex ultrasound; RAS, renal artery stenosis.

4755

4.6.3.1.3 Lower extremity artery disease in patients presenting with coronary artery disease

The co-existence of LEAD in CAD patients is associated with worse prognosis. In the REACH registry,¹ the 1-year rate of cardiovascular death/myocardial infarction/stroke/hospitalization for 4760 other atherothrombotic event(s) was 13.0% for patients with CAD alone, and 23.1% for patients with both conditions. LEAD is often under-recognized in CAD, as patients are largely asymptomatic; in patients with limiting angina, failure to recognize the condition may be because these patients exercise to a degree 4765 insufficient to evoke intermittent claudication. Therefore, a systematic approach, with ABI measurement, could lead to better identification of LEAD in patients with CAD.

In a cross-sectional study performed in primary care, ABI detected LEAD in 26.6% of 1340 patients with CAD and no 4770 other known location of atherothrombotic disease.³⁸¹ The prevalence of LEAD was increased significantly in patients with diabetes mellitus. Similar findings were reported in the Peripheral Arterial Disease Awareness, Risk, and Treatment: New Resources for Survival (PARTNERS) study.³⁸²

In different studies, the prevalence of ABI <0.90 can be estimated at 25-40% in patients hospitalized for CAD,³⁸³⁻³⁸⁵ while only <10% would be detected by clinical examination.³⁸⁶⁻³⁸⁸ Among patients with CAD, older age, intermittent claudication or atypical leg pain, smoking, diabetes, uncontrolled arterial hyper- 4780 tension, and elevated LDL cholesterol can be identified as factors suggestive of LEAD.

At any stage of CAD, the presence of LEAD is associated with a more severe and poorer prognosis. In 234 consecutive patients who underwent coronary angiography, Brevetti et al. found 4785 higher rates of multivessel CAD in patients with LEAD (60% vs. 20%, P < 0.01), which were associated with higher concentrations of C-reactive protein.³⁸⁹ In the Global Registry of Acute Coronary Events (GRACE), the in-hospital mortality of patients with acute

^aClass of recommendation. ^bLevel of evidence.

coronary syndromes (ACS) as well as the presence of cardiogenic
 shock was significantly higher in subjects with LEAD. At 6 months the rate of major cardiovascular events was 14.6% in patients with LEAD vs. 7.2% in those without.³⁹⁰ In the Monitoring of Trends and Determinants in Cardiovascular Disease (MONICA) study, the mortality rates in ACS were 18.8% and 13.1% in patients
 with vs. without LEAD, respectively.³⁹¹

The presence of LEAD is associated with a worse prognosis not only in patients with ACS but also in those with chronic stable angina as in the Coronary Artery Surgery Study (CASS), where the mortality rate was 25% higher in patients with PAD as compared with non-PAD patients, during a follow up of >10 years.³⁸⁶

4800 pared with non-PAD patients, during a follow up of >10 years.³⁸⁶ After percutaneous coronary intervention (PCI), patients with LEAD have a worse outcome. In a meta-analysis of eight studies, the HRs for 30-day, 6-month, and 1-year mortality were, respectively, 1.67, 1.76, and 1.46 (1.08–1.96) in patients with concomitant LEAD.³⁹² Similarly, the prognosis of CAD patients after CABG was

poorer in those with clinical or subclinical LEAD.^{393,394}

In summary, patients with LEAD associated with CAD are at twice the level of risk as those presenting with CAD alone. However, whether the management of CAD patients should differ in the case of concurrent LEAD is not obvious, because

- there are no specific trials related to this situation. To date, the co-existence of LEAD and CAD should only lead to closer attention, with a strict control of risk factors and the use of preventive treatments. Lowering the target for LDL cholesterol from 2.6 to
- 1.8 mmol/L should be considered. Regarding the use of antiplatelet therapy in stable CAD, given the greater benefits of clopidogrel vs. aspirin found in those with LEAD, clopidogrel rather than aspirin may be considered for the long-term treatment.³⁸ In a post-hoc analysis of the Clopidogrel for High Atherothrombotic Risk and
- 4820 Ischaemic Stabilization, Management and Avoidance (CHARISMA) study, there was a benefit of the combination of aspirin and clopidogrel in patients with LEAD.⁴⁰ Because of the post-hoc nature of this analysis, the benefit of such an approach needs confirmation. In the case of severe LEAD in CAD patients undergoing CABG,
- 4825 the use of venous bypass should be limited as far as possible, because this may lead to healing issues in the lower limbs, and because the venous material should be spared for potential *in situ* venous bypasses for the leg.
- 4830 4.6.3.2 Screening for and management of coronary artery disease in patients with peripheral artery disease
 Management of CAD in patients presenting with carotid disease and LEAD is addressed below.
- 4835 4.6.3.2.1 Screening for and management of coronary artery disease in patients presenting with carotid artery disease
 Few studies have systematically used coronary angiography to define the frequency of asymptomatic CAD in patients with carotid disease. In a landmark study performed over two decades ago, hae4840 modynamically relevant CAD was demonstrated in 40% of 200 patients while only 6% had absence of disease at angiography.³⁹⁸
 In a recent prospective investigation in 390 patients undergoing elective CAS, systematic coronary angiography showed the presence of one-, two-, and three-vessel disease and left main stenoses in 17, 15,
- 4845 22, and 7% of patients, respectively. Only 39% of the patients with significant coronary artery stenoses had cardiac symptoms.³⁹⁹

Recommendations for management of patients with LEAD and concomitant CAD

	Recommendations	Class ^a	Level ^b	Ref ^c		4050
	In patients with unstable CAD, vascular surgery should be postponed and CAD treated first, except when vascular surgery cannot be delayed due to a life- or limb-threatening condition.	I	С	-		4850 4855
	The choice between CABG and PCI should be individualized, taking into consideration the clinical presentation of CAD and LEAD, and comorbidities.	I	с	-		4860
	In the case of LEAD in patients with stable CAD, clopidogrel should be considered as an alternative to aspirin for the long-term antiplatelet therapy.	lla	В	38		4865
	In patients with CAD, screening for LEAD by ABI measurement should be considered.	lla	с	-		4870
	Prophylactic myocardial revascularization before high- risk vascular surgery may be considered in stable patients if they have persistent signs of extensive ischaemia or are at high cardiac risk.	ШБ	В	47, 395-397		4875
	lass of recommendation. evel of evidence.				_	
R	eferences. $M = apt kla_{\rm b} prachial index; CABC =$		tony hundre -	offing CAD -		4880

ABI = ankle-brachial index; CABG = coronary artery bypass grafting; CAD = coronary artery disease; LEAD = lower extremity artery disease; PCI = percutaneous coronary intervention.

The only study involving the management of patients undergoing 4885 CEA randomized 426 patients with no history of coronary disease and with normal cardiac ultrasound and electrocardiography into two groups, namely systematic coronary angiography and (if needed) revascularization, or no coronary angiography.⁴⁰⁰ No post-operative myocardial ischaemic events were observed 4890 among patients undergoing coronary angiography, while nine events were observed in the no-angiography group (P = 0.01).

In conclusion, patients with carotid stenosis have a high prevalence of CAD—even in the absence of cardiac symptoms—and are at risk of cardiovascular events. While CEA is considered as 4895 an intermediate-risk procedure, the cardiac risk associated with carotid revascularization may be lower with stenting than with endarterectomy.^{79,98} With respect to screening with coronary angiography and, if needed, coronary revascularization, before vascular surgery, the results of the four available randomized 4900 trials^{395–397,400}—none of them large-scale—have led to conflicting results, and no firm recommendation can be made at this point for patients undergoing carotid revascularization.

4.6.3.2.2 Screening for and management of coronary artery disease in patients presenting with lower extremity artery disease

4.6.3.2.2.1 Patients with lower extremity artery disease undergoing 4905 surgery

This topic has been addressed extensively in the ESC guidelines for pre-operative cardiac risk assessment and perioperative cardiac management in non-cardiac surgery.⁴⁷ Briefly, the goals of pre-

- operative screening are to ensure that the perioperative period 4910 is free of adverse cardiac events and to identify PAD patients with a poor long-term prognosis in whom treatment and risk factor modification may improve their outcome.
- In LEAD patients, screening offers the opportunity to initiate timely medication for secondary prevention of atherosclerotic 4915 disease; this improves both direct post-operative outcome and long-term survival. Factors that need to be considered in LEAD patients for screening, include:
 - (i) Emergency surgery: chronic cardiovascular medication should
- 4920 be continued during the procedure, and patients should be referred for surgery without delay.
 - (ii)Unstable cardiac conditions: deferral of the procedure and treatment of the underlying cardiovascular disease is recommended.
- 4925 (iii) Whether cardiovascular medications for secondary prevention of atherosclerosis (β-blockers, statins, ACE inhibitors, aspirin) are needed.
 - (iv) Whether work-up for the presence and extent of CAD is warranted.
- 4930 (v) How the results of the work-up will alter perioperative management.

The first step is to identify unstable cardiac conditions (ACS, arrhythmias, decompensated heart failure, severe valvular disease) that require immediate treatment. Patients with LEAD 4935 have a high risk for CAD: in a study of >1000 patients, only 8% had a normal angiogram.⁴⁰¹ Therefore, secondary prevention for atherosclerotic complications is recommended before high-risk surgery, including a low-dose, titrated β -blocker, statins, and

- aspirin. In patients with reduced left ventricular function, ACE 4940 inhibitors are recommended, according to the ESC guidelines.⁴⁷ Overall, the second step is to assess the level of surgical risk. However, peripheral vascular surgery is classified as high-risk surgery. The third step is to assess functional capacity. If the
- patient can achieve four or more metabolic equivalents without 4945 symptoms, then it is acceptable to proceed with surgery. Patients who have a functional capacity of less than four metabolic equivalents are at higher risk. A metabolic equivalent of less than four is equivalent to the inability to climb two flights of stairs or to run a
- short distance. Obviously, for patients with lower extremity arter-4950 ial insufficiency, this might not always be possible to assess. In patients with a low functional capacity, the cardiac risk of the procedure should be considered (Table 11).

Three randomized studies including patients with LEAD addressed the role of prophylactic coronary revascularization in 4955 stable patients scheduled for vascular surgery. The Coronary Artery Revascularization Prophylaxis (CARP) trial was the first to compare optimal medical therapy with revascularization (by either CABG or PCI) in patients with stable ischaemic heart

Table II Cardiac risk stratification for non-cardiac surgical procedures

High (reported cardiac risk often more than 5%) Aortic and other major vascular surgery Peripheral vascular surgery	4965
Intermediate (reported cardiac risk generally 1%–5%) Intraperitoneal and intrathoracic surgery Carotid endarterectomy Head and neck surgery Orthopaedic surgery Prostate surgery	4970
Low (reported cardiac risk generally less than 1%) Endoscopic procedures Superficial procedures Cataract surgery Breast surgery	4975
Ambulatory surgery	

From Poldermans et al., with permission.⁴⁷

disease prior to major vascular surgery.³⁹⁶ Of 5859 patients screened, 510 were randomized. Patients were included on the basis of a combination of cardiovascular risk factors and the detection of ischaemia on non-invasive testing. There was no difference 4985 in the primary endpoint of mortality at 2.7 years after randomization: 22% in the revascularization group vs. 23% in the no-intervention group. In addition, no difference in the rate of perioperative myocardial infarction was detected (12% vs. 14%, respectively). As a limitation, only a small proportion (8.9%) of 4990 screened patients were randomized, and patients with left main coronary disease were excluded by design from randomization.

DECREASE-V was a pilot study that applied a precise screening methodology and a more contemporary perioperative medical management.³⁹⁷ Patients at high risk for surgery underwent dobu-4995 tamine stress echocardiography or nuclear stress testing, and in the presence of extensive ischaemia were randomized to either revascularization or no revascularization. β-Blocker therapy was initiated and aspirin was continued during surgery in all patients. All patients (n = 101) had had a previous myocardial infarction, 5000 51% had ongoing angina, and 47% had congestive heart failure. Three-vessel or left main disease was present in 75% of cases and 43% had an ejection fraction of <35%. Both groups showed a very high 30-day death or myocardial infarction rate at 30 days (43% for revascularization vs. 33% for no revascularization; P =5005 not significant) and at 1 year (44% vs. 43%, respectively). The fact that all patients who were randomly assigned to the revascularization arm were compelled to undergo revascularization may have increased the risk associated with revascularization in patients with anatomy unsuitable for PCI and at high risk for CABG.³⁹⁷ 5010

A third study involved 208 consecutive patients scheduled for elective surgical treatment of major vascular disease who were at moderate to high cardiac risk for surgery. The patients were randomized to mandatory pre-operative coronary angiography and revascularization, if needed, or a selective strategy arm in which 5015 angiography was performed only if indicated based on the results

of non-invasive tests.³⁹⁵ The revascularization rates were 58% and 40% (P = 0.01), respectively. The in-hospital major adverse cardiovascular event rate did not differ between the two groups, but

5020 at a mean follow-up of 58 months patients subject to the systematic strategy of pre-operative coronary angiography had a statistically significant benefit in terms of freedom from major cardiovascular events as well as of survival.

LEAD patients scheduled for intermediate-risk surgery can be referred for surgery without additional testing for CAD. In patients 5025 scheduled for high-risk surgery, the number of cardiac risk factors should be assessed: angina pectoris, myocardial infarction, stroke or TIA, renal dysfunction (creatinine $>177 \mu mol/L$; 2 mg/dL), heart failure, and diabetes mellitus. In patients with three or more risk factors, additional cardiac testing for the presence and 5030 extent of CAD is recommended, if this will change management. In selected cases one might also consider additional cardiac testing as a means of patient counselling. If cardiac stress testing shows no or only mild stress-inducible myocardial ischaemia. additional invasive testing is not recommended. Again, all patients 5035 should be prescribed statins, low-dose titration of β -blockers before surgery, and aspirin; and those with systolic dysfunction

- should have ACE inhibitors. Patients with extensive stress-inducible myocardial ischaemia present a very difficult group. Optimal medical treatment including β -blockers and statins will not provide sufficient cardioprotection. However, preoperative prophylactic coronary revascularization is not generally associated with an improved perioperative outcome in this patient population. An individualized approach should be carried
- 5045 out for these patients, taking into account the very high cardiac risk of the planned surgical procedure and the possible harms of not performing surgery (i.e. risk of rupture in patients with abdominal aortic aneurysm). If it is decided to perform preoperative revascularization after multidisciplinary consultation, it

must be realized that the vascular surgical procedure should be postponed for \geq 14 days for balloon angioplasty, for 3 months for bare-metal coronary stent placement, and for 12 months for drug-eluting coronary stent placement.⁴⁷

In summary, perioperative cardiovascular complications are common in LEAD patients and result in significant morbidity following non-cardiac surgery. All patients require pre-operative screening to identify and minimize immediate and future risk, with a careful focus on known CAD or risks for CAD and functional capacity. The 2009 ESC guidelines⁴⁷ are clear that noninvasive and invasive testing should be limited to circumstances in which results will clearly affect patient management or in which testing would otherwise be indicated. β-Blockers, statins, and aspirin therapy should be continued in patients already on therapy and should be started in PAD patients undergoing

5065 intermediate- or high-risk surgery.

4.6.3.2.2.2 Patients with non-surgical lower extremity artery disease

Beyond the specific situation where a patient with LEAD will undergo vascular surgery, the goal of screening for CAD is to identify LEAD patients with a poor long-term prognosis in whom treatment and risk factor modification may improve their outcome. The co-existence of significant vascular lesions in different sites is a common feature of atherosclerosis, a systemic disease that can affect virtually any of the arterial vessels.^{384,402–404} The importance of prompt diagnosis and treatment of CAD has been repeatedly underscored. Half of patients with LEAD die from cardiovascular complications, and as early as 1 year after diagnosis; cardiovascular mortality rates are 3.7-fold higher than in patients without LEAD.⁴⁰⁵ One-third of PAD patients have significant CAD lesions. Of interest, asymptomatic CAD is usually independently associated with traditional risk factors but also with the severity and extent of non-surgical LEAD.

The pending question is whether such identification may improve clinical outcomes in patients who are already in secondary prevention programmes. Of importance, stable atherosclerotic 5085 patients without previous ischaemic events experienced significantly more events in the case of multisite artery disease,⁴⁰⁶ but this does not preclude any prognostic improvement in the case of prophylactic coronary revascularization. Screening asymptomatic CAD in patients with LEAD would be interesting if it leads 5090 to a different management from the one proposed for LEAD patients without CAD. Asymptomatic CAD in patients with LEAD is by definition stable, a situation in which coronary revascularization is controversial, given the negative results of the Clinical Outcomes Utilization Revascularization and Aggressive Drug 5095 Evaluation (COURAGE) trial,⁴⁰⁷ which failed to demonstrate the superiority of coronary revascularization over optimal medical therapy. However, this trial excluded situations in which revascularization was considered as necessary, especially patients with a poor ejection fraction and those with left main coronary artery ste- 5100 nosis >50%. These situations are not infrequent in patients with severe and extended LEAD, which is frequently associated with multisite artery disease. In the absence of any specific trial in LEAD patients, the screening and management of CAD may be considered after a multidisciplinary discussion for each case. 5105

5. Gaps in evidence

Solid evidence is still needed in many aspects of the management of PAD. In numerous situations, adequate trials are lacking and 5110 sometimes the management of PAD is extrapolated from data regarding CAD. In the field of interventional therapy, rapid changes in available therapeutic techniques create the situation in which clinical practice tends to follow technical developments without evidence from randomized trials. In addition, the random-5115 ized studies often yield conflicting results because of technical evolution and growth in the participants' experience. Moreover, PAD may involve multiple sites, creating a large number of clinical scenarios that are difficult to investigate in a systematic way. All of these aspects contribute to the broad spectrum of gaps in evi-5120 dence, of which the most relevant are listed below.

Carotid artery disease

- (i) The benefits of statins in patients with symptomatic carotid stenosis derive from the subgroup analysis of the SPARCL 5125 trial; the treatment goals for LDL cholesterol levels cannot be clearly defined. Even fewer data are available on the benefits of statins in asymptomatic carotid stenosis.
- (ii) The benefits of other preventive therapies, i.e. antiplatelet drugs and ACE inhibitors, are not well assessed in carotid 5130

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disease, especially in the case of carotid plaque with nonsignificant stenosis, which is the most frequent situation.

- (iii) The benefits of CEA in asymptomatic patients were proven in RCTs performed before the modern era of cardiovascular
 prevention, when medical therapy was almost non-existent and patients >80 years of age were excluded; thus, both CEA and CAS need to be evaluated against current optimal medical therapy in asymptomatic carotid stenosis, with a particular focus on elderly patients.
- 5140 (iv) The efficacy of EPDs during CAS has not been studied in adequately powered RCTs, and the available evidence is conflicting.
 - (v) The optimal duration of dual antiplatelet therapy after CAS is not well established.
- 5145 Vertebral artery disease
 - (i) Almost no data are available on the clinical benefit of revascularization of symptomatic VA stenosis, and on the comparison between surgical and endovascular revascularization.
- 5150 Upper extremity artery disease
 - (i) Almost no data are available on the clinical benefit of revascularization of symptomatic subclavian artery stenosis/occlusion, and on the comparison between surgical and endovascular revascularization.
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(ii) Little is known about the natural course in UEAD.

Mesenteric artery disease

- (i) No data are available on the comparison between surgical and endovascular revascularization for symptomatic mesenteric artery disease.
- (ii) No data are available on the potential benefits of revascularization for asymptomatic mesenteric artery disease involving two or more main visceral vessels.
- 5165 Renal artery disease
 - (i) Large-size trials are still necessary to clarify the potential benefits of RAS in patients with different clinical presentations of renal artery disease.
- 5170 (ii) Appropriate treatment of in-stent renal artery restenosis is not yet defined, although several trials are under way.

Lower extremity artery disease

 (i) The benefits of statins in LEAD patients derive mainly from small studies or from subgroup analyses of large RCTs focused on CAD patients; thus, the treatment goals for LDL cholesterol levels in LEAD patients cannot be defined clearly.

- (ii) Data on the benefits of the combination of 'supervised exercise training' and medical therapy are lacking.
- (iii) Data on the potential benefit of endovascular revascularization over supervised exercise for intermittent claudication are limited.
- (iv) The role of primary stenting vs. provisional stenting in aortoi- 5195 liac disease needs to be evaluated.
- (v) In the superficial femoral artery, the role of primary stenting in TASC II type C lesions, the potential benefit of covered stents for long superficial femoral artery occlusions, and the optimal treatment of in-stent restenosis need to be 5200 investigated.
- (vi) The role of drug-eluting stents and drug-eluting balloons in superficial femoral artery and below-the-knee interventions has to be established.
- (vii) Optimal treatment for popliteal artery stenosis needs to be 5205 addressed.
- (viii) The role of self-expanding stents for below-the-knee interventions is unclear.
- (ix) Benefits and/or adverse effects of $\beta\mbox{-blockers}$ in CLI must be further evaluated.
- (x) Optimal duration of dual antiplatelet therapy after LEAD stenting, as well as potential benefit of long-term dual antiaggregation therapy in patients with advanced CLI should be further investigated.
- (xi) The role of gene or stem cell therapy in CLI needs further 5215 studies.

Multisite disease

- (i) The need for prophylactic carotid revascularization in patients with asymptomatic carotid stenosis scheduled for CABG is still unclear.
- (ii) The preferred timing of CABG associated with carotid revascularization (synchronous or staged) is still unclear.
- (iii) If future studies confirm the benefits of carotid revascularization in patients undergoing CABG, the optimal treatment 5225 method (CAS vs. CEA) should be determined.

Acknowledgements

We thank Nathalie Cameron, Veronica Dean, Catherine Després, 5230 Jennifer Franke, Sanne Hoeks, Tomasz Jadczyk, Radoslaw Parma, Wojciech Wańha, and Piotr Wieczorek for their excellent technical assistance.



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The CME text 'ESC Guidelines on the diagnosis and treatment of peripheral artery diseases' is accredited by the European Board for Accreditation in Cardiology (EBAC). EBAC works according to the quality standards of the European Accreditation Council for Continuing Medical Education (EACCME), which is an institution of the European Union of Medical Specialists (UEMS). In compliance with EBAC/EACCME guidelines, all authors participating in this programme have disclosed potential conflicts of interest that might cause a bias in the article. The Organizing Committee is responsible for ensuring that all potential conflicts of interest relevant to the programme are declared to the participants prior to the CME activities.



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