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- 6 of the lower extremities
- 7 Draft

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9 This guideline replaces Note for Guidance on clinical investigation of medicinal products for the

treatment of peripheral arterial occlusive disease (CPMP/EWP/714/98 rev 1).

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	disease (LEAD), Chronic Limb Threatening Ischaemia (CLTI),
	Atherosclerosis

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Guideline on clinical investigation of medicinal products

for the treatment of peripheral arterial occlusive disease

of the lower extremities

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## **Executive summary**

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- 67 The main aim of the quideline is to address development of medicinal products for the treatment of
- atherosclerosis-related chronic ischaemia affecting the lower extremities.
- 69 This is the second revision of the NOTE FOR GUIDANCE ON CLINICAL INVESTIGATION OF MEDICINAL
- 70 PRODUCTS FOR THE TREATMENT OF PERIPHERAL ARTERIAL OCCLUSIVE DISEASE.
- 71 The main aim of the second revision was to explicitly address clinical development of medicinal
- 72 products intended to treat arterial lower extremity disease, to reflect updates in disease classification
- 73 and management, including the angiosome concept, but also provide guidance on estimands, definition
- 74 of clinical endpoints, and cover specific aspects related to the clinical development of advanced medical
- 75 therapies (ATMPs) in the setting of lower extremity arterial disease.

## 1. Introduction (background)

- 77 Lower extremity ischaemic disease (LEAD) is the most common clinical manifestation of chronic
- 78 peripheral arterial occlusive disease (PAOD) and is sustained by the obstruction of blood flow within the
- 79 limb arteries or the aortoiliac tract, commonly recognising atherosclerosis as etiopathogenetic cause.
- 80 The wide spectrum of clinical presentation, severity and anatomical distribution of the disease imposes
- 81 the use of classification schemes for patient management. Historically, the Fontaine and Rutherford
- 82 classifications have been most widely employed. The Fontaine categorization rates the LEAD in four
- 83 stages based on signs and symptoms: asymptomatic patients (stage I), intermittent claudication
- 84 (stage II, with a distinction between stage IIa and IIb referring to claudication at a distance > 200 m
- and < 200 m, respectively), rest pain (stage III), and trophic lesions including necrosis/gangrene
- 86 (stage IV). The Rutherford staging is based upon a combination of clinical symptoms and non-
- 87 invasive haemodynamic measures (i.e. treadmill test, arterial brachial indices). Like the Fontaine
- 88 scheme, it distinguishes four grades of disease including asymptomatic disease (Grade 0),
- 89 intermittent claudication (Grade I), rest pain (Grade II) and morphologic lesions (Grade III), with
  - a further subcategorization into six classes, based on objective criteria. With the introduction of the
- 91 new definition of the Chronic Limb Threatening Ischaemia (CLTI), which refers to the end-stage
- 92 manifestation of chronic atherosclerosis and better reflects the continuum of occlusive disease and
- 93 associated symptomatology, also including diabetic patients, the Global Vascular Guidelines (GVG)
- 94 recommend the application of the Wound, Ischemia, and foot Infection [WIfI] grading score. In
- 95 candidates for surgery, the GVG suggest the new Global Limb Anatomic Staging System (GLASS) as a
- 96 classification tool in the planning of revascularisation strategies: through an angiography-based
- 97 characterisation of the target arterial path (TAP) (that can also incorporate the so-called angiosome
- 98 concept) and the estimated limb-based patency (LBP), the GLASS rating reflects the probability of
- 99 success of the procedure and consequent clinical benefit of treatment in terms of limb salvage
- 100 probability.
- 101 The natural course of LEAD shows a high variability. In a recent trial (Hiatt WR et al, 2017) that mostly
- 102 recruited patients with Rutherford Grades I-III, the analysis of changes at 12 months from baseline in
- the Rutherford classification showed that the clinical symptomology remains unchanged in most

104	patients (63.7%), improves in 25.4% and worsens in 10.9% of the study population. A relevant
105	information for the purpose of preventative strategies came from the analysis of the group of
106	asymptomatic patients that reported disease progression with a rate of 26% over 1 year.
107	Almost 50% of patients who present with the most advanced stage of disease (i.e. CLTI) have no
108	history of prior LEAD. These patients are more likely older and male with pre-existing cardiovascular
109	morbidities and renal failure. Because of the neuropathy-associated symptomatology, diabetic patients
110	often remain underdiagnosed, and this explains the highest probability of presenting with de-novo
111	CLTI. Because of the high likelihood of peripheral arterial disease and the elevated prevalence of an
112	asymptomatic manifestation in this population (i.e. around 75%), learned societies now recommend
113	yearly screening in diabetic patients. Of note, the diabetic status predisposes to a more rapid
114	progression and increased severity of the disease. Data have been reported for a higher rate of major
115	and all amputations in type 1 compared to type 2 diabetics and differences in revascularization
116	strategies between the two groups (Jain N et al, 2022).
117	Generally, all LEAD patients (even if asymptomatic) are at increased risk of major adverse
118	cardiovascular events such as myocardial infarction, stroke or cardiovascular death and major adverse
119	limb events like amputations, chronic or acute lower limb ischaemia, lower limb revascularization.
120	Subjects not suitable for, or who failed revascularisation using surgical bypass or endovascular
121	methods, who often represent the target population of clinical trials testing ATMPs, are at high risk for
122	amputation and death (Norgren et al 2007).
123	The therapy of LEAD focuses on symptoms relief and the prevention of cardiovascular morbidity,
124	amputation and death. Given the association with conventional cardiovascular risk factors,
125	interventions aiming at controlling smoking, hypertension, dyslipidaemia and diabetes as well as the
126	use of antithrombotic agents, pain controllers and rehabilitation programs as appropriate, all concur to
127	the optimal management of patients undergoing either a medical treatment or surgical approach to
128	therapy.
129	2. Scope
130	Guidance is provided on the clinical development program of medicinal products intended to treat
131	lower extremity arterial disease. Acute ischaemia and peripheral vascular disorders of inflammatory or
132	immunologic origin such as Buerger's disease and necrotic vasculitis are not considered because these
133	diseases differ from arteriosclerosis obliterans in their clinical picture, in their evolution and in their
134	prognosis.
135	The current revision concerns the clinical development program of medicinal products, including
136	ATMPs, intended for an indication in LEAD, with specific reference to the definition of the study

population, choice of clinical endpoints for inference of efficacy, and estimation of treatment effect

(estimands) in confirmatory trials.

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## 3. Legal basis and relevant guidelines

- 140 This Guideline should be read in conjunction with the introduction and general principles of Annex I to
- Directive 2001/83/EC, as amended, and all other relevant EU and ICH guidelines. These include, but
- 142 are not limited to:

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- Guideline for good clinical practice (EMA/CHMP/ICH/135/1995 [ICH E6[R2]]);
- ICH Guideline E8 (R1) on general considerations for clinical studies (EMA/CHMP/ICH/544570/1998 Corr\*);
- Pharmacokinetic studies in man (1987);
- Note for Guidance on Population Exposure: the extent of population exposure to assess clinical safety (CPMP/ICH/375/95 [ICH E1]);
- Note for Guidance on Dose Response Information to Support Drug Registration (CPMP/ICH/378/95
   [ICH E4]);
- Note for Guidance on Statistical Principles for Clinical Trials (CPMP/ICH/363/96 [ICH E9]) and
  Addendum on estimands and sensitivity analysis in clinical trials to the guideline on statistical
  principles for clinical trials (EMA/CHMP/ICH/436221/2017 [ICH E9[R1]]);
- Points to consider on switching between superiority and non-inferiority (CPMP/EWP/482/99);
- Guideline on the choice of the non-inferiority margin (EMEA/CPMP/EWP/2158/99); Note for Guidance on choice of control group in clinical trials (CPMP/ICH/364/96);
- Note for Guidance on Studies in Support of Special Populations: Geriatrics CPMP/ICH/379/95
   (ICH E7) and Questions and Answers (EMA/CHMP/ICH/604661/2009 [ICH E7 Q&A]);
- Reflection Paper on assessment of cardiovascular safety profile of medicinal products (EMA/CHMP/50549/2015);
- Guideline on the evaluation of medicinal products for cardiovascular disease prevention (EMEA/CHMP/EWP/311890/2007);
- Guideline on human cell-based medicinal products (EMEA/CHMP/410869/2006);
- Reflection paper on stem cell-based medicinal products (EMA/CAT/571134/2009);
- Quality, non-clinical and clinical aspects of medicinal products containing genetically modified cells
   (CHMP/GTWP/671639/2008);
- Guideline on safety and efficacy follow-up and risk management of advanced therapy medicinal products (EMEA/149995/2008).

# 4. Development strategy

- 170 Throughout the clinical development programme, key aspects to be considered pertain to the selection
- of patients and definition of study objectives, which are expected i) to depend on the development
- phase of the medicinal product (see section 5 for a detailed guidance on study design) and ii) to

173 provide adequate representativeness of the target indication. This latter point is an important

requirement in confirmatory therapeutic studies, for which a disease stage-specific approach can be

adopted in the choice of the study population and clinical endpoints, thus making efficacy results fully

176 evaluable. In this context, a distinction between treatment and preventive objectives can be

177 considered, based on the intended indication. A list of acceptable clinical endpoints is reported in this

section, while the choice of valid primary and secondary endpoints according with both the

development and disease stage is discussed in section 6-7.

## 4.1. Selection of patients - General considerations

181 The criteria used for the diagnosis of LEAD in patients recruited for clinical trials must be clearly

defined. The diagnosis, type of occlusive lesion (stenosis, complete block) and its location must be

confirmed by objective means. Different classification systems are currently available for patient

staging, covering both the clinical and anatomical characterization of disease. The choice of the

different classification systems and their combination is expected to be adequate to the definition of a

study population that should be homogenous in terms of disease severity, localization, and risk of

disease progression. While a symptomatology-based grading supports the definition of stage of

disease, anatomical classification systems also incorporating, but not limited to the angiosome

concept, are essential elements to be considered in revascularization and wound healing treatments.

For instance, the angiosome-based approach implies the angiographic study of the affected territories

to establish the feasibility for either a direct flow restoration by re-establishing arterial patency in line

to the ischemic area, or the possibility to achieve an indirect revascularisation through collaterals. The

success of both techniques is influenced not only by technical aspects related to the surgical

intervention, but also to comorbidities (i.e. diabetes, smoking status) that influence the perfusion

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196 Since revascularisation should be offered to all CLTI patients, the definition of "no or poor

revascularisation options" needs to be tightly defined in the study protocol. Treatment groups should

be balanced in respect of patient demography, severity of disease, previous revascularisation

199 procedures and duration of symptoms, as well as concomitant medications and standardized

rehabilitation programs, as appropriate. This is usually ensured by adequate randomisation

procedures. Influences of potential confounders such as cardiovascular risk factors (e.g. smoking,

hypertension, hyperlipidaemia, diabetes mellitus) should be carefully taken into consideration in the

analysis plan, and respective therapeutic measures should not be changed during the course of the

trial unless ethically or clinically indicated (e.g. for participant safety) keeping in mind that an

appropriate strategy dealing with intercurrent events should be specified in the study protocol (see

section 5). A distinction between type 1 and type 2 diabetes should also be considered, based on the

reported differences in terms of clinical outcomes and therapeutic management between the two

208 populations.

## 4.2. Assessment of efficacy/ Methods to assess efficacy

## 4.2.1. Treatment endpoints

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11	4.2.1.1.	Improvement of	i walking	capacity
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- 212 Claudication distances should be assessed using a standardised, reproducible test methodology (i.e.
- 213 treadmill test or 6MWDT test). There are two internationally accepted treadmill protocols, i.e. the
- constant workload protocol using a constant speed and grade (mostly 3.2 km/h and 12% grade), and
- the graded test where the speed is kept constant, but the grade is varied, starting horizontally but
- then increasing in predefined steps (e.g. 2%) at predefined intervals (e.g. 2 min). The two tests differ,
- in that the relationship between workload and walking time follows a linear function with the constant
- 218 test but a curvilinear function with the graded test.
- 219 Both tests can be equally recommended for use in clinical trials but cannot be used in an
- 220 interchangeable way; a decision on the treadmill protocol and the treadmill settings must be made
- beforehand and should not be altered.
- The 6MWD test can also be considered as a method to assess clinical efficacy. It has been
- demonstrated to be representative of daily life walking functionality and is correlated with daily
- physical activity in contrast to the treadmill test (McDermott 2014).

#### 225 Treadmill testing

#### 226 Initial claudication distance (ICD)

- 227 From a clinical point of view, ICD compared to absolute claudication distance (ACD) as symptomatic
- 228 endpoint may be the more important variable, since patients seldomly force themselves to the extreme
- of ACD. On the other hand, ICD is more subjective.
- 230 If ICD is chosen as primary endpoint, ACD should be evaluated as a secondary endpoint.

#### 231 Absolute claudication distance (ACD)

- ACD can be used alternatively. However, if a graded treadmill protocol is used, ACD should be the
- primary efficacy variable. The reproducibility of ACD is superior to ICD with graded protocols.
- 234 If ACD is chosen as primary endpoint, ICD should be evaluated as a secondary endpoint.

### 235 Six Minute Walk Distance Test (6MWT)

- The 6MWT can be used as a primary endpoint, considering it has been demonstrated to be
- 237 representative enough of daily life walking functionality, and is even more correlated with daily
- 238 physical activity than the historically more often used treadmill testing (McDermott 2014). The 6MWT
- was found also to correlate with mortality outcomes (McDermott 2011).
- 240 The minimal clinically important difference (MCID) that is intended to be used in the inference of
- 241 efficacy of treatment requires to be pre-specified in the study protocol and is expected to be justified
- and relevant to the specific targeted population.

#### 4.2.1.2. Improvement of pain

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- Pain (if existing) should be quantified. The intensity of the pain should be assessed by means of
- 246 standardised methods (e. g. visual analogue scale). Pain relief should be defined as "complete relief of
- pain". Since pain at rest is an endpoint influenced by variables such as mood, motivation, and other
- factors, the standardisation of the trial methodology is of utmost importance. This does not only refer
- to the methodology used to quantify pain but does include factors such as the time of pain assessment
- 250 (same time of the day, preferably at drug trough levels), the personnel taking the measurement
- 251 (which should not change), and the assessment of analgesic consumption. The consumption and the
- type of analgesics should be measured and documented, although the comparison of patients on
- 253 different analgesics schemes may be difficult.

#### 254 **4.2.1.3.** Healing of ulcers

- 255 Ulcer healing must be defined as healing of all ischaemic ulcers in both legs (all ulcers epithelialized as
- assessed by an independent physician and documented by photography). Since quantification of partial
- 257 healing may be difficult to assess objectively and since the clinical relevance of partial healing remains
- unclear, only total healing of lesions should be reported as main efficacy criterion.

#### 259 **4.2.1.4.** Interventional/surgical procedures

- 260 Clinical parameters that should be considered include the rate of revascularisation procedures, minor
- amputations as well as frequency of major amputations.

#### 262 **4.2.1.5. Quality of Life (QoL) outcomes**

- 263 In trials with adequate sample size an assessment of quality of life may be performed by using
- 264 properly validated general and disease specific questionnaires. Measuring QoL adds a subjective
- perspective to the evaluation of treatment and complements the clinical evaluation (Kinlay, 2013).

## 266 **4.2.1.6. Response Based Endpoints**

- Response could be defined as the patient being alive, having both legs, having no wound or pain, and
- being off analgesics. This endpoint concept would consider the time period for which the response can
- be maintained, and is intended to study the overall medium-term/long-term outcome.
- 270 Clinical response can also be defined by a change in stage as defined by classification scoring systems.

#### 4.2.2. Prevention endpoints

#### 272 **4.2.2.1. Prevention of disease progression**

#### 273 **Amputation**

- The rate of major amputations can be considered as a primary endpoint or as a component of a
- 275 composite primary endpoint.
- Only major amputations, above the ankle, should be counted (unlike minor amputations, i.e below
- the ankle). Both legs must be considered for the assessment of amputation rates.

278	The criteria for major amputations are to be specified à priori in the study protocol to avoid relevant
279	centre-related effects (e.g. a more conservative or a more progressive attitude towards the indication
280	for amputation).
281	4.2.2.2. Prevention of CV/ischemic events
282	Mortality
283 284 285	All-cause mortality or cardiovascular mortality alone has rarely been used as a primary efficacy endpoint in prevention trials. However, either all-cause or cardiovascular death has always to be considered as an important component of a composite efficacy endpoint.
286	Cardiovascular morbidity
287 288	Cardiovascular morbidity alone should not serve as a single primary endpoint, but it should be incorporated into a composite primary endpoint which also includes mortality.
289	Composite Endpoints
290 291 292 293 294	The most adequate endpoint in prevention studies is a composite endpoint, if specified à priori, and if consisting of clinically relevant components. Such an endpoint may include cardiovascular morbidity (e.g. stroke, myocardial infarction) and all-cause mortality. Major amputation as a component of a composite endpoint can also be considered, and should be generally adopted in more severe stages (stages III/IV, CLTI).
295	5. Study design
296	5.1. Pharmacokinetics and Pharmacodynamics
297 298 299	For the purpose of investigating the pharmacokinetics and pharmacodynamics of a new investigational medicinal product intended for the treatment of peripheral arterial disease, reference is made to the available and general EMA guidance on the different aspects of clinical pharmacology (see section 3).
300 301 302 303	Pharmacodynamic endpoints should be product-specific, defined based on the mechanism of action of the investigational medicinal product with the intention to provide a "proof-of-concept" and evidence of the pharmacological activity of the drug, as well as a characterisation of the exposure-response relationships.
304	5.2. Exploratory therapeutic studies/dose finding
305 306	The purpose of this development phase is to prove the therapeutic activity of the drug under investigation and to establish suitable therapeutic dose ranges.
307 308 309	The dose and therapeutic schedule should be selected according to the results of previous studies.  These studies should be carried out in selected patients with strict inclusion and exclusion criteria (see section 4 for general considerations on patient selection).

310	A randomised, double-blind, parallel group, placebo-controlled design is recommended. Primary
311	assessment criteria depend on the aim of the study (e.g. walking distance in claudication trials, relief
312	of rest pain and ulcer healing as symptomatic endpoints in critical limb ischaemia).
313	In general, the treatment period should be in the range of 2 to 3 months. However, the overall
314	duration of dose response studies may vary and should be properly justified considering the
315	mechanism of action and the main endpoint of the study. A run-in period is recommended to verify the
316	stability of the patient's conditions, e.g. comedication, stability of the claudication distances.
317	5.3. Confirmatory therapeutic studies
318	5.3.1. Statistical aspects
319	Studies aiming at the proof of efficacy must have a confirmatory statistic approach – e.g. a
320	demonstration of superiority, equivalence or non-inferiority must be pre-specified in the protocol.
321	The design and analysis should be performed in accordance with the available methodology guidelines.
322	Generally, studies which investigate the possibility to reduce the risk of several serious events
323	(prevention studies) and which therefore may use a composite endpoint as a primary variable should
324	be designed to either show a significant difference between the treatment groups or be designed as
325	non-inferiority studies with suitable comparators.
326	However, efficacy in the composite endpoint should be coupled with evidence that none of the
327	components is negatively influenced.
328	It is recommended to plan the study in such a way that the results indicate a clinically relevant effect
329	in addition to its statistical significance.
330	The duration of the trial depends on the aim of the study and the endpoint(s) chosen.
331	However, the length of exposure to the drug should be sufficient to investigate the potential of
332	tolerance developing.
333	5.3.2. Confounding factors
334	There are several confounding factors which could influence the results of therapeutic clinicaltrials in
335	LEAD.
336	Regular physical exercise improves symptoms of intermittent claudication. Thus, the frequent use of
337	repeated exercise testing in clinical trials may lead to an improvement in exercise capacity independent
338	of drug treatment. This should be considered in the design and analysis of such trials. At the same
339	time, adherence of patients to standardized supervised rehabilitation programs should be taken into
340	account. To this respect, exclusion criteria may limit recruitment to those with training capacity based
341	on respiratory or cardiovascular conditions, or major gait disturbance.

Regular physical exercise and cessation of smoking are of much importance in the treatment of

intermittent claudication and have significant impact also on the outcomes of revascularisation.

344 Advice on smoking cessation and physical exercise should be given before patients are included in a 345 clinical trial. Respective effects should be documented. 346 Even if claudication distances were considered clinically stable and stability was proven during the run-347 in phase of a clinical trial in LEAD stage II patients, a marked placebo effect cannot be avoided. 348 Experiences from previous trials indicate that observed variability of claudication distances between 349 trials varies considerably. In addition, the distribution of these endpoints is often skewed. These 350 factors should be considered when calculating the sample size and planningthe analysis strategy 351 (e.g. considering logarithmic transformation). 352 In patients hospitalised for critical limb ischaemia there is a high response rate as regards bothrest 353 pain and ulcer healing during placebo treatment. If this situation is not accounted for, the number of 354 patients enrolled to a clinical trial may be inadequate for inference of treatment effect relative to 355 placebo. **5.3.3.** How to estimate effect (Estimands) 356 357 An estimand is a precise description of the treatment effect reflecting the clinical question posed by a 358 given clinical trial objective. A disease-specific approach should be adopted for estimation of treatment 359 effect, with the definition of the main clinical endpoints to be driven by the intended use of an 360 investigational drug and target population, as defined by disease stage and anatomical localization. As 361 a general consideration for LEAD patients, the primary outcome should be either the symptomatic 362 relief or a preventative effect on cardiovascular events and amputations. Differences between the 363 active treatment and control arms would provide inference of treatment effect. Variables such as the 364 actual adherence to treatment or intercurrent events should be considered in the estimation of the 365 effect through appropriate sensitivity analyses. 366 Intercurrent events expected to be potential modifiers of treatment effect in the context of LEAD 367 include drug discontinuation, changes in background therapies with effects on the perfusion status or 368 drug-to-drug interactions, as well as terminal events (i.e. death or leg amputation when not included 369 in the clinical study endpoints) or unplanned revascularisation procedures. The nature of the specific 370 intercurrent events and their chance to occur vary depending on the target population, as defined by 371 disease severity, anatomical distribution, and presence of comorbidities. It is expected that the study 372 protocol identifies and clearly defines relevant strategies to handle pre-specified intercurrent events. 373 Moreover, protocol violations and deviations should be considered. 374 As a general consideration, a treatment policy strategy should be regarded as the preferred approach

for providing an unbiased estimate of treatment effect in the target population. An appropriate analysis

plan should be conceived to handle intercurrent events through specifications of censoring rules,

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imputation of missing values and sensitivity analyses.

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#### 6. Studies evaluating treatment outcomes (e.g. walking 378 distance, pain, ulcer healing) 379 380 For studies evaluating treatment of symptoms of LEAD, the specific claim of the clinical benefit put 381 forward in the product information is expected to be clearly supported by relevant study population 382 and primary endpoints. 6.1. Design elements 383 384 A randomised parallel group, double-blind, placebo-controlled design is generally required. Placebo 385 should be used for the control group since suitable reference substances have not yet been established 386 in the symptomatic treatment of intermittent claudication, as well as in more advanced stages of 387 disease. Active drug-controlled trials without a placebo arm may only be considered if the comparator 388 drug has consistently shown superiority over placebo (assay sensitivity, see ICH E10). 389 A run-in phase of 2-6 weeks is recommended to verify the stability of the patient's conditions, e.g. 390 comedication, stability of the claudication distances. 391 The active treatment phase should last for a minimum of 6 months for administration of an 392 investigational medicinal product intended for chronic therapy. Shorter lengths are expected to be 393 adequately justified depending on the nature of the product, mode of action, intended indication. 394 Depending on the duration of active treatment, the length of the follow-up period may vary. Generally, 395 the cumulative duration of active treatment phase and follow-up period should not be less than 6 396 months for a controlled trial. Double-blindness should be maintained during the whole period. 397 The allowed medication during the active treatment phase and follow-up period should be standardised 398 as much as possible, and should be comparable for the treatment groups. 399 The follow-up period should be specified à priori in the study protocol. 400 A disease stage-specific approach should be adopted. Across the spectrum of CLTI patients, the 401 unequivocal characterization of disease stage in the study population should be provided to estimate 402 treatment effect (i.e. revascularisation success). 403 6.2. Patient selection/target population

- The stage of disease should be clearly outlined in the study protocol using appropriate classification scoring systems and depending on the clinical outcomes and intended therapeutic indication for the experimental medicinal product to be tested.
- For claudicant patients, a history of typical intermittent claudication lasting for at least 6 months to ensure clinical stability is expected. The clinical diagnosis of LEAD should be confirmed by objective evidence (e. g. reduced ankle systolic blood pressure). It is recommended that patients with high variability in the walking distance be excluded. For this purpose, at least two treadmill tests should be performed with a timeinterval of ≥ 1 week. The maximum change in the claudication distance should not exceed a predefined threshold [e.g. 25 % for the absolute claudication distance (ACD)]. Walking training is considered the first treatment option in this patient population. Therefore,

414 it is expected that this therapeutic measure is tried for all patients before they are considered for 415 entering the trial, unless\_otherwise justified. If walking training is applied during the study, some 416 advantages may exist to use supervised, structured protocols. 417 Claudication studies should not include patients suffering from illnesses limiting their exercise capacity 418 (angina pectoris, heart failure, respiratory disease, orthopaedic disease, neurological disorders ...). 419 Only patients with rest pain due to chronic critical limb ischaemia, that is persistent recurrent pain at 420 rest requiring analgesics for more than 2 weeks should be acceptable. 421 Generally, patients eligible for surgical/interventional reconstruction should not be included. However, 422 patients with a high perioperative/periinterventional risk for ischaemic complications may be included, 423 provided that the study design guarantees that necessary invasive procedures are notdelayed. 424 As per all patients with CLTI, the diagnosis currently requires objectively documented atherosclerosis, 425 based on a combined evaluation of pressure measurements and Doppler arterial waveforms 426 corroborated by the vascular imaging, combined with rest pain for at least 2 weeks. 427 In diabetic patients, macroangiopathy (rather than microangiopathy or neuropathy) should be the 428 leading cause for the lesion(s). Patients with skin lesions of mixed arterio-venous origin or patients 429 suffering from a vasculitis should not be included. 430 It is strongly recommended to study diabetic patients and non-diabetic patients in separate trials or to 431 use appropriate stratification. 432 CLTI patients who present with "poor or no revascularisation option", and this justifies the placebo use 433 in the control arm. Since revascularisation should be offered to all CLTI patients and be prioritized with 434 respect to experimental treatments, conditions leading to "poor or no revascularisation option" needs 435 to be tightly defined in the study protocol. To this end, it should be considered that treatment decisions 436 also reflect the local expertise of clinical specialists, patient's access to specific surgical techniques and 437 technologies, as well as the availability of a multidisciplinary and interdisciplinary team-based care to 438 optimize patient's outcomes at a given facility. In this context, a centralized independent adjudication 439 committee that would revise individual clinical cases may be considered under certain circumstances to 440 ensure consistency in the recruitment process and fully adherence to the inclusion criteria across study 441 centres.

## 6.3. Choice of endpoints

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It is recommended that a disease stage-specific approach is adopted for the choice of the clinical endpoints to be used in confirmatory studies evaluating symptomatic outcomes, also depending on the intended drug indication. The chosen endpoints should be clinically meaningful and consistent with the expected drug effect according to its mechanism of action. For specific methodological aspects regarding these clinical endpoints, reference is made to the sections 5.

## 448 6.3.1. Primary Endpoints

- Acceptable primary treatment efficacy goals are those providing inference of treatment with regards to
- improvement in walking capacity, control of pain, and wound healing.
- 451 For claudication studies, walking distance should be the primary symptomatic endpoint.
- 452 For patients with rest pain, the main symptomatic efficacy endpoint is the relief of pain at rest. It must
- be shown that the drug under investigation has no analgesic properties.
- 454 In the presence of skin lesions, the main symptomatic efficacy endpoint is complete healing of all
- 455 necrosis and ulcerations.
- Wound healing, pain control and limb salvage should be the primary goal for ATMPs in patients with
- 457 CLTI who have no options for revascularization.
- 458 A response-based approach can be applied assuming that evidence of maintenance of effect in the
- 459 overall medium term/long-term can be provided. This endpoint concept would consider the time period
- 460 for which the response can be observed.

### 461 **6.3.2. Secondary Endpoints**

- 462 Secondary endpoints should focus on clinically relevant data supporting the study aim. These include
- 463 walking distance, haemodynamic measures, interventional/surgical procedures, quality of life,
- 464 consumption of analgesics.

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# 7. Studies evaluating prevention of disease

## progression/prevention of ischemic events

### 7.1. Design elements

- 468 A randomised, parallel group, double-blind, controlled study design is generally required. Placebo
- and/or active drug-controlled trials may be adequate.
- 470 Treatment should last for a minimum of 12 months, but longer periods are recommended. For specific
- 471 considerations, reference is made to the Guideline on the evaluation of medicinal products for
- 472 cardiovascular disease prevention (EMEA/CHMP/EWP/311890/2007).

## 7.2. Patient selection/target population

- In general, all patients with a proven diagnosis of LEAD are eligible for clinical trials of the prevention
- of ischaemic events. Thus, patients may present with a history of intermittent claudication, previous
- 476 peripheral (lower extremity) vascular intervention such as surgical endarterectomy, bypass grafting or
- 477 abdominal aortic aneurysm repair, transcutaneous endoluminal procedures (PTA, stenting), minor or
- 478 major amputations because of LEAD, or may be asymptomatic if LEAD has been proven by objective
- means (e.g. haemodynamic and non-invasive imaging studies or angiography).
- 480 Regarding the cardiovascular prognosis, it is recommended that patients to be included in a clinical
- 481 trial should be homogeneous. Patients with intermittent claudication and CLTI should be studied

distinction should serve as stratification factor.
Diabetics and non-diabetics should be studied separately or, if included in the same study, this
population to be recruited.
anatomical disease classifications should be adopted for an unequivocal description of the study
interventions or treatment in patients not suitable for revascularization), additional clinical and
depending on the aim of the investigational treatment (i.e. post-procedural or peri-procedural
separately, or appropriate stratification techniques should be applied; within the CLTI group and

#### Background therapy

- 490 Vasoactive substances other than the test drug, haemodilution or rheological therapy may be 491 considered prohibited medications as per protocol and any use be recorded during the study. Other 492 pharmacotherapy which is considered relevant for the treatment of LEAD or relevant for the prevention 493 of cardiovascular events in general must be documented. It should be maintained during the course of 494 the study. If this is not possible, e.g. during follow-up or deterioration, the study design must consider 495 this appropriately (for specific considerations. reference is made to the statistical section on 496 estimands). All other medicinal products can be given, as long as they have no established effect on
- 497 the investigated parameters. However, their administration must be fully documented.
- 498 In CLTI, basic local treatment (e. q. local wound treatment, removal of necrotic tissue, antibiotics)
- 499 must be documented and should be standardised as much as possible.
- 500 Because LEAD patients have a high risk for cardiovascular events, it is recommended to use
- 501 antiplatelet agents and statins as background therapy as well as to optimise diabetic control and
- 502 implement smoking cessation in patients prior to study entry. It is desirable to avoid any change in
- 503 medication.

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### 7.3. Choice of endpoints

### 7.3.1. Primary endpoints

- 506 Since the goal of preventative trials is the reduction of atherosclerosis-associated morbidity and 507 mortality events, cardiovascular morbidity (e.g. myocardial infarction, stroke), major amputation and 508 death are the clinically most meaningful endpoints. They may be used in isolation or in form of a 509 composite, combined endpoint. The components of a composite endpoint will depend on the clinical 510 stage of LEAD. Whereas cardiovascular morbidity/mortality and all-cause mortality will be the most 511 appropriate components for trials in mild to moderately severe diseased patients (i.e. Fontaine stages I 512 and II), the rate of major amputations should also be considered in trials including CLTI patients (i.e.
- 513 Fontaine stages III and IV).
- 514 The question on whether to use all-cause mortality or cardiovascular mortality as a component of a 515 composite endpoint will depend on the estimated frequency and the possibility to identify 516 cardiovascular death. The trial hypothesis, whether this is superiority on non-inferiority, may also play 517 a role, particularly if the incidence of cardiovascular- and non-cardiovascular death does differ 518 substantially. However, generally, all-cause mortality should be given preference as long as there are 519 no persuading arguments for the use of cardiovascular mortality. If a composite endpoint is used and

520 521	significantly influenced by the drug under investigation, it is expected that the single components will move in the same direction. However, a significant effect in the composite endpoint should be coupled
522	with evidence that none of the components is negatively influenced.
523	7.3.2. Secondary Endpoints
524	Components of the composite endpoint
525 526	If the primary endpoint is a composite endpoint, the components of this composite endpoint should be evaluated as secondary endpoints.
527	8. Safety aspects
528 529 530	All adverse effects occurring during clinical trials should be fully documented. Any groups especially atrisk should be identified. Special efforts should be made to assess potential adverse effects that are characteristic of the class of drug being investigated.
531 532	Adverse drug events occurring during the treatment should be carefully recorded throughout all study phases, including data about their nature, frequency, intensity, and relevance.
533	Particular attention should be paid to the following specific side effects:
534	8.1. Blood pressure and heart rate
535 536	This may be either symptomatic or asymptomatic. Special attention should be paid to orthostasis and first-dose phenomenon.
537 538	8.2. Neurohumoral activation and pro-arrhythmic and/or pro-anginal effects
539 540 541	Depending on the particular pharmacodynamic properties of the new agent, measurement of effects on neurohumoral compensatory mechanisms, heart rate, ECG and Holter monitoring should be performed at frequent intervals throughout the study.
542	Effects on cardiac conduction (PR, QRS, QT and QTc) should be documented.
543	8.3. Rebound, withdrawal phenomena
544 545	Withdrawal phenomena, especially rebound phenomena should be studied in selected cases depending on the mode of action of the investigational product and specific concerns.
546	8.4. Mortality, cardiovascular morbidity
547 548 549 550	If not investigated as efficacy endpoint, a separate analysis on all-cause mortality, cardiovascular morbidity / and vascular death should be made on basis of the pivotal clinical trials. A new agent in LEAD is only acceptable for registration if there is no negative impact on mortality and cardiovascular morbidity.

## 9. Studies in special populations

## 9.1. Studies in elderly patients

- The prevalence of LEAD increases with age in both sexes, with the prevailing group of patients being in
- 554 the elderly population. It is relevant to generate evidence on clinical pharmacology, efficacy and safety
- that are representative of this subgroup and the different old age categories, considering that it will
- 556 constitute the predominant target population.

## 9.2. Studies in paediatric patients

- 558 Atherosclerosis-related peripheral vascular disease is very rare in paediatric patients but can affect
- 559 children with specific underlying predisposing conditions leading to an early disease manifestation also
- within the paediatric age (i.e. nephrotic syndrome, type 1 diabetes, familial hypercholesterolaemia)
- (Lavie G et al. 2014; Virani SS et al. 2020; Akinyosoye G et al. 2022 and 2023). The epidemiological
- scenario in this setting is still under definition and might change over time. Extrapolating data from the
- adults to the paediatric setting is a possibility that requires considerations on a case-by-case basis.

## 10. Advanced Therapy Medicinal Products (ATMPs)

- 565 ATMPs comprise gene therapy, somatic cell therapy medicinal products and tissue engineered
- products. In the setting of peripheral arterial disease, regenerative treatment aims at promoting limb
- perfusion and skin lesion healing through restoration of the vascular function (i.e. vasodilation) and
- 568 structure, as well as phenomena of vascular regeneration mediated by angiogenesis, arteriogenesis,
- and vasculogenesis. The regulation of the inflammatory milieu and tissue regeneration are additional
- 570 mechanisms of action potentially contributing to the regenerative function of ATMPs in the CLTI
- 571 condition.

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- 572 While the general recommendations on study design already provided in other sections of this
- 573 guidance all apply to ATMP products tested in this clinical condition, peculiarities can be recognised
- 574 pertaining to the study design, patient selection and clinical endpoints that warrant specific
- 575 considerations. For the general principles of quality and clinical experimentation of ATMPs, reference is
- 576 made to the relevant EMA guidelines that should be consulted in conjunction with the present
- 577 recommendations.

### 10.1. Study Design

- Whilst it is acknowledged that dose-dependency generally does not apply to ATMPs, the chosen
- therapeutic dose should be fully substantiated.
- 581 Certain ATMPs can undergo a single administration; however, the follow-up period should not be less
- than 6 months in controlled trials. The length of follow-up may vary but should be appropriate for the
- assessment of functionality and structural aspects of the repaired and/or regenerated tissue (target
- limb arteries), as well as its persistence in the human body which is expected to be considerably longer
- than 6 months. The follow-up should also enable assessment of secondary efficacy endpoints or

important safety variables, such as the frequency of reconstructive measures, morbidity and mortality
and the rate of major amputations. Double-blindness should be maintained during the whole period.
10.2. Patient selection
Across the spectrum of CLTI patients, the unequivocal characterization of disease stage in the
study population should be provided to estimate treatment effect (i.e. revascularisation success).
The anatomical disease characterization, also adopting the angiosome terminology, should be
considered in the study design to ensure a standardised route of drug administration, and may
support the assessment of the regenerative properties of tested ATMPs, even though it is worth
noting that for previously re-vascularised patients the angiosome concept might not be directly
applicable given the post-surgery anatomical rearrangement.
In addition to the diabetic status, the smoking habit could impact on the regenerative effect of
ATMPs. It is strongly recommended to study diabetic and non-diabetic as well as smokers and
non-smokers in separate trials or to use appropriate stratification schemes.
10.3. Criteria for Efficacy
10.3.1. Primary Endpoints
Wound healing, pain control and limb salvage should be the primary goal for ATMPs in patients
with CLTI who have no options for revascularization. For specific methodological aspects
regarding these clinical endpoints, reference is made to the relevant sections 4-7. For prognostic
endpoints including amputation reference is made to the section on prevention trials.
10.3.2. Secondary Endpoints
In addition to the secondary endpoints listed in the section 6 and 7, demonstration of to the
vascular regenerative properties and the mode-of-action of ATMPs appears particularly relevant
to ATMPs trials. To this end, vascular imaging modalities are regarded as particularly valuable in
the confirmation of clinical efficacy. It is acknowledged that objective imaging-based
methodologies are not available for quantification of vascular changes. However, it is expected
that all relevant measures will be taken to support the claimed mechanistic model.
10.4. Criteria for Safety
ATMPs carry additional risks that should be addressed according to current relevant EMA
recommendations (Guideline on safety and efficacy follow-up and risk management of advanced
therapy medicinal products [EMEA/149995/2008 rev.1]). Moreover, adverse events of special interest
should be identified and pre-specified in the study design, including but not limited to
allergic/immunologic reactions, severe infections, adverse events of CRP increases, specific AEs of

MACE, MI/UA, strokes, AEs of rest pain increase and/or ulcer worsening, incidence of tumours. An

appropriate risk management plan is always required to monitor events in the post-marketing phase,

considering the limited length of follow-up in registration clinical trials.

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## 660 12. Definitions

- 661 ACD: Absolute Claudication Distance;
- 662 ATMP: Advanced Therapy Medicinal Products;
- 663 CRP: C-reactive Protein;
- 664 CLTI: Chronic Limb Threatening Ischaemia;
- 665 ECG: Electrocardiography;
- 666 GVG: Global Vascular Guidelines;
- 667 GLASS: Global Limb Anatomic Staging System;
- 668 ICD: Initial Claudication Distance;
- 669 LBP: Limb-Based Patency;
- 670 LEAD: Lower Extremity Ischaemic Disease;
- 671 MCID: Minimal Clinically Important Difference;
- 672 6MWDT: 6-Minute Walk Distance Test;
- 673 PAOD: Peripheral Arterial Occlusive Disease;
- 674 PAD: Peripheral Arterial Disease;
- 675 PTA: Percutaneous Transluminal Angioplasty;
- 676 QoL: Quality of Life;
- 677 TAP: Target Arterial Path;
- 678 WIfI: Wound, Ischemia, and foot Infection (grading score).