Peripheral arterial disease (PAD), presenting as intermittent claudication (IC), affects a large proportion of the general population [1, 2] and is one of the major causes of disability in middle age and elderly [3, 4]. Despite its relevance, PAD treatment still remains a challenge. This is essentially due to the fact that, for a long time, the pathophysiology of this condition remained unknown, and the narrowing of a proximal artery was considered the only factor responsible for the reduced oxygenation in the skeletal muscle. Conversely, the restriction of blood flow in a proximal artery is only the primum movens of a sequence of microcirculatory alterations accounting for the mismatch between metabolic supply and demand in the affected skeletal muscle.

In PAD, the ischemia induced by maximal exercise increases oxidative stress, inflammation and endothelial dysfunction. Perturbation of the endothelial homeostasis results in increased adhesiveness of leukocytes and platelets, and in reduced vasodilator capability. These events, expression of the interplay between inflammation and endothelium, provoke an obstacle in the microcirculation with a reduction in the nutritive blood flow, leading to acidosis and impaired energy metabolism in skeletal muscle, with consequent reduced exercise tolerance.

ET counteracts these effects by improving walking ability and quality of life in patients with intermittent claudication, thus representing the gold standard in the treatment of PAD.

Keywords: peripheral arterial disease, intermittent claudication, microcirculation, skeletal muscle energy metabolism, exercise training.

lial dysfunction [10-15]. In particular, in affected individuals exercise induces increase in plasma levels of thiobarbituric acid-reactive substances [16], accompanied by an increase in thromboxane and interleukin-8 [10, 11, 17], and elevated plasma levels of intercellular adhesion molecules-1 (sICAM-1), vascular cell adhesion molecules (sVCAM-1), von-Willebrand factor (vWF), E-selectin, endothelin-1 (ET-1), thrombomodulin, and circulating endothelial progenitor cells [18-25]. Furthermore, a maximal working load induces depressed artery brachial flow-mediated endothelial-dependent dilatation (FMD, an index of endothelial dysfunction).

In addition to the above described systemic inflammatory response, and probably even more important with respect to the impairment of microvascular circulation in the affected limb, acute exercise incites a local inflammatory response. In 17 patients with one-sided PAD, Neumann et al. [26], found that immediately after claudication, total neutrophil number, neutrophil flexibility and the proportion of activated neutrophil were increased in the venous draining from the affected leg versus the arterial blood. These venous-arterial differences were not observed in the circulation of the exercising contralateral healthy leg. The aforementioned alterations, typically inflammatory in nature, may act so that a certain number of neutrophils remains entrapped in the microcirculation of the affected limb causing vessel plugging. Furthermore activated leukocytes release thromboxane A2, which in addition to be a vasoconstrictor, promotes platelet aggregation [27], so inducing the phenomenon of platelet plugging. To this phenomenon in claudicants may also participate overexpression of P-selectin [28, 29] which mediates platelet-endothelium interaction [30]. Consistent with the observation by Neumann et al. [26], our group observed that in the claudicant limb, but not in the healthy legs of the control group, maximal exercise increased the transmural venous-arterial difference of the neutrophil myeloperoxidase (MPOx) content [31], an index of neutrophil activation and a well established marker of inflammation [32]. MPOx is an enzyme which, when released by activated neutrophils, exerts very harmful effects on the endothelium [32]. In particular, it uses nitric oxide (NO) as a physiologic substrate, thereby reducing the bioavailability of this gas which is fundamental for the endothelial homeostasis. In addition to MPOx, activated neutrophils release various noxious substances, among which elastase, that has been shown to have harmful effects for the endothelium in vitro [33], and show a progressive increase from normal, asymptomatic PAD patients, to claudicants [34]. Furthermore, in claudicants, its levels have been shown to further increase with exercise [35]. Disturbed endothelium results in a reduced vasodilator capability and in the already mentioned increased adhesiveness to leukocytes and platelets which causes a physical obstruction to microcirculation. An additional mechanism that may favour the adherence of neutrophil to endothelium is impaired shear stress, which also involves an inflammatory reaction. Indeed, the CD11b/CD18-ICAM-1 adhesion mechanism is very sensitive to shear stress as low as 0.5 dyne/sec [36]. Obviously, distal to an obstruction there is considerable turbulence so that there is a strong likelihood of a low shear stress zone distally to the obstruction. Finally, endothelial cell swelling, favoured by acidosis [37], may cause a further obstacle to microcirculation [38, 39]. All these events, expression of the interaction between inflammation and endothelium, by reducing the nutritive blood flow, leads to acidosis and impaired energy metabolism with consequent reduced exercise tolerance (Fig. 1).

3. Oxygen delivery to the tissue

One of the most important functions of the blood circulation is oxygen delivery to the tissues. The classical scheme of tissue oxygenation assumes that oxygen passes to the tissue via the capillaries. However, blood entering capillaries is only 50% saturated [40], thus implying that half of O2 gathered by the lungs exits the circulation prior to arriving to the capillaries include oxygen shunting from arterioles to parallel venules, arteriolar-capillary oxygen diffusional shunting and periarterial tissue consumption.

3.1. Arterioles and tissue oxygenation

The first demonstration that capillaries are not the sole source of oxygen for the tissues dates back to 1970 when Duling and Berne found that significant amounts of oxygen exit the arteriolar network [40]. More recently, the phosphorescence decay technology [41] provides means for evaluating simultaneously intra- and extravascular pO2, and therefore it is a direct method of evaluating both longitudinal vessel pO2 gradients (or blood oxygen saturation gradients) and the perivascular pO2 tissue gradients that presumably determine the rate of oxygen exit from blood vessels. Results confirm that a major portion of blood oxygen is delivered to the tissue by the arterioles. Furthermore, this technique allows an accurate mass balance between the decrease of oxygen content in the arterioles and the diffusion flux of oxygen out the microvessels are determined by the oxygen gradient measured in the surrounding tissue. These measurements support the hypothesis that the exit of oxygen from arterioles is due to a high rate if oxygen utilization by the vascular wall.

3.2. The arteriolar wall as an oxygen sink

When endothelium is removed from the dog hind limb, oxygen consumption decreases by 34% [42]. Indeed, endothelium is capable of metabolic activity which can be 100-fold that of other cells [43]. It is the site of chemical synthesis and metabolic processes that require oxygen. These include synthesis and secretion of renin, prostaglandins, collagen, NO, endothelin, prostacyclin, interleukin, degradation of bradykinin and prostaglandins, clearance of proteins, lipids and lipoproteins, expression of adhesion molecules. Moreover, endothelial cells have an active actin/myosin-based contractile system that also may consume O2 [44]. Therefore, it is conceivable that during the metabolic stress induced by ischemia more oxygen is consumed by the endothelium and less becomes available to the metabolic requirements of the muscle. This pathophysi-
logic mechanism could contribute to the alteration of oxidative machinery occurring in the skeletal muscle during claudication.

4. Skeletal muscle metabolism

Mitochondrial Adenosine TriPhosphate (ATP) regeneration is the principal mechanism in the skeletal muscle for meeting energy requirements for sustained force production. It depends on the activity of tricarboxylic acid (TCA) cycle that is fuelled by acetyl-CoA formed primarily from oxidation of pyruvate and fatty acids. The rate of these metabolic pathways is coupled to the rate of muscle work, and conversely, muscle work is coupled to the supply of oxygen and the rate of oxidative phosphorylation. At low work loads (defined as those below the lactate threshold), muscle metabolism is primarily aerobic, with fatty acids as the primary substrate [45]. Conversely, at high work loads (above the lactate threshold), carbohydrates serve as the primary substrate for exercising muscle [46], and thus acetyl-CoA is formed from pyruvate through the action of pyruvate dehydrogenase (PDH). However, with high intensity exercise, the rate of acetyl-CoA formation exceeds the maximal capacity of the TCA. As a consequence, acetyl-CoA tends to accumulate within the mitochondria, increasing the acetyl-CoA/CoA ratio and thus inhibiting PDH. The reduced oxidative utilization of glucose leads to increased lactate formation. Under these conditions the role of carnitine is crucial. It serves as a “buffer” of the metabolically critical mitochondrial acetyl-CoA pool [47]. Through the action of carnitine acetyl transferase, carnitine depletes acetyl-CoA and releases free CoA and acetyl-carnitine, which, unlike acetyl-CoA, may be transported out of the mitochondria and released in the blood stream (Fig. 2). Indeed, increased levels of acetyl carnitine occur in muscle and plasma of normal subjects performing maximal exercise [32].

4.1. Effect of ischemia

Muscle ischemia adversely affects muscle metabolism. In the ischemic muscle during exercise the levels of inorganic phosphate are very high as the breakdown of ATP and phosphocreatine is elevated. Therefore, the ratio of inorganic phosphates to phosphocreatine increases to a greater extent than in normoperfused muscle. The result is a higher rate of glycolytic metabolism as indicated by the marked drop in pH (Fig. 3) These changes observed by 31P magnetic resonance spectroscopy [7] are in accordance with biopsy studies, which demonstrated increased accumulation of lactate and depletion of cellular ATP during exercise in claudicants compared to normal subject [8]. Reduced ATP synthesis results from the reduced vascular perfusion which decreases both oxygen delivery (thus impairing mitochondrial oxidation) and the washout of toxic metabolites from the muscle (hence inhibiting many enzymatic activities and cellular proton efflux). In particular, patients with claudication accumulate acetylcarnitine (thus reflecting acetyl-CoA accumulation) and lactate (as a consequence of PDH inhibition). Importantly, there is an inverse correlation between both plasma and muscle levels of acetylcarni-
EXERCISE TRAINING AND INTERMITTENT CLAUDICATION

Figure 2. Schematic depiction of skeletal muscle substrate metabolism. Note that accumulation (↑) of acetyl-CoA within the mitochondria inhibits (↓) the activity of PDH. Carnitine removes the acetyl-CoA excess by forming acetylcarnitine. PDH= pyruvate dehydrogenase; CPT= carnitine palmitoyl transferase; CAT= carnitine acetyl transferase.

Figure 3. 31P MRS data in exercising muscle of normal subjects and patients with peripheral ischemia (note that in the normoperfused muscle, in the early phase of the exercise there is a slight increase in the oxidative ATP synthesis). In the ischemic muscle during exercise the levels of inorganic phosphate are very high as the breakdown of ATP and phosphocreatine is elevated. Therefore, the ratio of inorganic phosphates to phosphocreatine increases to a greater extent than in normoperfused muscle. The result is a higher rate of glycolytic metabolism as indicated by the marked drop in pH. ATP= adenosine triphosphate; Pi= inorganic phosphate; PCr= phosphocreatine; ADP= adenosine diphosphate; VO2= oxygen consumption.
the decrease observed before treatment [50]. This implies that a corresponding amount of acetyl-CoA was removed from mitochondria with consequent stimulation of PDH. Indeed, in patients with IC, carnitine administration results in a reduction of the lactate concentration in the venous blood leaving the exercising ischemic muscle [51].

5. Benefits of exercise training (ET) in PAD

In 1957, Foley [52] described the beneficial effects of daily walks and physical training in patients with IC. Several subsequent reports confirmed these findings, and in particular, a meta-analysis from the Cochrane Collaboration [53], concluded that exercise improves maximal working time by an average of 150%, an extent that would appear to exceed that obtained with current pharmacological therapies.

Although the mechanisms by which ET yields improvements in walking ability remain speculative, many changes that occur during supervised physical activity presumably simulate an adaptative response that ultimately reduces claudication symptoms [54]. One of the mechanisms may be the increase in blood flow consequent to the process of collateralization. This is traditionally viewed as a recruitment of pre-existing collaterals (arteriogenesis) or an expansion of microvessels (angiogenesis) into ischemic areas. Following a program of ET, humans with heart failure will increase expression of vascular endothelial growth factor (VEGF) messenger RNA in skeletal muscle [55], and animal models of limb ischemia reveal an increase in collateral development [56, 57]. However, in humans, studies demonstrating collateral development or increase in calf blood flow in response to ET are limited and conflicting [58-63]. In PAD, Gardner et al. [61] showed that six months of exercise increased maximal calf blood flow by 30%, and this change was associated with improved walking ability (r=0.38, P<0.05). On the other hand, a number of other studies did not document increased leg blood flow in patients with IC, although all of them reported an improvement in exercise performance [58-61]. Hence, other mechanisms must account for the large exercise-induced improvements in function and symptoms that occur in patients with claudication.

While acute exercise incites a systemic and local inflammatory response, ET reduces free radical damage, neutrophil activation, plasma levels of plasma markers such as C-Reactive Protein (CRP), Serum Amyloid A (SAA), CD11b [64], and increases endothelial NO synthase (eNOS) expression and phosphorylation [65]. The consequence of these changes is an increased activity of NO, the synthesis of which is reduced in claudicant patients [66]. In animal models of peripheral ischemia, ET improves NO activity [66], and thus it is presumably that it counteracts the microcirculatory alteration described above. Unfortunately, studies of ET on vasodilator endothelial function in patients with claudication are limited. An uncontrolled study [67] found that training induced improvements in walking ability was associated with an increase in FMD. Furthermore, in 22 PAD patients Andreozzi et al. [17] measured baseline and post-exercise FMD, before and after ET. Before training the pre-exercise FMD was 7.6±2.9 and post-exercise 5.3±3.3 (−33.2%) (P<0.01). After training the pre-exercise FMD was 10.3±4.04 and post-exercise 7.8±2.6 (−18.97%) (P<0.01). Thus, ET increased both baseline and post-exercise FMD. Furthermore, it halved the difference in FMD between the pre- and post-exercise observed before and after training, although this was not statistical significant.
Finally, ET induces multiple beneficial effects in the structure and metabolism of skeletal muscle. It increases the percentage of oxidative type 1 fibers and decreases the density of type 2B glycolytic fibers. Furthermore, it increases the volume density of mitochondria [68]. In a phosphorus-31 nuclear magnetic resonance spectroscopy study, after physical training, the phosphocreatine depletion and the increase in adenosine diphosphate (ADP) during exercise were reduced significantly although there were no significant changes in the response of muscle pH to exercise. The substantial correction of the impaired oxidative metabolism of skeletal muscle was confirmed by the finding that in untrained subjects the initial rate of phosphocreatine resynthesis after exercise (a measure of the rate of oxidative ATP synthesis and the inferred maximal rate of mitochondrial ATP synthesis) were reduced compared with rates in control subjects (P<0.003) and both were significantly increased (P<0.05) by training [69].

To the best of our knowledge, only one study investigated the effects of ET on skeletal muscle histology and metabolism in PAD. Hiatt et al. [70] performed gastrocnemius muscle biopsies at rest and before and after ET. At 12 weeks training did not alter type 1 or type 2 fiber areas and did not increase citrate synthase activity. Improvements in exercise performance with training were associated with a correlative decrease in plasma (r=-0.067) and muscle (r=-0.59) short chain acylcarnitine concentration. As reported above, such a decrease of resting plasma levels of short-chain acylcarnitine implies a greater availability of carnitine with consequent amelioration of energy metabolism [70].

**Conclusions**

Although the pathophysiology of IC is primarily accounted for a flow-limiting stenosis of a conduct artery, a large body of evidence indicates that impairments in microcirculation and energy metabolism play a relevant role in the reduced working ability of affected individuals. In microvessels, disturbed endothelium results in increased adhesiveness of leucocytes and platelets which causes physical obstruction. The consequent reduction in nutritive blood flow leads to acidosis and impaired energy metabolism with secondary decrease in exercise tolerance. By counteracting these effects, ET is considered by the American Heart Association the treatment of choice for improving walking ability and quality of life in IC patients [71].

**Riassunto**

Sebbene nell’arteriopatia obliterante degli arti inferiori (AOAI) la causa principale dell’inaugnata apporto di sangue all’arto affetto durante l’esercizio fisico (EF) sia una stenosi in un condotto arterioso, molte evidenze indicano che alterazioni del microcircolo e del metabolismo energetico del muscolo scheletrico giocano un ruolo rilevante nella ridotta autonomia funzionale dei pazienti con claudicatio intermittens (CI).

Lo scopo di questa rassegna è stato quello di far luce su questo argomento, ponendo maggiore attenzione sui benefici derivanti dall’EF nel trattamento dei pazienti arteriopatici.


L’EF, neutralizzando questi effetti, migliora l’autonomia di marcia e la qualità della vita nei pazienti con CI, rappresentando così il gold standard nel trattamento dell’AOAI.

Parole chiave: arteriopatia obliterante degli arti inferiori, claudicatio intermittens, microcircolo, metabolismo energetico del muscolo scheletrico, esercizio fisico.

**ABBREVIATION LIST**

PAD: Peripheral Arterial Disease  
IC: Intermittent Claudication  
FMD: Flow-Mediated Dilation  
MPOx: Myeloperoxidase  
NO: Nitric Oxide  
ATP: Adenosine TriPhosphate  
TCA: TriCarboxylic Acid  
PDH: Pyruvate dehydrogenase  
ET: Exercise Training

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