Buerger’s disease: providing integrated care

Peter Klein-Weigel¹
Theresa Sophie Volz¹
Leonora Zange²
Jutta Richter³

¹Clinic of Angiology, ²Clinic of Cardiology and Nephrology, HELIOS Klinikum Berlin-Buch, Berlin,
³Medical Faculty, Department of Rheumatology and Hiller Research Unit Rheumatology, Heinrich-Heine-University Duesseldorf, Duesseldorf, Germany

Abstract: Buerger’s disease, also known as thromboangiitis obliterans (TAO), is a segmental inflammatory disease affecting small- and medium-sized vessels, which is strongly associated with tobacco use. Although the etiology is still unknown, recent studies suggest an immunopathogenesis. Diagnosis is based on clinical and angiomorphologic criteria, including age, history of smoking, clinical presentation with distal extremity ischemia, and the absence of other risk factors for atherosclerosis, autoimmune disease, hypercoagulable states, or embolic disease. Until now, no causative therapy exists for TAO. The most important therapeutic intervention is smoking cessation and intravenous prostanoid infusions (iloprost). Furthermore, effective analgesia is crucial for the treatment of ischemic and neuropathic pain and might be expanded by spinal cord stimulation. Revascularization procedures do not play a major role in the treatment of TAO due to the distal localization of arterial occlusion. More recently, immunoadsorption has been introduced eliminating vasoconstrictive G-protein-coupled receptor and other autoantibodies. Cell-based therapies and treatment with bosentan were also advocated. Finally, a consequent prevention and treatment of wounds and infections are essential for the prevention of amputations. To achieve better clinical results, integrated care in multidisciplinary and trans-sectoral teams with emphasis on smoking cessation, pain control, wound management, and social care by professionals, social workers, and family members is necessary.

Keywords: Winiwarter-Buerger’s disease, Winiwarter–Buerger, thromboangiitis obliterans, immunoadsorption

Introduction

In 1879, Winiwarter,¹ a young assistant physician of Theodor Billroth in Vienna, published the clinical course and pathologic examination of a lower limb amputation of a 57-year-old male describing “a peculiar kind of angiitis and endophlebitis with gangrene”. Although this is considered to be the first case report of thromboangiitis obliterans (TAO), the disease is currently more exclusively linked to the American surgeon Buerger², whose systematic work on clinical and pathological aspects of the disease constituted our modern understanding of the disease.

TAO is an inflammatory vascular pathology affecting small- and medium-sized arteries and veins leading to vessel occlusions by the formation of a mononuclear cell-rich thrombus.² Its etiology is still unknown, but it is inseparably linked to tobacco use. Due to an undulating clinical course, normal vessel segments and different stages of lesions (acute to chronic types) might be found together in the same patient.²
Patients with Buerger’s disease usually present with acute ischemic or infectious acral lesions (ulcers, gangrenes, subungual infections, phlegmonous) and/or thrombophlebitic nodules. Skin discolorations such as Raynaud’s phenomenon, acrocyanosis, or livedo-like pictures are often seen.3–5 Rarely, a nonerosive arthritis might precede ischemia for months or years.6

Epidemiology

Buerger’s disease occurs worldwide and is more prevalent in males, but an increasing prevalence in females has been reported in different countries.7–9 Disease characteristics and prognosis do not differ between males and females.9 In contrast to North America and Western Europe, the Mediterranean, the near and far East, and the Indian subcontinent are high prevalence regions.3–5 Thus, prevalence rates among in-hospital treated patients with peripheral arterial occlusive disease were reported to range from 0.5% to 5.6% in Western Europe, 45%–63% in India, and 16%–66% in Korea and Japan.10 In the meanwhile, the formerly often cited extremely high prevalence rate in Ashkenazi Jews was identified as a scientific error as it referred to the response rate of an invitation to participate in a study and did not reflect the true prevalence in this ethnic group.11 Reported prevalence of TAO seems to decline during the past decades due to a decrease in tobacco use or – as others believe – due to an increase in socioeconomic conditions.12–14

Etiologic, pathologic, and pathogenetic aspects

There is a very tight correlation between the manifestation, flaring, and recurrence of Buerger’s disease (no tobacco, no Buerger’s disease).3–5,10 Thus, tobacco must be considered to be the dominant risk factor. Besides potential differences in regional smoking habits, regional and ethnic differences in the prevalence of the disease might point toward a genetic background determining individual susceptibility. Human–leukocyte–antigen-linked factors may play a role; nevertheless, human leukocyte antigen association studies revealed heterogeneous findings.15–18 Published genetic polymorphisms consist of CD14 T7T polymorphism, eNOS gene 894 T/T polymorphism as a protective factor, and MyD88 rs7744 A-G polymorphism, coding for a Toll-like receptor signaling adaptor.19–22

Chronic infectious disease – especially periodontal disease – was found to be associated with TAO.23,24 On the other hand, in a particular disease group of the disease (ie, low social status and excessive smokers), periodontal disease can be expected to be very high triggering a close correlation, which does not necessarily imply a causative linkage. Nevertheless, smoldering infections such as periodontitis might trigger autoimmune mechanisms and coagulation.24

Signs of endothelial activation and proliferation as well as the presence of immunocompetent cells are seen in acute type lesions. Immunoglobulin and complement deposition as well as CD4+ and CD8+ T-lymphocytes, CD 20+ B-lymphocytes, and S-100-positive dendritic cells were found alongside the lamina elastica interna, which becomes structurally altered but is typically preserved.25–30 Giant cell formation and the appearance of the so-called microabscesses within the mononuclear cell-rich thrombus may occur.2

Analysis of cytokine activation in patients with TAO revealed a pattern of elevated pro- and anti-inflammatory cytokines.31,32 Various kinds of autoantibodies have been identified in patients with TAO, including anti-endothelial antibodies, antibodies directed against vessel wall structures such as elastin and collagen, anticardiolipin antibodies, and antineutrophil cytoplasmatic antibodies.33–39 More recently, agonistic autoantibodies directed against G-protein-coupled receptors were identified as potentially promoting vasospasm, compromising microcirculation, damaging vessel structures, and inducing proliferative processes.40

Overall, the findings are consistent with the assumption of an immunopathogenesis of TAO. A first model of this new paradigm has recently been published by Ketha and Cooper.41

Social and psychosomatic aspects

Buerger’s disease typically occurs in patients with a low social status.14 Some authors even described a Buerger-type personality with manipulative and autoaggressive tendencies often matched with denial, negligence, or tendencies to minimize their illness, while others even presumed typical morphological characteristics.42,43 However, no systematic work has been performed in this field, and the preliminary results do not allow differentiating between premorbid traits and conditions and psychological consequences of the chronicity and severity of the disease or implications of chronic drug intake such as morphine or opioids in the affected patients.

Diagnostic criteria

Diagnosis is usually based on clinical and angiomorphic criteria published by Olin et al and Shionoya.15,44 The latter is based on only five criteria and thus easy to remember (Table 1). Combined upper and lower extremity involvement is present in ~20%–25% of the cases.5,46 An isolated affection of only one limb strongly argues against Buerger’s
Coronary, and visceral arteries have been published. Nevertheless, case reports of typical lesions even in cerebral, coronary, and visceral arteries have been published. Thrombophlebitis – if present – is often migratory type and precedes or parallels arterial disease activity.

Establishing diagnosis

Typically, a young heavy smoker presents with a more or less symmetrical distal ischemic syndrome or a crural–acral or antebrachial–acral type of arterial occlusion in two or more extremities. Distal pulses are usually absent or diminished, but can be normal in the case of exclusive acral disease manifestations. Allen’s test often reveals an upper extremity disease. Proximal arterial involvement is rarely present. Nevertheless, case reports of typical lesions even in cerebral, coronary, and visceral arteries have been published. Thrombophlebitis – if present – is often migratory type and precedes or parallels arterial disease activity.

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Ankle brachial index or forearm–brachial index is usually reduced, but might be normal in cases with more distally located disease. Digital pulse recordings are characterized by low amplitudes and delayed slopes, anarchic or silent pulse curves.

Angiographically, a typical but not pathognomonic pattern (Figure 1) has been described, which substantiates the diagnosis. This pattern might also be identified by magnetic resonance imaging or a careful duplex ultrasound examination.

Embolic disease – if suspected – can be ruled out by transesophageal echocardiography, computer tomography angiography, and duplex ultrasound of the proximal extremity arteries.

Capillary video microscopy is often limited by infection or hornification. It often reveals capillary loss and unspecific morphologic changes. Nevertheless, capillary microscopy is a useful tool in the differential diagnosis.

There are no specific biomarkers for TAO. Systemic inflammatory markers such as C-reactive protein are usually absent or only slightly elevated and are therefore unsuitable for the assessment or monitoring of disease activity. Nevertheless, laboratory tests are important to exclude other entities such as diabetes, connective tissue disease, vasculitides, or congenital or acquired thrombophilia.

If biopsy can be performed without endangering the limb or if an amputation is performed, diagnosis should be confirmed by histopathologic examination. Fresh thrombophlebitic nodules are suitable for this purpose.

Therapy

In the past decades, therapeutic efforts concentrated on pain and infection control, revascularization, or amputation.

Smoking cessation

Nevertheless, the most important therapeutic intervention in Buerger’s disease is smoking cessation. Its overwhelming effect for the prevention of consecutive limb amputation was impressively shown. Patients with TAO should be prevented not only from active smoking but also from alternative consumption mode and passive exposure. Tobacco dependency is usually considered to be exceptionally strong in patients with Buerger’s disease, but in the only prospective study addressing this question the degree of tobacco dependence was similar to that in patients with coronary artery disease.

Individual strategies for smoking withdrawal have to be discussed, including in- and outpatient treatment in specialized institutions with multidisciplinary teams. Unfortunately, the percentage of patients who maintain smoking cessation despite is low. In one study, the continuous abstinence rate decreased from 29% at the end of the treatment to 18.5% at the 12-month follow-up. Best results seem to be achieved by structured and guided peer group and anti-smoking programs starting during hospital stay or shortly thereafter. Medication support by nicotine replacement therapy, bupropion, or varenicline might be provided. Whether replacement therapies or adjuvant therapies might prolong disease activity was – to our knowledge – not addressed, but should be examined in further studies.

As was shown by the TEMPO study, work and family circumstances, co-occurring substance use, and psychological difficulties may influence smoking cessation in the typical age groups of patients with Buerger’s disease. Factors specifically associated with a low probability of smoking cessation were job strain and symptoms of hyperactivity/inattention, while occupational grade was associated with smoking relapse. Prostanoid therapy and antiplatelet drugs

The effectiveness of prostanoid therapy was elucidated in two older randomized trials and in two more recently published trials prospectively assessing clinical outcome in addition to smoking cessation, aspirin, or compared with sympathectomy. Iloprost, a prostacyclin analog, is considered the drug of choice. Fiessinger and Schäfer randomly allocated

## Table 1: Diagnostic criteria for TAO

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<th>Criteria</th>
<th>Description</th>
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<td>Disease onset before the age of 50 years</td>
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<td>Smoking</td>
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<tr>
<td>Absence of other atherosclerotic risk factors</td>
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<tr>
<td>Infrapopliteal arterial occlusions</td>
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<tr>
<td>Upper limb involvement or phlebitis migrans</td>
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Note: Data from Shionoya

Abbreviation: TAO, thromboangiitis obliterans.

## Prostanoid therapy and antiplatelet drugs

The effectiveness of prostanoid therapy was elucidated in two older randomized trials and in two more recently published trials prospectively assessing clinical outcome in addition to smoking cessation, aspirin, or compared with sympathectomy. Iloprost, a prostacyclin analog, is considered the drug of choice. Fiessinger and Schäfer randomly allocated
152 patients with Buerger’s disease and pain from critical leg ischemia to iloprost intravenously or low-dose aspirin for 28 days in a double-blind trial. Fifty-eight (85%) of 68 iloprost-treated patients showed ulcer healing or relief of ischemic pain versus eleven (17%) of 65 in the aspirin-treated group. Forty-three (63%) patients on iloprost treatment had complete relief of pain, compared with 18 (28%) on aspirin. Unfortunately, these striking results could not reproduced in the largest randomized trial, including 319 patients, when iloprost was administered orally in two dose regimes and compared with placebo. In a study published by the Turkish Buerger’s Disease Research Group, complete ulcer healing rate was 61.9% in those receiving iloprost and 41% in the lumbar sympathectomy group at 4 weeks and 85.3% versus 52.3% at 24 weeks. In a prospective Turkish multicenter observational trial in 158 patients with Buerger’s disease suffering from rest pain and/or ischemic lesions, complete ulcer healing without residual rest pain or major amputation was met by 60% of the patients treated with iloprost. Pain scale values decreased significantly after 4 weeks and 24 weeks. Ulcer size reduction at 4 weeks and 24 weeks, as well as clinical status, investigator, and observer grading, was also significantly improved at both time points.

A recently published Cochrane review emphasized low-quality evidence concerning medical therapy in Buerger’s disease and stated that high-quality trials assessing the effectiveness of pharmacological agents in people with Buerger’s disease are urgently needed. Although widely used, there is no proven evidence for platelet function inhibitors such as aspirin or clopidogrel in TAO. Same is true for oral anticoagulants.

Analgesia
Effective analgesia is crucial as ischemic and neuropathic pain in Buerger’s disease is usually severe. Therefore, co-treatment by pain therapy specialists is essential. Combinations of morphine or opioids and peripheral analgetics are often required in high doses. Antidepressants might be of additional value. Epidural anesthesia, neuronal block, or local analgesia is often applied. In the selected cases, spinal cord stimulation might improve not only pain control but also perfusion by inducing sympathicolysis and via antidrome mechanisms.

Revascularization procedures
Due to the distal localization of arterial occlusions and the absence of recipient vessels, interventional or surgical
revascularization is impossible to perform in the majority of cases. Nevertheless, especially in the older literature, series of peripheral bypass procedures in Buerger’s disease have been published with acceptable results in highly selected patients (revascularization rate: 4.6%–17.7%) and highly specialized centers, reporting up to 48.8% and 62.5% at 5 years, and 43.0% and 56.3% at 10 years, respectively.75–78

Endovascular therapy might also be effective even in extended femorotibial occlusions, but the reported numbers are small and the role of endovascular therapy in Buerger’s disease has yet to be defined.79–82

**Sympathectomy**

As revascularization procedures are often impossible, surgical or chemical sympathectomy is often considered, despite a lack of valid data supporting this practice. In a more recent publication of a cohort of 216 Turkish patients, sympathectomy was preferred over open surgical reconstruction or bypass procedures (81% versus 19%). Clinical outcome following sympatholysis was rated “improved” in 52.3%, “stable” in 27.8%, and “worse” in 19.8% of the patients, while seven major and 36 minor amputations were performed.83 On the other hand, lumbar sympathectomy was reported to be inferior to intravenous iloprost applications in a randomized trial by the same group.68 Thus, currently, there is no proven indication for primary sympathectomy in Buerger’s disease despite the fact that it is still widely used in many countries.

**Immunosuppressive drugs**

Although widely used in former times, there is no proven evidence for the use of steroids or cyclophosphamide therapy.43,84,85

**Progenitor cell therapy**

In the past decade, cell-based therapies with autologous progenitor cells harvested from bone marrow or peripheral blood have been advocated in critical limb ischemia, including Buerger’s disease. The cell suspensions are usually applied by intramuscular injections alongside the vascular bed of the limbs or by intra-arterial injection.86–90

Meta-analyses confirmed practicability and safety as well as positive therapeutic effects (including pain control, ulcer healing, pain-free walking ability, amputations-free survival) of cell-derived therapies in critical limb ischemia.88,91,92 Patients with Buerger’s disease responded better than patients with atherosclerotic peripheral arterial disease in some, but not all studies.93–95 There seems to be a significant time lag of 4–8 weeks until an improvement of microcirculation becomes evident in responders after bone marrow cell transplantation.96 This lag might be especially problematic in case of severe ischemia demanding a more urgent improvement of perfusion. Results of randomized double-blinded studies are awaited, but the hype about progenitor cell therapy already seems to be over.

Intramuscular or intra-arterial progenitor cell therapies compete against surgical concepts of stimulating angiogenesis and arterialization in patients with TAO based on tibia bone distraction or fenestration, or implantation of a Kirschner wire in the tibial intramedullary canal. These procedures were also reported to result in improved outcomes including pain scores, ulcer healing, and walking distances.97–99 Nevertheless, they might be hampered by side effects as the operation takes place in an ischemic environment. Controlled and comparative studies are missing.

**Immunoadsorption**

Immunoadsorption (IA) is an extracorporeal procedure clearing plasma from immunoglobulins and circulating immune complexes approved in many immune-mediated diseases. Based on the hypothesis that Buerger’s disease is immune-mediated with humoral factors playing a major role, IA was successfully introduced in a pilot study conducted by Baumann et al100 and later introduced in clinical routine care.101 More recently, a possible effective mechanism of IA was elucidated as IA eliminates vasoconstrictive α- and endothelin receptor agonistic autoantibodies that seem to cluster in patients with Buerger’s disease.102 The pilot study revealed a fast improvement of pain, a steep increase in tcpO2-levels and decrease in tcCO2-levels, an improvement in ulcer healing, and a high return-to-work rate of the patients.100 Overall, these positive results were reproduced in a clinical routine setting.101 IA is being performed on five consecutive days for 5–6 hours per session aiming for a clearance of ~2.5-fold of patient’s plasma volume.100–102 It might be followed by the substitution of polyvalent immunoglobulin to ameliorate infectious risks in patients with active gangrene or ulcers.100

**Bosentan**

Referring to a first positive case report, another Spanish group published their results of a pilot study introducing the endothelin-receptor-blocking agent, bosentan, in the treatment of digital ulcers in patients with Buerger’s disease.103,104 Dosing was derived from the approved application for prophylaxis of digital ulcers in patients with scleroderma. Despite the promising results, ~1/6 of the patients had to undergo minor digital amputations: a finding, not necessarily arguing against the effectiveness of bosentan as minor amputations might have already been inevitable at presentation or might even have been made successfully possible by the treatment, and a finding that was also observed in our IA patients.101
Wound management and infection

Local wound management in ischemic lesions in Buerger’s disease is based on modern wound care standards with surgical debridement and selected wound dressings according to the wound’s stage and condition. As ischemic wounds – if at all – tend to heal very slowly, a cross-sectional and multidisciplinary concept is crucial. Wound, soft tissue, and bone infections might cause serious clinical problems and relapses as they occur in often highly ischemic states. Bacterial species and resistance spectra vary widely with gram-positive species dominating our own series (unpublished data). Starting calculated antibiotic therapy, one has to take anaerobic species and multiple resistances into account.

Outcome and social consequences

According to an older literature survey conducted by Börner and Heidrich,105 amputations were performed in 6.9%–75% of patients with TAO within 3–10 years of follow-up. Minor amputations predominated; nevertheless, major amputation rate was reported as high as 31%. The high amputation rates in the relatively young patients significantly contribute to the financial and social burden of the disease, which additionally includes job loss, early retirements, divorces, and subsequent social isolation.105

Perspective

Many decades from Buerger’s landmark report the disease he dedicated himself to remains an important health issue not only in high prevalence regions as it affects young people and induces a high social and financial burden. Hopefully, the new paradigm of an immunopathogenesis of Buerger’s disease might improve knowledge and prognosis in the future. To achieve better clinical results, integrated care in multidisciplinary and trans-sectoral teams with emphasis on lifestyle changes such as smoking cessation, pain control, wound management, and social care by professionals, social workers, and family members is necessary.106,107

Disclosure

The authors report no conflicts of interest in this work.

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