Lymphedema: a mini-review on the pathophysiology, diagnosis and treatment

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Abstract

Lymphedema (LE) is a complex chronic debilitating condition resulting from inadequate lymph flow leading to significant physical and psychological morbidity. Recent studies of lymphedema diagnosis and therapy are promising. The bedside diagnosis of lymphedema requires high level of suspicion and identification of specific findings on physical examination. Noninvasive diagnostic tools are helpful in confirming the diagnosis. Treatment with decongestive lymphatic therapy provides improvement in patient symptoms and volume reduction. This mini-review article highlights the pathophysiology, diagnosis and treatment of lymphedema.

Abbreviations: CAM: Complementary and Alternative Medicine; CR: Compression bandaging; CDT: Complete Decongestive Therapy; CG: Compression garments; CT: Computed tomography; IPC: Intermittent Pneumatic Compression; LE: Lymphedema; MLD: Manual lymphatic drainage; MRI: Magnetic resonance imaging; US: Ultrasound

Introduction

Lymphedema (LE) is a complex edematous condition that occurs when tissue homeostasis is jeopardized due to insufficient lymph transport [1]. It is characterized by the accumulation of regional interstitial protein-rich fluid [2]. Lymphedema is either primary or secondary. Primary lymphedema is classified according to the age when edema appeared. Congenital lymphedema occurs at birth and is apparent at two years of life. Lymphedema praecox occurs at puberty and lymphedema tarda is seen at those above the age of 35 years. Secondary lymphedema results from disease processes, post-surgery and axillary node clearance and after radiotherapy and irradiation. It is sometimes confused with filariasis.

Pathophysiology

LE is considered as the pathologic outcome of inadequate lymphatic outflow. Lymphatic aplasia, hypoplasia, valvular insufficiency, obliteration/disruption of lymphatic vessels and primary decreased lymphatic contractility are all direct causes of the pathology [3]. This leads to lymphatic hypertension and decreased contractility of lymph. As a result lymphostasis and accumulation of lymph, interstitial fluid, proteins and glycosaminoglycans takes place within the skin and subcutaneous tissue. This eventually stimulates collagen production by fibroblasts, disruption of elastic fibers, and activation of keratinocytes, fibroblasts and adipocytes where skin thickening and subcutaneous tissue fibrosis is the result [3-5]. Studies have shown that stasis of lymph causes accumulation of protein and cellular metabolites in the extracellular space that raises the tissue colloid osmotic pressure, causing water accumulation, formation of edema and increased interstitial hydraulic pressure [4,5].

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Treatment

LE can be treated in various ways. The Complete Decongestive Therapy (CDT) has been widely known and implemented in the treatment of LE and specifically post cancer related LE [15]. According to the Medicare Evidence Development and Coverage Development Committee, CDT includes 1) manual lymphatic drainage (MLD); 2) compression bandaging (CB) and/or garments (CG); 3) exercise; 4) skin care; and 5) sequential pumps [16]. The American LE Framework Project also showed that CDT is an effective intervention in reducing LE [17,18]. After CDT, self-maintenance is crucial involving self-MLD, day-time CG, CB, night-time exercise, skin care and nutrition [6]. To achieve this, patient education and awareness regarding LE management modalities is important including hygiene, moisturizing, sunscreen, avoidance of blood flow constriction and use of tourniquet [6]. Manual lymphatic drainage (MLD) is a massaging technique that enables the sequestration and lymph transport. The congested lymph moves from the restricted locations to the patent and intact nodal basins. Experts also recommend the use of CB during exercise [6]. Exercise should be tailored to the patient’s physical condition and abilities. In conclusion, CDT, MLD and CB controls symptoms and improves the quality of life for LE patients [15].

Surgical treatment for LE includes excisional/debulking, lymphatic reconstruction, or tissue transfer [19]. Excisional procedures involve removing the fibrofatty tissue that has formed secondary to sustained lymphatic fluid stasis [15]. Procedures include debulking, liposuction, and amputation are considered an option when standard LE treatment, such as CDT, has failed [15]. Studies have shown that excisional procedures had the highest LE volume loss (91.1%) compared to lymphatic reconstruction (52.9%) and tissue transfer (45.6%) [19]. It is worth noting that the majority of the procedures will be followed up with lifelong CG to preserve post-operative results [19].

Intermittent Pneumatic Compression (IPC) Therapy is well tolerated in low to moderate pressures and studies show that they may be part of a supervised multi-modality approach for home based patients with LE [20].

Complementary and Alternative Medicine (CAM) has been adopted as a means for LE management where patients practice mindfulness-based stress reduction and studies showed that it improves the quality of life and reduces psychological distress [21]. There is also a higher patients’ adherence rate with CAM and aqua lymphatic therapy [22]. Yoga, acupuncture/moxibustion, and Tai Chi breathing with arm exercises were shown to be safe by pilot data, but still more studies are needed to understand their role in LE management [23].

Conclusion

Lymphedema is a chronic condition that occurs due to insufficient lymph flow leading to limbs or genital swelling. It is important to properly diagnose LE to understand the extent and severity of extra fluid to tailor treatment accordingly. There are several diagnostic tools available and each has a specific method of implementation. LE is classified in general into primary or secondary. The treatment of lymphedema varies from complete decongestive therapy, surgery, pneumatic compression to complementary and alternative medicine.

Declaration

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