Lymphatic Capillaries in Aging

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Keywords
Lymphatic vessels · Aging · Atherosclerosis · Endothelial cells

Abstract
The lymphatic system is responsible for fluid drainage from almost every organ in the body. It sustains tissue homeostasis and is also a central part of the immune system. With the discovery of cell-specific markers and transgenic mouse models, it has become possible to gain some insight into the developmental and functional roles of lymphatic endothelial cells (LECs). Only recently, a more direct regulatory role has been assigned to LECs in their functions in immunity responses and chronic diseases. Here, we discuss the changes occurring in aged lymphatic system and the role of lymphatic capillaries in some age-related diseases and experimental animal models.

Introduction
The lymphatic system spans nearly the entire body. Lymphatic vessels are responsible for the drainage and unidirectional flow of fluid from the interstitial space back into the blood circulation. In contrast to capillaries of the blood system, lymphatic capillaries have so-called “button-like” junctions, that form loose cell-cell contact, and lack a continuous classical basement membrane. Moreover, the cell-cell junctions are disjoined and have a unique overlapping architecture which forms flaps that cause an increase in permeability, thus allowing for passive uptake of waste products and other macromolecules from the interstitium, as well as the entry of immune cells into the lymph. In addition, capillaries lack surrounding mural cells and are anchored to the extracellular matrix with the help of filaments that also assist them in their drainage function. Lymphatic capillaries start from endothelial occluded sacs. The lymph taken up by capillaries flows into the collecting lymphatic vessels, which have tighter “zipper-like” junctions and are wrapped around with mural cells [1]. Identification of lymphatic specific markers, such as the transcription factor prospero-related homeobox protein 1 (Prox1) and the cell surface proteins, namely lymphatic vessel endothelial hyaluronan receptor 1 (LYVE-1), vascular endothelial growth factor receptor 3 (VEGFR-3), and podoplanin, have enabled a great understanding of the lymphatic system [2–7]. These studies using mouse models have also enabled mapping of the tissue-specialized properties of initial and collecting vessels in different parts of the body. For instance, initial capillaries in the skin form a three-dimensional network, whereas lymphatic vessels in the lacteals of the...
gut are tube shaped that then fuse into the mesenteric collectors.

In contrast to the cardiovascular system, where circulation is powered by continuous pumping by the heart, the lymphatic system relies on either extrinsic forces such as muscle contractions from the surrounding tissues or on the intrinsic contraction by collecting vessels themselves [8–10]. Moreover, under physiological conditions, endothelial cells produce nitric oxide that maintains intrinsic pump activity [10, 11]. Lymph that has entered the lymphatic vessels is prevented from retrograde flow by lymphatic valves. Lymphatic vessels carry the lymph through the body via lymph nodes and eventually empty the fluid into subclavian or internal jugular veins via the right lymphatic duct or the thoracic duct, and the lymph rejoins the blood [1, 12]. On the luminal side, the lymphatic endothelial cells (LEC) are covered by proteoglycans and glycoproteins, which are anchored to the endothelial cell membrane, and that bind to free-floating plasma proteins and endothelium-derived molecules via their side chains. This meshwork of proteoglycans and glycoproteins together with soluble components bound to their branched side chains, is collectively called the glycosalx, and functions as a protective barrier [13]. Interestingly, the thickness and the coverage of the glycosalx declines with age within the collecting vessels [14].

The lymphatic system is an important part of the immune system. It carries immune cells, antigens, and pathogens to draining lymph nodes. Although it was historically viewed as just a passive carrier of various components of the immune system, research in the last two decades has revealed the vast immunoregulatory roles it plays. Cancer cells take advantage of lymphatic vessels as a route to metastasize. On the other hand, a well-functioning drainage system is pivotal for a strong immune response against pathogens or any other kind of trauma that disrupts tissue homeostasis, which could lead to abnormally expanded lymphatic vessels and lymphedema. During aging, the lymphatic system undergoes considerable changes with potentially pathological consequences [14–16]. In this review, we aim to summarize how lymphatic vessels impact age-related changes in three main sites of the body which are of special interest from a gerontological perspective, namely the skin, the arteries, and the brain. There are significant age-dependent changes in the appearance and function of skin. Atherosclerosis is an age-related disorder of arteries. Furthermore, the central nervous system undergoes age-related degradation. In each section, we will discuss the physiological function of lymphatic capillaries as deduced using experimental animal models and as well as their role in the pathobiology of age-related diseases (Fig. 1).

There are of course additional important points with regard to lymphatic vessels and aging, such as the impact of smooth muscle cells and contractility, mast cells, oxidative stress, and also the diverse tissue specific roles in organs such as the thoracic duct, ovaries, prostate and eye, to mention a few. Those topics, however, are beyond the scope of this Viewpoint article, and can be found elsewhere [17].

Skin

The skin is the body’s largest organ and the foremost target for external insult. Aging in the skin is manifested by loss of elasticity, wrinkles, and age spots. The mecha-
The mouse model K14-VEGFR-3-Ig that lacks dermal lymphatics has proved to be useful for studying skin drainage. In one study in which trypan blue was injected intradermally or skin was painted with FITC, K14-VEGFR-3-Ig mice were found to have impaired drainage and dendritic cell migration to skin draining lymph nodes, compared to wild-type animals. Moreover, these mice displayed an altered lymph node architecture, with B cells scattered throughout the nodes as opposed to clustering into B-cell follicles as in normal lymph nodes. Moderately aged K14-VEGFR-3-Ig mice (1 year old) had increased titers of IgG1, IgG3, and IgA antibodies in the serum as well as antibody deposits in the skin. This study suggests that dermal lymphatics are important in antigen and cell drainage from the skin, and if not functioning properly, could contribute to an age-associated autoimmune phenotype [23].

As mentioned above, lymphatic drainage function declines with aging. Using luminescent dyes that were injected intradermally into the ear of young (2 months), middle-aged (7 months), and old (18 months) mice, it was shown that clearance of the dye was impaired in the older animals. Reduced drainage was directly related to fewer and smaller LYVE-1+ vessels [15]. Moreover, site-specific delivery of VEGF-C to skin, with an engineered fusion protein where VEGF-C was fused to an antibody which recognizes an inflammation-induced component of fibronectin, expanded the lymphatic vessel area in inflamed skin. The authors used two models to induce psoriasis in mice, and in both cases, the fusion protein improved lymphatic drainage and reduced accumulation of inflammatory T cells [24]. In photoaged skin, the number of lymphatic vessels is reduced, and these aged vessels in addition show enhanced permeability. Exposure of a single burst of UVB to mouse skin was followed by enhanced macrophage infiltration and downregulation of VEGF-C. Edema in these mice was rescued by treatment with an intradermal injection of a mutant VEGF-C that specifically binds to VEGFR-3, and induces lymphangiogenesis [25].

Thus, as far as skin lymphatics are concerned, they could either contribute to sustain homeostasis through drainage, inhibit T cell responses in a melanoma setting, or allow for more distant metastases in aged skin. Therefore, it seems that dermal lymphatic vessels could play either a beneficial or harmful role. In aged skin, contractility and drainage function declines, whereas permeability increases, and collectively the homeostatic function of lymphatic capillaries and collectors decreases.
Arteries

Arteries are large, macroscopically visible blood vessels of the body that flow into arterioles and finally capillaries. They consist of three layers: tunica intima, the innermost monolayer of endothelial cells in direct contact with the blood, tunica media, a muscular layer, and the tunica adventitia, the outermost layer attaching the vessel to the surrounding tissue. The adventitia contains both blood and lymphatic vessels under physiological conditions, with the latter necessary for sustaining homeostatic drainage. Normal function of the blood vasculature includes distribution of oxygen, nutrients, and hormones throughout the body. Moreover, immune cells, blood cells, and platelets also travel through the cardiovascular system. Arteries regulate the blood pressure through molecules within the renin-angiotensin-aldosterone system, a cross-talk mechanism between the kidneys and arteries that is in direct relation to fluid and salt intake [26]. Regular exercise, a physiological means to activate lymphatic circulation, has a beneficial impact on numerous aspects of the cardiovascular system [27].

Atherosclerosis is a multifactorial, chronic inflammatory disease of the arteries that is characterized by abnormal immune cell accumulation, lipid uptake as well as calcification and necrosis within the arterial tunica intima. In the beginning of atherogenesis, mononuclear cells, such as T cells and macrophages, enter the arterial intima via adhesion molecules which are expressed at predisposed sites on stressed blood endothelial cells [28]. We have previously shown that autoreactive T cells first recognize endogenous heat shock protein 60 (HSP60) expressed on the surface of blood endothelial cells [29]. However, other autoantigens have also been proposed to drive disease, such as oxidized low-density lipoprotein and apolipoprotein B-100 [30]. Classical atherosclerosis risk factors, such as smoking, obesity, hypercholesterolemia, and low laminar blood flow, all act as endothelial cell stressors and induce HSP60 expression on the surface of endothelial cells, thus becoming a target for preexisting autoreactive T cells [28, 29]. In relation to classical risk factors and blood pressure, we have previously demonstrated that high amounts of salt can also directly stress cultured endothelial cells and cause ectopic surface HSP60 expression [31].

Although the disease starts at a young age and takes decades to develop, atherosclerosis manifests itself clinically at an older age. As the disease progresses, the arteries thicken and a nutrient and oxygen-depleted core is formed [28]. To combat this, under physiological conditions, small blood vessels, so-called vasa vasorum (Latin: the vessels of the vessels) originating from blood vessels that are found in the adventitia are formed by the angiogenic process. Blood vessels formed in plaques are often leaky, and neovascularization within the plaque is considered to contribute to unstable plaques with increased risk of rupture [32]. In addition, advanced plaques also develop lymphatic vessels within the intima and media [33].
Almost 50 years ago, Wolinsky and Glagov [34] showed that *vasa vasorum* also form in the tunica media of seemingly healthy arteries that contain 29 or more layers of smooth muscle cells. This finding, exactly reproduced in our own studies, has curiously remained unknown to the larger atherosclerosis community. Moreover, this phenomenon is age- and species-specific, as smaller species, such as the mouse, never seem to have more than five layers, whereas a human infant is already born with 34 layers of smooth muscle cells in aortas [34]. Based on the findings of Wolinsky and Glagov, we speculated that lymphatic vessels might also be present within normal arteries with >29 layers of smooth muscle cells. To this end, we collected normal and early lesion arterial vessels of the ascending aortae from patients undergoing surgery (unpublished results). Indeed, all samples contained more than 29 smooth muscle cell layers, with 50% having podoplanin+CD31− lymphatic vessels in the media (Fig. 2).

The function of lymphatic vessels within the arterial media can only be speculated upon based on studies in mice. LEC conduits in the adventitia beneath advanced plaques in mice were shown to be essential in draining cholesterol buildup [35]. Specifically, macrophage reverse cholesterol transport, the movement of cholesterol from tissues back to the liver was impaired when aortae from *ApoE−/−* mice were transplanted into donor mice that were treated with anti-VEGFR-3 antibody. The antibody treatment inhibited lymphatic vessel regrowth and in turn cholesterol uptake by LECs, leading to cholesterol buildup in the plaque [35]. Another study showed that treating *Ldlr−/−* mice that had diet-induced atherosclerosis with VEGF-C, a factor that promotes lymphangiogenesis, stabilizes both the plaque and the drainage function of lymphatics [36]. Both aged mice and humans develop tertiary lymphoid organs (TLOs) with lymphatic vessels within the adventitia, directly adjacent to plaques [37, 38]. In secondary lymphoid structures, such as the lymph nodes, lymphatic vessels function as a cell entry route to the afferent lymphatics and as a cell exit route at the efferent lymphatics [39]. Consequently, lymphatic vessels in disease-associated TLOs might have a similar function. In addition, LECs have been shown to directly modulate immune response by presenting antigens to both CD4 and CD8 T cells, inducing tolerance or deleting auto-reactive cells, respectively [40, 41]. Lymph node LECs express high levels of the T cell inhibitory molecule PD-L1 [42], although to our knowledge, no such finding has yet been reported in arterial LECs. In our studies, we could not observe lymphatic vessel-associated mononuclear cells in the media, and therefore assign a draining function to these vessels. However, a study in mice showed that adventitial lymphatic capillaries could be a route for T cell exit from the plaque, in a CXCR4/CXCL12-dependent manner [43]. Thus, experimental studies in mice and postsurgical studies in humans generally suggest that blood vessels within arterial plaques are detrimental to disease outcome, whereas lymphatic vessels seem to improve the pathophysiology of atherosclerosis [44]. Future studies are necessary to translate the findings in mouse arterial lymphatic vessels to human conditions under both physiological and inflammatory settings. Specifically, in relation to our finding that roughly half of the cohort of early lesion arteries had lymphatic vessels within the media, it would be of importance to find out if these lymphatic *vasa vasorum* are a prerequisite for stable plaques.

### The Brain and the Central Nervous System

Although meningeal lymphatic vessels had been mentioned as early as the end of the 18th century, their existence has been largely forgotten by the scientific community. In 2015, meningeal lymphatics in the brain were rediscovered in rodents by two separate groups [45, 46], and later also described in the spinal cord [47], thanks to advanced imaging technology. Up until then, the brain was thought to be an organ devoid of a lymphatic drainage system. The presence of such a system in humans was confirmed 2 years later [48]. Immune cell trafficking to and from the brain was thought to occur only through the blood vasculature. Drainage was assigned to both the blood vasculature and the lymphatic system [49, 50]. The new findings show that brain lymphatic vessels are located in the dura mater and are able to drain cerebrospinal fluid (CSF) into the deep cervical lymph nodes. They express LYVE-1, Prox1, and VEGFR-3. The studies used fluorescent tracers to exclude uptake by venous vessels [45, 46]. Development of the meningeal lymphatics was shown to be dependent on VEGF-C and its receptor VEGFR-3, and in mice these were fully developed 1 month after birth [47]. Interestingly, deletion of VEGFR-3 specifically in LECs in adult mice led to a regression of the meningeal lymphatics, as well as reduced drainage of CSF. Therefore, both development and function of brain lymphatics is dependent on VEGFR-3 signaling [47].

Dural lymphatics have prompted a whole new take on age-related cognitive function and brain disorders, such
as Alzheimer’s disease. Drug-induced disruption of the brain lymphatics in a mouse model led to cognitive impairment, and a reduction in CSF and brain interstitial fluid drainage [51]. In aged mice, a decrease in the meningeal lymphatic vessel diameter and an increase in amyloid-β accumulation in the CSF suggest the dural lymphatics as a significant route to drain amyloid-β [51]. Another study suggested the lymphatic vessels in the brain as the main site for draining CSF, and also observed a reduction in CSF draining in aged mice [52]. However, brain fluid clearance is mainly managed by the perivascular and glymphatic system [53]. In addition to amyloid-β, both intracellular and extracellular tau have been implicated in Alzheimer’s disease and other neurodegenerative diseases. In a study where mice that lack dural lymphatics were used (K14-VEGFR3-lg mice), injection of fluorescently labeled tau into the parenchyma showed significantly lower drainage from the brain compared to wild-type mice [54]. In a multiple sclerosis model, trafficking of autoreactive T cells through dural lymphatic vessels in a CCR7-dependent manner was directly observed [55]. Fluid uptake was first demonstrated mainly on the dorsal site [45, 46]. This has now been expanded to also include the basal part of the skull, where the structure of meningeal lymphatic vessels seems to be different from that in the dorsal region, containing both capillary and collecting vessels [56]. The authors also concluded that the basal vessels are much better in the uptake of CSF than the dorsal vessels. Lastly, decreased function of basal lymphatic vessels has been reported in aged mice [56].

Furthermore, dural lymphatics have been implicated in brain swelling (hydrocephalus), multiple sclerosis, stroke, and various neurodegenerative disorders. As the field of brain lymphatics is still young, there are more questions waiting to be answered by future studies [57].

Summary

The availability of new reagents that enable identification of LECs by various imaging methods has provided unexpected insights into the function of the lymphatic transport system in healthy young and adult animals and humans. Interestingly, there are very few studies addressing age-dependent alterations of the lymphatic system and its functional consequences. Here, we briefly review the most important findings on this topic in relation to the brain, the skin, and the vascular system. The brain is the target for a number of neurodegenerative conditions, such as Alzheimer’s disease, where impaired lymphatic drainage seems to play an important pathogenetic role. The skin obviously is the organ where age-related changes are most visible and where studies of morphological and functional alterations can be readily conducted. Finally, in the field of cardiovascular research, very little work has been done on the role of the lymphatic system in the development of atherosclerosis, the most frequent age-dependent disease. Here, we present data showing the existence of lymphatic vasa vasorum not only in atherosclerotic lesions but also in the wall of normal, healthy arteries. Thus, lymphatic vessels obviously play important roles in normal and pathological aging, and it is therefore surprising that this interesting topic has so far received only scarce interest in basic and applied gerontological research.

Acknowledgments

The authors would like to thank former members of the LAI, as well as Dr. Christoph Krapf and Prof. Michael Grimm from the Department of Cardiac Surgery, Innsbruck, Austria, for making human surgical arterial specimens available to us and Christian Höpperger and Prof. Christian Ensinger from the Department of General Pathology at the Medical University Innsbruck, for their help with histological staining.

Statement of Ethics

Patients undergoing routine aortic vessel and arterial bypass surgery were recruited and the samples were collected with written informed consent, according to the regulations of the Ethics Committee of the Medical University of Innsbruck Ethics (Approval No.: UN5215 328/4.26 [3404b]).

Disclosure Statement

The authors have no conflicts of interest to disclose.

Funding Sources

This research was funded by the Austrian National Bank (Grant No. 15953) (G.W), the Science Fund of the State of Tyrol (Grant No. UNI-0404/1395) (B.J.) and the Lore-and-Udo Saldow Donation (B.J.).

Author Contributions

B.J. and G.W. wrote the first draft of the manuscript. D.K. contributed with expertise scientific advice. All authors contributed to the editing of the final manuscript.
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Gerontology 2020;66:419–426
DOI: 10.1159/000508459

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