Current Knowledge and New Challenges in Exercise Immunology

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SUMMARY

Exercise has a significant effect on the regulation of the immune system. Acute bouts of exercise induce an intensity-dependent leukocytosis, followed by redistribution of effector cells into peripheral tissues. These processes are a result of the activation of the sympathetic nervous system and the hypothalamic-pituitary-adrenal axis, increased hemodynamics, and the release of stress hormones such as catecholamines and glucocorticoids.

During periods of intensive training, athletes frequently report increased symptoms of upper respiratory tract infections (URTI) which may be a result of a stress overload associated with an impairment of mucosal immunity. In contrast, single bouts of short-term or moderate intensity as well as regular exercise training are mainly “immuno-enhancing”.

In this regard, exercise positively affects the composition of the T cell compartment and the function of various leukocyte subpopulations. Regular exercise of moderate intensity also exerts immune-regulating effects during inflammation-associated diseases, such as obesity, type 2 diabetes, or cardiovascular diseases, and counteracts hallmarks of immunosenescence.

Consequently, exercise represents a powerful behavioral intervention that has the potential to improve both immune function and health outcomes in subjects of all ages in prevention and therapy. The present review summarizes the current knowledge, new challenges and future directions in the field of exercise immunology.

KEY WORDS:
Inflammation, Physical Activity, Lymphocytes, Mucosal Immunity, Neutrophils, NK Cells

Introduction

Exercise immunology is a dynamic developing sub-discipline of sports medicine. Although the knowledge about the diverse relationships between exercise and the immune system is profound, the complexity of immunity and its bidirectional interactions with other physiological systems continuously represents an enormous challenge. However, the present review summarizes the current knowledge, new challenges and future directions in the field of exercise immunology.
Effects of Exercise on Infection Risk

Since early reports on a higher incidence of infections of the upper respiratory tract (URTIs) in endurance athletes after periods of intensified training and competitions and the development of the open window theory (37, 65), immunology research focused primarily on the acute and chronic effects of different types of exercise on immune system parameters and the identification of potential key risk factors. During the last years, there is increasing evidence, which challenges this theory as discussed later on in this chapter. On the other hand, it is generally accepted that regular and moderate physical exercise improves the hosts immunocompetence and protects against infectious diseases. Therefore, in this section a distinction is made between the athlete’s population and a public health perspective.

The most frequent illnesses in athletes are URTIs and these infections are more likely to occur during intense training cycles and after tapering or after competitions than having seasonal patterns. There are early reports showing a 2–6-fold increase in the reported symptoms of URTI for several weeks after running a marathon and an association with the individual running time (65). Most of the evidence initially originated from studies, which involved running disciplines, but a series of data has confirmed the association between heavy exercise and URTI in other sports (65). Subsequently, much effort has been spent to the identification of potential mechanisms and risk factors for the increased URTI susceptibility in athletes. However, it is challenging to interpret the clinical impact of changes in single immune system parameters because the immune system as a whole developed great redundancy. Nevertheless, a number of studies have shown that an acute bout of exercise, depending on mode, intensity and duration, is associated with a transient change in numerous immune system parameters such as peripheral leukocyte numbers and function or salivary immunoglobulin concentration (see section 5). A genetic basis for a higher susceptibility to URTI has been proposed by both exercise and non-exercise studies demonstrating that the regulation of several anti-inflammatory pro-inflammatory cytokines is characteristic for illness prone individuals (20). Findings from studies investigating mucosal immune system (MIS) integrity are more controversial, as some reported on associations between saliva Immunoglobulin A (sIgA) concentrations and URTI incidence while others did not (65).

However, since many studies from the past monitored URTI incidence by self-reports but neither verified an infectious cause by a laboratory identification of pathogens nor by a physical examination, the open window theory has been called into question. It is still debatable whether the URTI symptoms, which particularly occur in high performance athletes, are actually caused by infectious agents (7, 65). Further, it has been shown that there is a discrepancy between the physician’s URTI diagnosis frequency and verified infections by pathological tests (11). The latter demonstrate that bacterial and viral pathogens coincide URTI episodes in only 5% and 30–40% of cases of URTI episodes, respectively (11, 54). To date, there is increasing evidence that rather other exercise-induced inflammatory stimuli or a viral reactivation (7, 21) are the major cause of the upper respiratory symptoms (URS). Cox et al. (11) examined the effectiveness of a prophylactic administration of an antiviral agent for the control of Epstein Barr Virus (EBV) reactivation and URS in elite distance runners. They showed that the treatment resulted in a significant (∼-82%) reduction in the detectable EBV load in saliva without any difference in the URS incidence compared to the control group. A series of studies demonstrated that URS might be caused by a local airway inflammation instead. It has been speculated that the latter could be mediated by various mechanisms such as physico-chemical strains, asthmatic reactions and the migration of inflammatory cytokines, e.g. originating from the damaged skeletal muscle to the respiratory system (65, 66). Taken together, these data raise reasonable doubt on the validity of the open window theory and suggest other non-infectious causes for URS in elite athletes. Apart from that, even if opportunistic infections are causative for URS in athletes, it is believed that additional/or other internal and external factors which precede or parallel episodes of intensified training or competition, such as mental stress and anxiety, sleep disruption, traveling, nutritional deficiencies, and exposure to climatic changes etc. likely play a key role in the impaired immune competence (7, 66). Additionally, endurance competitions per se increase the risk of acquiring novel infections due to the exposure to human crowds (7).

Apart from the detrimental effects of exhaustive exercise periods, there is limited evidence that master athletes are predisposed to have rather a more efficacious immune defense as they suffer from fewer URTI episodes, which allows them to realize higher training volumes and thereby to compete more successfully (66). It is speculated that beside genetic factors (62) this population is characterized by behavioral factors such as a more effective hygiene routine, diet, sleep and stress management (66). Nevertheless, illness frequencies in athletic populations do not appear to be higher than in the general population (7) but they do not show such a typical relationship of symptom episodes to seasonal patterns (11).

Beside from athletic populations and competitive sports, regular exercise training is a major protective factor as it ameliorates immune function, whereas a sedentary lifestyle depresses immunity (43, 44). Both experimental and epidemiological research studies indicate a negative relationship between activity level and both, URTI incidence (43, 44) and the severity of URTI symptoms (43). In a 12-month observational study Matthews et al. (37) evaluated the activity level of 547 healthy adults (age 20–70 years). After adjustment for a number of potential confounders including age, education, psychosocial measures and dietary factors, they showed that moderate levels of physical activity reduce the risk of URTI about 20%. Nonetheless, due to methodological limitations in the majority of research studies, a Cochrane review from 2015 declares that it is impossible to determine the effectiveness of exercise in altering the occurrence, severity, or duration of URS to date (22).

Effects of Acute Exercise on Leukocyte Numbers and Function

Acute bouts of exercise are well-known to modulate the innate as well as adaptive immune response by affecting immune cell numbers and functions (29, 65). Basically, the innate immune response represents the first line of defense against harmful pathogens and consists of cellular components such as granulocytes, macrophages, natural killer cells (NK), dendritic cells (DC) as well as soluble factors such as
complement proteins among others (13). As an essential part of the innate immune response, granulocyte neutrophils represent a type of granula-secreting phagocytes that migrate to the inflamed tissue as first-responders to inflammatory cells (60). Acute exercise induces a first rapid neutrophilia, followed by a second delayed rise of blood neutrophil count a few hours later, depending on the duration and intensity of exercise (65). Novel findings let assume that the priming of neutrophils may be implicated in skeletal muscle inflammation after high-intensity exercise (45). On a functional level, unstimulated neutrophils respond to acute exercise stress with increased degranulation, oxidative burst and phagocytosis (65).

Additionally, after endurance exercise, the concentration of some complement proteins, such as C5a, is increased in the peripheral blood, promoting the chemotaxis of neutrophils (57). Monocytes are the largest type of blood leukocytes that can differentiate into macrophages or DCs after migration into damaged tissues and serve phagocytosis, antigen-presentation and cytokine production (72). With regard to monocytes, acute exercise induces a transient monocytosis of approximately two hours which is most likely due to a shift of monocytes from the marginalized to the circulating pool (65). Preferentially, acute exercise stimulates the mobilization of the proinflammatory CD14+/CD16+ monocyte subtype relative to classical CD14+/CD16- monocytes (29). To date, there is still a lack of studies investigating the exercise-induced mobilization of DCs, which act in the initiation of immune response by antigen presentation (24).

Interestingly, a recent study demonstrated a near fourfold increase of the number of monocyte-derived DCs (mo-DCs) after dynamic running exercise until volitional exhaustion without comprise of functional parameters (34). Since DCs are characterized by their unique ability to cross-present viral and tumor antigens, repetitive bouts of acute exercise might promote beneficial effects for health (24). NK cells are a minor subpopulation of lymphocytes which exhibit cytotoxicity and secrete cytokines promoting defense mechanisms against tumor- and virus-infected cells (70). Similarly to other blood leukocytes, NK-cells show a rapid mobilization in response to an acute bout of exercise (29). Preferentially, the cytotoxic CD56dim NK cell subset rather than the INF-Y producing CD56bright subtype appears to be mobilized by exercise (65). Moreover, functional changes in NK cell cytotoxicity depend on the intensity and duration of exercise. While short bouts of low- to moderate intensity exercise induce an increase, NK cell cytotoxicity expressed on a per cell basis can be reduced for hours after exhausting and prolonged exercise, probably favoring an increased susceptibility to infectious diseases (29, 65).

Contrary to the innate immune response, the adaptive immunity is characterized by antigenic specificity and immunological memory and is composed of two main types of lymphocytes termed T cells and B cells (13). T cells are derived from the bone marrow and migrate to the thymus, where MHC-receptors are formed on their surface, allowing them to identify specifically foreign antigens. Furthermore, the differentiation process includes the maturation into diverse subpopulations such as CD4+ T-helper, CD8+ cytotoxic and regulatory T cells (Treg) (46). B cells are the carriers of the humoral immune response and are the only cells capable of producing antibodies after differentiation into plasma cells (3). Acute exercise results in transient biphasic changes in the numbers of circulating lymphocytes marked by lymphocytosis during and immediately after exercise, while numbers of cells sinking below basal levels during the early recovery period (26).

Studies indicate that the initial increase of lymphocytes may be the result of mobilization from the marginal pool, the spleen as well as other lymphoid tissues, while the exercise-induced lymphopenia in the post-exercise period seems to be due to redistribution in lymphoid- and non-lymphoid organs and apoptosis (27, 28). Particularly, exercise-induced mobilization of T cells seems to be affected by differentiation stage. Accordingly, highly differentiated T cells show a more frequent mobilization response to acute exercise compared to low differentiated T cells (51, 53). Unlike T cells, B cell mobilization does not seem to be triggered by effector status. In response to acute exercise, the immature B cell subpopulation displays the largest increase, followed by memory B cells and naive B cells (63).

However, recent studies reveal that the behavior of the exercise-induced mobilization of differentiated T cells is highly dependent on exercise intensity, training status, and individual predispositions such as CMV serostatus and age (12, 33, 51). Thus, high-intensity loads induce greater apoptosis in highly differentiated T cells than moderate intensity exercise (27). Moreover, high physical fitness individuals show a greater decrease of mobilized highly differentiated effector memory T cells (mTeff) in the early post-exercise period, possibly indicating more efficient clearance mechanisms (12). Independently of exercise intensity CMV-seronegative athletes mobilize more low- and medium differentiated T cells than CMV-seropositive (33). Further, the amount of CD34+ hematopoietic progenitor cells and angiogenic T cells (TANG) is inversely correlated with age and older adults display attenuated exercise-induced mobilization of these cell types (51).

To date, only a limited number of studies are available investigating the effects of acute exercise on Treg cells. Tregs are a specialized subgroup of T cells that modulate the activation of the immune system by suppressing the activation and proliferation of effector T-cells as well as maintaining tolerance to self-antigens (6, 7). Novel studies demonstrated that Treg cells are mobilized in response to high-intensity exercise stress (12, 27). In this context, a higher level of physical fitness appears to promote a stronger mobilization (12). On the one hand, Treg mobilization amplifies exercise-induced immunosuppression, but on the other hand, it may also have immune regulatory effects (67).

Novel studies suggest that chronic exercise has the potential to modulate both the number and function of cells of the innate immunity (1, 8, 50, 59). Although exercise training seems to reduce the neutrophil count only under conditions of chronic inflammation, regular exercise has a significant impact on neutrophil function (65). Recently it was reported that four weeks of hypoxic exercise training in sedentary healthy men increased the expression of adhesion molecules and opsonic receptors, and also enhanced the bactericidal capacity and apoptosis of neutrophils at basal conditions and after acute exhausting exercise (8).

However, exercise modalities seem to have powerful impact on markers of neutrophil function, as normoxic exercise training did not show any effect (8). In response to
chronic exercise training, it was demonstrated that physically active individuals show a lower percentage of the pro-inflammatory CD14+CD16+ monocyte subtype (29, 56). It seems likely that this reduction contributes to the anti-inflammatory effects of chronic exercise. The evidence for the effects of exercise training on DCs is weak, as to date only results from animal models are available. In rat model, DC numbers increased after chronic exercise. Moreover, regular exercise amplified major histocompatibility complex II (MHC II) expression on rat DCs and increased their production of IL-12, a cytokine that modulates the differentiation of naïve T cells and stimulates T cell growth and function (9, 35). Indeed, exercise-induced increases in DC number may protect against cancer and virus infections because of their ability to present viral and tumoral antigens supporting balanced regulation of immune function.

Nevertheless, future human studies that verify the behavior of DCs in response to chronic exercise are essential. With regard to the effects of exercise training on the frequency and cytotoxicity of NK cells, the results of many studies are controversial due to methodological discrepancies and different exercise regimens (type, duration, intensity) (71). While some studies indicated an increased NK cell frequency and higher NK cell cytotoxicity in athletes compared to sedentary controls, others reported no effects (43). In contrast to studies with healthy young subjects, most studies including healthy older subjects have shown no effects of chronic exercise on the cytotoxicity of NK cells (71). Regarding the impact of chronic exercise on peripheral lymphocyte numbers, most longitudinal studies have failed to demonstrate any major changes (65).

Though, current cross-sectional studies reported a VO2max dependent decrease of the proportion of total peripheral lymphocytes as physiological adaptation to high-volume endurance exercise training (2, 41). Consistently, T and B cell function appears to be modulated by regular exercise. Especially, well-trained athletes undertaking high-intensive training regimens showed decreased numbers of cytotoxic CD8+ T cells, decreased T cell proliferation and a decline in stimulated B cell immunoglobulin (Ig) synthesis (2, 32, 64). Therefore, athletes who train for long periods of time in high-volume and high-intensity exercise loads may be more prone to infectious diseases through reductions in T cell functionality.

However, recent studies demonstrated an increased Treg cell frequency in athletes (67). Due to their anti-inflammatory properties, Tregs could at least partially compensate impaired immune function during periods of heavy exercise loads. Moreover, lymphocyte apoptosis sensitivity is associated with aerobic endurance training status (2, 26). After ex vivo cultivation, lymphocytes of athletes showed reduced sensitivity to specific apoptosis inducers such as PHA-L. Although lymphocyte apoptosis was similar under basal conditions between athletes and non-athletes, molecular signatures differed in lymphocytes of highly endurance trained athletes characterized by a balanced modulation of pro- and anti-apoptotic apoptosis proteins and micro-RNAs such as XIAP, BAK1, miR-23a and miR-27a in athletes (2).

Hence, it is suggested that lymphocytes adapt to exercise training by increasing their resistance to apoptosis and facilitating lymphocyte homeostasis. By this, lymphocytes of athletes may be better prepared to combat the transient immune suppression post-exercise (2). Accordingly, it has been reported that lymphocytes adapt to chronic exercise by up-regulation of cellular defense systems such as heatshock protein 72 (HSP72) (16). In addition, recent studies show that the proportions of low differentiated and highly differentiated T cell subpopulations are also modulated by regular exercise (12, 59). Thus, at baseline, the proportions of mTeff were lower, while memory regulatory T cells (mTreg) were higher in high physical fitness subjects (12). Considerably, a study discovered changes in naïve and memory T cells in elite swimmers even during season (59). During the initial training period characterized by a gradually increasing load, CD4+ naïve, CD8+ naïve, gamma-delta effector-memory (γδ EM) and central-memory (CM) T cells increased in peripheral blood, while γδ terminal effector (TEMRA) T cells decreased. Though, there was an increase in CD8+ TEMRA.
Effects of Exercise on Mucosal Immunity

The MIS represents a network of mechanical, cellular and humoral factors which are integrative regulated to exert protection against invading antigens. MIS can be mainly found at the nasal passages, the respiratory tract, and the intestines. It offers a protective role at all mucosal surfaces, and the effectiveness of its immunity is based on mucosal epithelial cells (ECs) covered by mucus and antimicrobial products and fortified by both innate and adaptive components of host defense (39, 40, 46).

High rates of infections in athletes have led to an increasing focus on the mechanisms of mucosal homeostasis in the respiratory tract during exercise. It is well-known that neuronal signals and stress responses are important regulators of the immunity at mucosal sites. These include psychological stress, various environmental stressors, and exercise. The majority of studies in the field of exercise and mucosal immunity have focused on the respiratory tract. Most studies presented changes in the secretion of SIgA in saliva after exercise and training (65). These results are partly inconsistent, because methods of saliva collection and diurnal variations importantly affect the results. However, after acute intensive endurance exercise most studies observed a reduction of SIgA which seems to be associated with an increased risk of URTI in athletes. In contrast, exercise and training of moderate intensities can increase IgA levels and might be associated with a reduced incidence of infections (20). Accordingly, SIgA has shown moderate diagnostic value, while its utility is limited due to large intra- and inter-individual variation (65). Since SIgA is affected by vegetative nerve stimulation, it is speculated that exercise of long duration and high intensity induces the activation of the sympathetic nervous system, followed by an inhibition of salivary IgA synthesis, exocytosis, and transcytosis (20). Saliva concentrations of α-amylase and lysozyme increase after intensive exercise, but no association to athletes’ infection rates have been found so far (65).

Regarding the effect of exercise on intestinal mucosal surfaces there are less data available. While some recent data provide evidence for health promoting effects of exercise on microbiota and gut barrier function, some human studies demonstrated perturbations of the mucosal immune homeostasis and an increased intestinal permeability especially after exercise of long duration in hot environment (73). In these subjects a modest circulatory endotoxaemia was observed which was accompanied by a pro-inflammatory cytokinaemia. However, the underlying mechanisms and the clinical relevance is still discussed (18).

Various molecular mechanisms have been identified which mediate acute and chronic exercise effects on the adaptive and innate immune function. These mechanisms are versatile, but also integrative and contiguous. An important process seems to be the increased hemodynamics during exercises, including an increase in cardiac output, vascular vasodilatation, and blood flow. The greater mechanical forces on the endothelium induce leukocytes demargination and redistribution throughout the body. An important driver of this process is the individuals stress response mediated by the sympathetic nervous system (SNS) and the hypothalamic–pituitary–adrenal axis (HPA axis). The activation of both axes produce signals for receptors on a variety of immune cell subtypes (14). Norepinephrine and epinephrine represent ligands for adrenergic receptors which can be found on T cells, monocytes, and NK cells. After stimulation, important cellular functions, such as proliferation, differentiation, cytokine production, migration properties and trafficking, antibody production, or cytotoxicity are affected, impacting a major role for adrenergic mechanisms for regulating central immune cell functions (14). Mobilization of hematopoietic progenitor cells seems to depend on β2-adrenergic mechanisms as well (1).

Moreover, the action of catecholamines seems to have immune-regulating effects, since adrenoceptor activation downregulates the lipopolysaccharide (LPS)-induced production of inflammatory cytokines. The release of cortisol from the adrenal cortex affects leukocyte mobilization, redistribution, and exhibits anti-inflammatory effects by inhibiting various immune cell functions (40). Recent data provide evidence that cytokines, such as IL-6 and IL-7, and proteases are also important factors for immune cell activation, proliferation and survival. However, their particular role on the immune cell function during exercise is not quite clear (58). An important role for IL-6 and matrix metalloproteinases (MMPs) was proved for leukocyte redistribution during pathological conditions, such as cancer (49). Similarly, epigenetic mechanisms gained progressively interest in exercise immunology.

Accordingly, exercise affects epigenetic regulation of gene expression by various mechanisms, such as DNA methylation, histone modification and alterations in microRNA profiles. Some recent data showed immunological health benefits due to exercise training through epigenetic mechanisms. However, actually this research is a growing field and has to be integrated in the current knowledge in exercise immunology (17, 52). Regarding intracellular signaling, the exercise-induced stress stimulus has been shown to affect various intracellular signaling mechanisms, such as an altered Ca2+ signaling, changes in metabolic intermediates such as the NAD+ /NADH and AMP/ATP ratio, phosphorylation of kinases, cellular repair and recycling processes, and oxidative balance (58).

With regard to oxidative stress, regular exercise training increases the body’s antioxidant defense system followed by the prevention of oxidative damage to cellular structures. Due to the close bi-directional relationship between oxidative stress and inflammation an exercise-induced increase of anti-oxidative enzymes, such as superoxide dismutase (SOD), might be a favorable immune-regulating adaptation in response to training (58).
The term immunosenescence describes the process of progressive dysfunction during aging. Age-decline of various immunological processes contributes to the increased susceptibility of elderly persons to acute and chronic infections, autoimmune diseases, systemic inflammatory diseases, and cancer. Central functions of both the innate as well as the adaptive immune system decline in older people, and the adaptive response seems to be more affected (46, 48).

While T cell number remains constant during aging, the quality of the T cell compartment undergoes an important remodeling process. Accordingly, aging is associated with a slight and progressive decline of the CD4/CD8 T-cell ratio alongside a diminution in numbers of naïve T cells. Instead, more differentiated memory cells accumulate, which can be most profoundly observed within the CD8+ T cell compartment (48). Here, an increased percentage of T cells lacking expression of the lymphoid homing receptor CCR7 and the costimulatory molecules CD27 and CD28 alongside reexpressing CD45RA, can be found. These cells, termed T effector memory cells re-expressing CD45RA (TEMRA) cells, have a reduced ability to proliferate upon antigen contact, but produce large amounts of pro-inflammatory cytokines allowing them to potentially participate in immune pathology (48, 50).

Cellular changes are accompanied by profound modifications of the systemic cytokine network. While levels of IL-2 slightly decrease, aging is accompanied by a two- to four-fold increase in circulating cytokines, chemokines, and proteases, collectively termed ‘gerokines’. The progressive increase in gerokines, such as TNF-α, IL-6 and C-reactive protein (CRP) (48) reflects a generally increased basal activity of the entire inflammatory network that has recently been termed ‘inflamm-aging’ (48). Inflamm-aging is amplified by the existence of other immunological risk factors, such as obesity or smoking, and the blood stream represents an important distributor of inflammatory signals throughout the body.

Consequently, systemic inflammatory signals affect local, age-associated cellular changes to a senescence-associated secretory phenotype (SASP) that can have deleterious effects on the tissue microenvironment (5, 30, 31). Increased systemic inflammation seems to be inversely associated with a decline of immune function which mainly affects the T cell compartment (30, 31) (Figure 2). During exercise, older subjects were reported to mobilize substantially fewer naïve CD8+ T cells compared to younger people. These changes are attributed to a reduction in thymic output and to a reduced sensitivity of β2-ARs during aging, which negatively affects catecholamine-induced immune cell redeployment (28).

Recent studies proved that exercise is able to reverse hallmarks of immune senescence, specifically with regard to T cells. It is widely accepted, that regular endurance exercise such as running or walking, increases CD4+/CD8+ ratio in older adults (65). Our own group demonstrated that regular endurance training is followed by proportional increases of naïve and central memory T cells, while proportions of CD8+ TEMRA cells decreased in older pre-diabetic subjects (50). Acute and chronic exercise are followed by an increase of circulating haematopoietic progenitor cells (HPCs) which have been shown to decline during aging (27, 29). The mobilization of cells might reflect an increased production of myeloid lineage cells and a higher lymphoid potential. Animal studies showed that regular treadmill running exerts direct effects on the microenvironment of the bone marrow niche. Pro-inflammatory signals were down regulated and phenotype and function of HPCs were improved (15). Finally, not just younger, but older active individuals (>65 years) benefit from exercising by an increasing T cell proliferation, NK cell cytotoxicity, and neutrophil phagocytic activity (65) (Figure 3).

Some remarkable clinical benefits of exercise, such as a decreased risk of infection and an improved immunological response to vaccination, have been also proved in older adults (25, 67). Accordingly, active elderly women reveal a higher induction of antibodies against Flu B in their 18-month second vaccination follow-up (69). Similarly, regular exercising of moderate intensity for 25–30 minutes at three days per week for a period of ten months induced a significant increase in antibody titer to influenza vaccination (25).
Aktueller Erkenntnisstand in der Sportimmunologie

Regulatory Effects of Exercise on Inflammatory Status

The prevalence of obesity and chronic diseases rises worldwide. There is a huge body of evidence that a chronic low-grade inflammation is etiologically linked to the pathogenesis of many of these chronic diseases such as type 2 diabetes, specific types of cancer, dementia, cardiovascular diseases (19, 31, 65). This state is characterized by a 2- to 3-fold elevation in plasma concentrations of several inflammatory cytokines (IL-1beta, TNF-alpha) and acute phase proteins like CRP. During the last 2 decades many additional mediators of inflammation have been proposed (e.g. Adiponectin, leptin, resistin, visfatin, specific microRNA, autoantibodies, inflammasome & DAMPS) raising the question about the origin of these inflammatory signal molecules. Overweight and obesity are independent risk factors in developing a series of non-communicable diseases and there is growing evidence that malfunctioning adipocytes, especially those constituting to visceral fat mass, are the inducers of a chronic inflammatory state. Since adipose tissue (AT) is composed of various cell types, such as fibroblasts, endothelial cells, and leukocytes, the relative amounts of inflammatory so-called adipokines produced by adipocytes themselves remain often unknown.

In lean AT, the number of immune cells is low and mainly restricted to anti-inflammatory cell types such as M2 macrophages and Tregs. During the progression of obesity, there is an excessive fat energy storage which leads to an enlargement of fully differentiated adipocytes and pathologic expansion of AT. These processes are accompanied by both changes in the phenotype of various AT resident immune cells and the invasion of additionally recruited immune cells from the blood. This metabolically orchestrated inflammation, also known as metaflammation, is characterized by a spillover of pro-inflammatory mediators from AT resident leukocytes to the blood (23, 31).

For instance, the progressive infiltration of M1 macrophages enhances the secretion rate of pro-inflammatory mediators resulting in an increase of these cytokines in plasma. Besides, NK cells and T cells increase in AT during obesity while the number of inflammation regulating cells like M2 macrophages and Tregs decreases (23, 31, 45).

Epidemiological cross-sectional studies demonstrate both an association between physical inactivity and low-grade systemic inflammation and that self-reported physical activity and physical fitness is negatively associated with the presence of inflammatory biomarkers (47). Moreover, some longitudinal studies demonstrate that exercise training interventions may suppress systemic low-grade inflammation (38), most human RCTs are inconclusive or failed to show a decrease in the inflammatory level due to methodological limitations though (10). Multiple pathways by which exercise acts as an anti-inflammatory intervention have been discussed. First of all, regular exercise induces a negative energy balance and alters body composition in terms of a reduced fat mass (19). Second, exercise reduces the pro-inflammatory state of AT by means of a reduced recruitment of leukocytes to AT (19) and inducing a switch to an anti-inflammatory phenotype of many immune cells (macrophages, monocytes, T-cells) (19, 45).

Third, upon work skeletal muscle produces many cytokines and other peptides with anti-inflammatory properties. In this regard, IL-6 and other myokines act in a hormone-like fashion and execute endocrine effects in other organs and tissues than the skeletal muscle. IL-6 induces the production of anti-inflammatory cytokines such as IL-1ra and IL-10 (55), soluble TNF-alpha receptors (29), and inhibits the endotoxin-induced TNF-alpha production (55). Fourth, exercise stimulates the release of stress hormones with anti-inflammatory properties such as cortisol and adrenaline by activating both the SNS and the HPA axes (19). Last, recent evidence shows a close relationship between the integrity of gut microbiota and immune function and overall health. Since there is emerging research data which indicates that exercise and physical fitness ameliorates the gut microbiota (in terms of a higher diversity and balanced composition of microorganisms, see 4) limited evidence indicates that exercise-induced alterations of the gut microbiota account for a better immune function (4, 36). Taken together, it is believed that the direct and indirect anti-inflammatory effects of each bout of exercise, which accumulate over time, mediate a high proportion of both the preventive and therapeutic efficacy of exercise.

Figure 3: Cellular and molecular features of immunosenescence and the effects of exercise on the immune system of aged individuals (β2-ARs=beta-2 adrenergic receptors).
Most of the available data indicate that exercise is a powerful behavioral intervention to improve immune and overall health outcomes in subjects at all ages. Given that the global burden of chronic diseases is steadily rising, the development and promotion of strategies to a long-term increase of daily physical activity levels at the societal level are necessary. In parallel, scientists are tasked with deepening their understanding of molecular mechanisms in exercise immunology, optimize exercise regimes, and to investigate the effects of combined lifestyle modifications including exercise, nutritional, pharmacological, and other behavioral strategies. Thus, there is a need of well-controlled, randomized interventional training studies using different exercise regimens with multiple end points in order to develop guidelines for immune-enhancing training strategies. In parallel, exercise immunology should make use of innovative biomedical techniques to connect applied research questions more efficiently with basic science being the next step to translational science.

Conflict of Interest
The authors have no conflict of interest.
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