Evolutionary Aspects of Human Exercise – Born to Run Purposefully

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Abstract

This article is intended to raise awareness of the adaptive value of endurance exercise (particularly running) in the evolutionary history of humans, and the implications of the genetic disposition to exercise for the aging populations of modern technology-driven societies. The genome of Homo sapiens has evolved to support the svelte phenotype of an endurance runner, setting him/her apart from all other primates. The cellular and molecular mechanisms underlying the competitive advantages conferred by exercise capacity in youth can also provide a survival benefit beyond the reproductive period. These mechanisms include up-regulation of genes encoding proteins involved in protecting cells against oxidative stress, disposing of damaged proteins and organelles, and enhancing bioenergetics. Particularly fascinating are the signaling mechanisms by which endurance running changes the structure and functional capabilities of the brain and, conversely, the mechanisms by which the brain integrates metabolic, cardiovascular and behavioral responses to exercise. As an emerging example, I highlight the roles of brain-derived neurotrophic factor (BDNF) as a mediator of the effects of exercise on the brain, and BDNF’s critical role in regulating metabolic and cardiovascular responses to endurance running. A better understanding of such healthspan-extending actions of endurance exercise may lead to new approaches for improving quality of life as we advance in the coming decades and centuries.

Introduction

The take home message of this article can be appreciated from the following few sentences in which I comment on daily life in modern societies. Ah, this is the life – no need to run, nary even to walk. We curse the ‘inconvenience’ of a broken elevator and the ‘annoying’ bicycle rider who slows our drive home. We work with our fingertips and relax by watching television, while consuming large amounts of omnipresent tasty morsels. As a consequence, our body and brain experience a chronic positive energy balance. Unless we are motivated to rectify this dangerous lifestyle, obesity and/or insulin resistance occur, hastening the development of age-related diseases including diabetes, cardiovascular disease, cancers and neurodegenerative disorders. Because this scenario has become hauntingly common, our society must endure the increasing burden on the health care system and the workforce as the time window of productivity of persons in positive energy balance (PPEBs) shrinks.
Recently, several investigators have placed the ongoing epidemic of obesity and associated diseases in the context of the evolutionary history of Homo sapiens (Booth and Lees, 2007; O’Keefe et al., 2010). The idea is that a sedentary lifestyle betrays the evolutionary history encoded in our genes, a genetic code for periodic endurance running and intermittent feeding. Here I focus on the notion that for optimal fitness our genetic code demands challenges to the cells within which it resides, challenges that stimulate the expression of ‘survival genes’ that encode for proteins that enhance the ability of the cells to withstand oxidative and metabolic stress. The latter ‘use it or lose it’ hypothesis falls within the broader concept of hormesis which we have defined previously as “an evolutionarily conserved process in which a low dose of a stressful stimulus activates an adaptive response that increases the resistance of the cell or organism to a moderate to severe level of stress” (Calabrese et al., 2007). Though adaptive stress responses occur in cells throughout the body, I will comment on the adaptive responses of muscle and nerve cells to the stress of physical exercise.

**Homo sapiens are unique among primates in their capability for endurance running**

Before delving into an exploration of the molecular events occurring within muscle and brain cells during and after exercise that may protect against aging, I will briefly review some of the emerging information concerning the evolution of the remarkable capacity of humans for endurance exercise. An analysis of the structural and physiological underpinnings of endurance running ability among mammals revealed several features of the genus Homo that evolved within the past 2 million years that endow humans with superior sustained long distance running capability (Bramble and Lieberman, 2004). Humans are the only primates capable of sustained endurance running. One structural feature of humans that is believed to facilitate energetic efficiency during endurance running is legs with long spring-like tendons (e.g., the Achilles tendon) that attach short muscle fibers to leg and foot bones (Figure 1). The plantar arch of the foot may also be an adaptation for endurance running in humans which functions as a spring that returns up to 20% of the energy generated during the weight-loading phase of the running stride. Additional adaptations for endurance running in humans (compared to lower primates and quadruped mammals) include: long legs and stride length; relatively small feet with short toes (compared to other nonhuman primates) (Rolian et al., 2009); slow twitch muscle cells; large gluteus maximus muscle; structural modifications of the hips and shoulders that generate counter-balancing forces to enable smooth transitions between strides; sweat glands, reduced body hair and an elongated body form for heat dissipation (Marino, 2008); and mouth breathing.

A range of genetic and environmental factors may determine endurance running capabilities among individuals. A recent example is a study reporting that a polymorphism in the gene encoding α-actinin 3 was present in significantly higher frequencies in both elite endurance athletes and centenarians compared to elite power athletes and non-centenarians (Fiuza-Luces et al., 2011). Another example is the level of testosterone experienced by the developing fetus. A low ratio of the length of the second digit (index finger) to the fourth digit (ring finger), which has long been known to be associated with relatively high levels of fetal testosterone, was recently reported to be associated with superior endurance running (Manning et al., 2007).

**A brain – endurance connection?**

Interestingly, the increased size of the human brain relative to other primates, and its resultant cognitive capabilities, may have played an important role in the endurance running phenotype. Consistent with the latter possibility, a study in which brain size and maximum...
metabolic rate (MMR; a proxy for exercise capacity) were measured in a range of mammals revealed a positive correlation between brain size and MMR across a wide range of species (Raichlen and Gordon, 2011). Why might this be so? One reason is that distance running in our human ancestors was purposeful and required complex cognitive processes. The retention and recall of the details (topography, potential food sources, water sources, etc.) of large areas of land was likely required to maximize the acquisition of resources that were ‘spread thin’ in a timely manner. Individuals who possessed superior cognitive processing ability and endurance running capacity would be expected to have an advantage over those with lesser mental and endurance capabilities. The nervous system controls all aspects of body movements over a large time scale from milliseconds to hours, days, months and years. It is the planning of locomotion/behaviors that is mediated by nerve cell circuits housed in the evolutionarily expanded regions of the cerebral cortex in humans.

Not only does exercise strengthen muscle cells, it also strengthens brain cells (Figure 2). Indeed, it is now well-established that endurance exercise can stimulate the growth of brain cells and can improve cognitive function (see articles by Stranahan et al. and van Praag et al. in this issue). Running stimulates the production of neurotrophic factors, most notably brain-derived neurotrophic factor (BDNF), which promotes the growth of dendrites, the strengthening of synapses and even the production of new nerve cells from stem cells in some brain regions (van Praag et al., 1999; Duman, 2004; Stranahan et al., 2009; Kobilo et al., 2011a, 2011b). The latter scenario makes sense from the standpoint of optimizing the ability of the animal or human to acquire, store and protect resources. For example, during a 1 hour run many different potential sources of food or materials that could be used for shelter could be encountered, and up-regulation of the expression of BDNF at synapses activated during the encounter with the resource would promote strengthening of those synapses and long-term retention of the memory of the resource and its location. The cellular and molecular mechanisms by which BDNF expression is increased in response to exercise, and the signal transduction pathways activated by BDNF and some of the genes induced by BDNF that strengthen neurons are described below and have been reviewed in more detail recently (Lu et al., 2008; Greenberg et al., 2009; Cohen-Coray et al., 2010).

Interestingly, emerging evidence indicates that BDNF plays prominent roles in the regulation of energy metabolism via actions at multiple sites in the nervous system (Noble et al., 2011). For example,

**Exercise retards age-related decrements in muscle strength and endurance via hormesis**

While it is well-known that muscles atrophy if not used, and that regular exercise will increase muscle strength and endurance, the underlying molecular mechanisms have only recently emerged. As reviewed by Jackson and McArdle (2011), in young animals exercise induces superoxide and nitric oxide in the muscle cells, and the latter reactive oxygen species (ROS) stimulate several transcription factors (AP-1, HSF-1 and NF-κB) which, in turn induce the expression of antioxidant enzymes and protein chaperones. The increased levels of the latter cytoprotective proteins extend well beyond the exercise period such that the stress during the exercise results in a long-lasting resilience of the muscle cells. The importance of ROS-mediated signaling in the beneficial effects of exercise on muscle cells is supported by data showing that treatment of human subjects with antioxidants (vitamins E and C) abolish the ability of exercise to activate PGC-1α and enhance insulin sensitivity in muscle cells (Ristow et al., 2009). The oxidative stress that occurs in muscle cells during exercise can also trigger mitochondrial biogenesis, a process mediated by PGC-1α, a master regulator of the growth and division of mitochondria (Wright et al., 2007). Endurance exercise engages cytoprotective responses in all of the major subcellular organelles. For example, the endoplasmic reticulum (ER) responds to exercise by engaging a sequence of
events called the unfolded protein response (UPR) involving the up-regulation and/or activation of multiple ER proteins (e.g., glucose-regulated protein 78 and calreticulin) which ensure that newly synthesized proteins are properly folded. The latter adaptive response to exercise is mediated by PGC-1α and coactivation of ATF6α (Wu et al., 2011). In addition to these cell-autonomous signaling events that occur in muscle cells in response to exercise, those cells also produce factors that are released and promote the growth and resilience of adjacent muscle cells; such factors include insulin-like growth factor 1 (Harridge, 2003) and vascular endothelial cell growth factor (Olfert et al., 2009).

The ability of muscle cells to respond adaptively to exercise is compromised during the aging process. Thus, it was found that levels of protein chaperones and antioxidant enzymes were not increased in skeletal muscle in response to exercise in very old (28 month-old) rats (Vasilaki et al., 2002, 2006). Interestingly, muscle cells in very old animals exhibit elevated constitutive activation of the redox-sensitive transcription factor NF-κB, and some antioxidant enzymes and pro-inflammatory cytokines (Jackson and McArdle, 2011). Exercise also fails to induce mitochondrial biogenesis in muscles from very old animals. The combination of increased basal oxidative stress/damage and an inflammatory state may render the old muscle cells incapable of responding adaptively to exercise. Because regular exercise during midlife and continuing into old age can delay the onset of age-related muscle atrophy (Rogers and Evans, 1993; Peterson et al., 2010), it is likely that the ROS-mediated signaling pathways that mediate the beneficial effects of exercise on muscles are preserved as a result of the midlife exercise. However, numerous studies of human subjects have shown that endurance exercise benefits muscles and the cardiovascular system even when initiated in elderly subjects who were relatively sedentary in midlife (Hollmann et al., 2007; Williams and Stewart, 2009).

**Neurotrophic factors mediate anti-aging effects of endurance exercise on the brain**

It has become clear that there is a widespread activation of signaling pathways involved in adaptive stress responses in cells of many different organ systems that occurs during and after endurance exercise (Radak et al., 2008). Nerve cells in the brain are remarkably responsive to exercise. Studies of animal models and of human subjects support the hypothesis that exercise induces the expression of neurotrophic factors which, in turn, promote structural and functional plasticity of neurons and their resistance to injury and disease (Gomez-Pinilla, 2008; Mattson and Wan, 2008; Zoladz and Pilc, 2010). BDNF has been the most intensively studied exercise-induced neurotrophic factor. Recent studies of human subjects have shown that age-related decrements in cognitive performance are associated with reduced circulating BDNF levels (Erickson et al, 2010) and that aerobic exercise training can elevate BDNF levels, increase the size of the hippocampus and improve memory in elderly subjects (Erickson et al., 2011). Two mechanisms by which BDNF expression is induced by running are synaptic activity-mediated activation of the transcription factor CREB (cyclic AMP response element-binding protein (CREB; Conti et al., 2002; Chen and Russo-Neustadt, 2009) and energetic stress-mediated activation of the transcription factor NF-κB (Marini et al., 2004; Kairisalo et al., 2009). A major signaling pathway upstream of CREB involves activation of synaptic glutamate receptors which results in calcium influx and activation of a calcium/calmodulin-dependent protein kinase (Hu et al., 1999). CREB can be considered a stress sensor because it not only induces BDNF expression, but also expression of multiple genes that encode proteins that protect neurons against oxidative stress, including the DNA repair enzyme APE1 (Yang et al., 2010) and Bcl-2 (Meller et al., 2005).
Multiple intercellular signaling pathways have evolved that coordinate the adaptive responses of the brain to exercise. For example, in addition to BDNF, exercise also induces the expression of fibroblast growth factor 2 (FGF2) and VEGF in the brain (Gomez-Pinilla and Kesslak, 1998; Fabel et al., 2003). FGF2 can protect neurons against oxidative, metabolic and excitotoxic injury (Cheng and Mattson, 1991; Mattson et al., 1995) and also promotes the growth of astrocytes, a type of glial cell that provides metabolic support to neurons (Petroski et al., 1991). FGF2 also promotes the proliferation of neural progenitor cells and may, together with BDNF, play a key role in exercise-induced neurogenesis (Bull and Bartlett, 2005). In addition, it has been shown that running increases the expression of VEGF and enhances angiogenesis in the hippocampus of rodents (Fabel et al., 2003; Kerr et al., 2010), which would be expected to increase the supply of nutrients to the brain cells. During endurance exercise, energy utilization by skeletal and cardiac muscle are greatly increased, but it is critical that the energy supply to the brain be maintained because nerve cells require a constant supply of glucose and will quickly become dysfunctional if glucose levels are reduced as occurs in hypoglycemia and ischemic conditions. Endurance training can, therefore, enhance the availability of glucose to neurons.

In order to begin to understand the scope and integration of adaptive changes that occur in brain cells in response to endurance exercise in the context of aging, we performed a large-scale gene array analysis of brain tissue samples from old mice that were life-long runners compared to age-matched non-runner control mice (Stranahan et al., 2010). Prior to euthanizing the mice, they were trained in a water maze to stimulate activity in nerve cell circuits involved in learning and memory processes. Compared to the more sedentary mice, cells in the runners brains exhibited greater activation of genes involved in mitochondrial function and synaptic plasticity, and lower levels of activation of genes involved in oxidative stress and lipid metabolism. One interesting pathway modified by running was the pathway activated by the Wnt protein (Stranahan et al., 2010). It was recently reported that mice lacking the Wnt receptor frizzled-related protein exhibit reduced running exercise performance (Lories et al., 2009), although future studies will be required to establish whether Wnt signaling in the brain (or other tissues) is critical for endurance running.

BDNF has emerged as a pivotal regulator of energy metabolism and a mediator of many different adaptive responses of the brain and body to endurance exercise (see Pedersen et al., 2009; Noble et al., 2011 for review). Mice with reduced levels of mice exhibit hyperphagia, and develop obesity and diabetes (Kernie et al., 2000; Duan et al., 2003). Increased expression of BDNF may mediate, in part, the enhanced insulin sensitivity that occurs in response to endurance exercise. The latter possibility is consistent with studies showing that infusion of BDNF into the brain can reduce plasma glucose levels and ameliorate diabetes in mice (Tonra et al., 1999; Nakagawa et al., 2000). A recent study provided evidence that hypothalamic BDNF signaling can induce the generation of brown fat cells within white adipose deposits by a mechanism involving modulation of the autonomic innervation of the white fat (Cao et al., 2011). Cardiovascular adaptations to endurance running may also involve BDNF signaling in the nervous system. We found that intermittent food deprivation (which is known to extend lifespan in rodents) results in reductions in resting heart rate and blood pressure, and improved cardiovascular adaptation to stress in rats (Wan et al., 2003). These effects of intermittent food deprivation are similar to those that occur in response to endurance running, and involve increased parasympathetic tone and increased heart rate variability (Mager et al., 2006). More recently we provided evidence that BDNF signaling in the brainstem enhances parasympathetic tone and reduces heart rate (Griffioen et al., 2010), suggesting a role for BDNF as a mediator of the effects of exercise on the heart. Thus, the emerging picture of the brain – body interface and endurance includes a prominent role for BDNF as an integrator of many of the major...
systems that regulate neural, endocrine and cardiovascular adaptations to and for endurance exercise.

Conclusions

In the not too distant past pizzas were not delivered, there were no drive-through fast-food restaurants, and no automobiles. Quite the contrary, during our evolutionary history, there was selection for genes that ‘posed the questions’: Why sit when you can walk and why walk when you can run? From the perspective of optimal health, we have over-engineered our lives, not only from the perspective of technologies that reduce the necessity of exercise, but also in more subtle ways. One example is the running shoe industry, which has promulgated the myth that cushion and toe compartment stability reduce injuries. In fact, it has recently been directly determined that barefoot runners experience less strain on their legs than do shod runners because the barefoot runners are ‘toe strikers’ and the shod runners are ‘heel strikers’ (Lieberman et al., 2010). Humans exhibit numerous adaptations that enable endurance running that include features of musculoskeletal anatomy and physiology, cardiovascular regulation, and metabolic efficiency. A fascinating aspect of the evolution of endurance running capability is the apparent co-evolution of signaling pathways that regulate neuroplasticity and peripheral adaptations to exercise. I have highlighted BDNF signaling as one such pathway, and it will be of interest to delve deeply into the evolutionary origins of BDNF and its receptor trkB to determine if and how it influenced selection for endurance running phenotypes. Other signaling mechanisms that integrate the response of multiple organ systems to endurance running will undoubtedly be discovered. Modern methods for genome-wide DNA sequencing, and analysis of epigenetic modifications (e.g., methylation and acetylation), will enable discovery of the inherited and acquired molecular factors involved in endurance capacity. In the more distant future, knowledge of the various pathways that promote endurance phenotypes may lead to novel interventions that promote optimal health.

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References


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HIGHLIGHTS

- Humans are anatomically and metabolically suited for running long distances
- Endurance running improves health by enabling cells to cope with stress
- The human brain evolved to process complex information related to resource acquisition
- Brain-derived neurotrophic factor mediates multiple effects of exercise on the brain
Figure 1.
Adaptions of humans for endurance running.
Figure 2.
Brain-derived neurotrophic factor (BDNF) is an integrator of adaptive responses of the brain and body to endurance running.