

Impact of Physical Exercise on Gut Microbiome, Inflammation, and the Pathobiology of Metabolic Disorders

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■ Abstract

BACKGROUND: The gastrointestinal tract (GIT) harbors a complex and diverse microbial composition that outnumbers our own body cells and their gene contents. These microbes play a significant role in host metabolism and energy homeostasis. Emerging evidence suggests that the GIT microbiome significantly contributes to host health and that impairments in the microbiome may cause the development of metabolic diseases. The microbiome architecture is shaped by several genetic and environmental factors, including nutrition and physical activity. Physical exercise has preventive or therapeutic effects in respiratory, cardiovascular, neuroendocrine, and muscular diseases. Yet, we still have little information of the beneficial effects of physical exercise on GIT health and microbial composition. Furthermore, we are not aware whether exercise-derived benefits on microbiome diversity can beneficially influence other tissues and body organs. **OBJECTIVES:** The aim of this article is to review the available literature on exercise-induced microbiome

changes and to explain how these changes may induce inflammatory, immune, and oxidative responses that may contribute to the improvement of metabolic disorders. **METHODS:** A systemic and comprehensive search of the relevant literature using MEDLINE and Google Scholar databases was conducted during fall 2018 and spring 2019. The search identified sixty-two research and review articles that discussed exercise-induced microbiome changes. **RESULTS:** The review of the relevant literature suggests that exercise-induced microbial changes affect the host's immune pathways and improve energy homeostasis. Microbes release certain neuroendocrine and immune-modulatory factors that may lower inflammatory and oxidative stress and relieve patients suffering from metabolic disorders. **CONCLUSIONS:** Exercise-induced changes in microbial diversity are able to improve tissue metabolism, cardiorespiratory fitness, and insulin resistance.

Keywords: diabetes · exercise · microbiome · microbe · inflammation · immune response · obesity · oxidative stress

1. Introduction

During the last century the mortality index indicated a gradual shift from infectious diseases towards non-communicable diseases (NCDs) [1]. Simultaneously, the importance of healthy lifestyle has been emphasized. Besides genetics, diet and lifestyle are the two most important factors associated with the risk of obesity-

related metabolic disorders (ORMDs). Consumption of a calorie-rich diet and sedentary lifestyle have contributed to the rising incidence of ORMDs in industrialized countries. In particular, cardiovascular diseases, diabetes, and obesity have been associated with sedentary lifestyle.

Research on the gastrointestinal tract (GIT) microbiome has increased our understanding of the bi-directional communication between host

and microbiome which is carried out through neuroendocrine signals and genetic transfer, and increased knowledge on “microbiome perturbations” in the pathogenesis of ORMDs [2]. Cumulative evidence suggests that the GIT microbiome is an essential mediator between different factors such as genetics, diet, exercise, and environment and the pathophysiology of the ORMDs [3]. Genetics, dietary habits, and host environment may influence the microbiome and eventually affect host metabolism. While this interaction has been studied extensively in diet-induced microbiome perturbations and host physiological and pathological processes [4], little attention has been given to other factors such as physical exercise.

GIT microbiome research has emerged as a potential exploratory field with special focus on ORMD pathogenesis. Microbiome dysbiosis (derangements in microbial ecology) in ORMDs has been reported [5]. Because of the presence of highly diverse microbial populations, the GIT represents the primary site of immunological activity. Lifestyle and physical activity may act as important factors to reshape gut microbiome and immunological responses [6], and provide the opportunity to influence disease development by therapeutic measures.

Therefore, in this article, we collected information on the impact of exercise on GIT microbiome and discussed how this information may relate to major NCDs, such as obesity and diabetes. We have previously reported changes in microbiome diversity in a type 1 diabetes rat model [7], and we have outlined the changes in microbiome architecture in diabetes and obesity [8]. In this review, we also analyzed exercise-associated changes in the microbiome that are able to modify body physiology and homeostasis.

2. Methods

The review of the literature was intended to identify taxonomic and community changes in the GIT microbiome in response to physical exercise. We explored exercise-mediated alterations in gut microbiome and metabolites that could be related to systemic inflammation, oxidative stress, and impaired immune responses. A systematic and comprehensive search of MEDLINE and Google Scholar was conducted between fall 2018 and spring 2019 to find relevant research articles published from 2000 to 2019. The following keywords were used in different combinations to retrieve all relevant research articles published in the English language:

Abbreviations:

AMPK	5' AMP-activated protein kinase
BRFSS	Behavioral Risk Factor Surveillance System
BWG	body weight gain
CFU	coliform unit
GABA	γ -aminobutyric acid
GIT	gastrointestinal tract
GLP-1	glucagon-like peptide-1
GPR	G protein receptor
GPR	G-protein-coupled receptor
HFD	high-fat diet
HPA	hypothalamus-pituitary-adrenal
IL	interleukin
LDL	low-density lipoprotein
LPS	lipopolysaccharides
NCD	non-communicable disease
NF- κ B	nuclear factor kappa light-chain enhancer of activated B cells
ORMD	obesity-related metabolic disorder
PYY	peptide YY
SCFA	short-chain fatty acid
SGLT1	sodium-glucose cotransporter 1
Th cell	T helper cell
TLR	toll-like receptor
Treg cell	T regulatory cell

- Exercise
- Physical activity
- Microbiome
- Metabolic disorders
- Inflammation
- Immunity
- Obesity
- Diabetes

References and bibliographic lists in published and review articles were also reviewed to identify relevant studies that may have been overlooked during database searches. All original research articles that presented data on intestinal microbiome, exercise, and/or metabolic disorders in humans and animals were included in the study. The literature search was not restricted to:

- Pattern, intensity, or duration of exercise
- Participants' gender or age
- Sampling pattern or
- Study design

We retrieved 51 original articles and 10 review papers published during 2008-2019 (**Figure 1**). The research articles were divided into the following two categories:

1. Articles that studied exercise-induced microbiome diversity.

2. Generic articles that linked exercise-induced microbiome changes with ORMDs.

For each study, we retrieved data on exercise pattern, microbiome diversity, metabolic responses, and immune-inflammatory markers. The primary outcome of this effort was in the form of a data base on the following aspects:

- GIT microbiome taxonomy and metabolic profile
- Immune-inflammatory responses
- Oxidative stress

A few studies discussed the molecular mechanisms involved in exercise-mediated alterations of the microbiome and systemic inflammation and immunity [9, 10], while the majority of studies simply described phylogenetic changes observed in response to exercise. To pursue the objective of this review, namely to find out more about the relation between microbiome changes and exercise with respect to metabolic diseases, we further explored review articles and mechanistic studies that related systemic inflammation and immunity to microbiome and exercise.

3. The microbiome as a virtual body organ

Most body surfaces of mammals harbor a robust microbial colonization that is established to protect the body against potential pathogens; it is collectively called the microbiome. The microbiome is composed of archaea, bacteria, eukaryotes, viruses, and their genetic material. Both cell number and genetic material of the microbiome outnumber our own body cells and their genetic materials [11]. The host's genetics and several habitual and environmental factors act together in shaping host microbiome (**Figure 2**). Beyond genetics, a number of habitual and environmental factors contribute to shaping the host microbiome, including mode of delivery at birth, breastfeeding as opposed to for-

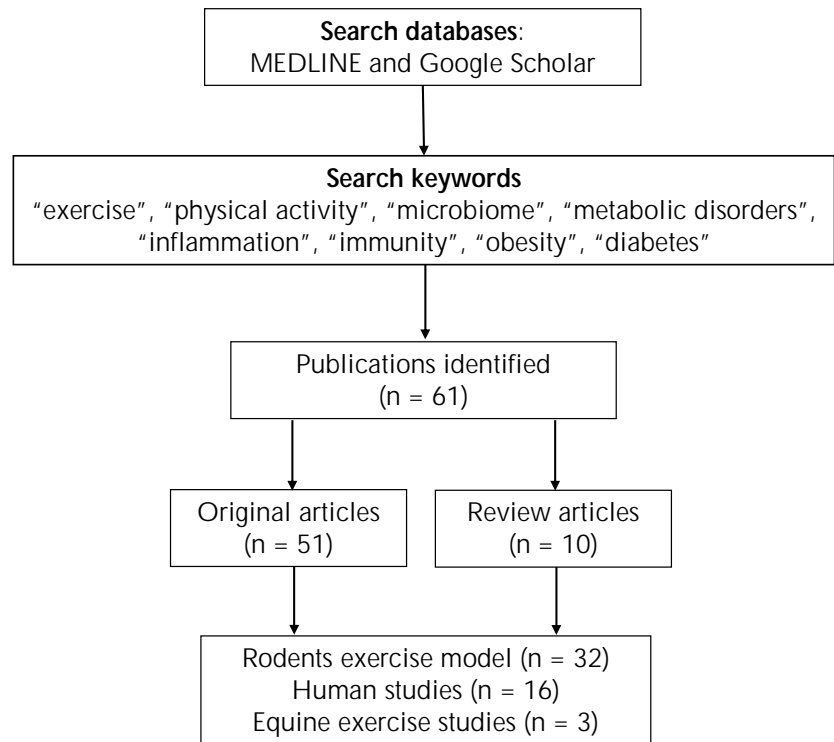


Figure 1. The figure illustrates the search method used to collect research articles from Medline and Google Scholar databases.

mula feeding, adult dietary habits, hygiene, physical activity, antibiotics or xenobiotics exposure, and geographical location [12-14].

The microbiome has essential structural, protective, metabolic, and endocrine functions that include food digestion, pathogen displacement, and nutrient synthesis [15, 16]. The GIT is the primary site of interaction between our body's largest component of the immune system and microbial colonization. The commensal microbes not only tune and train this local immunity, they also impact systemic immunity. Microbes interact with the nervous and endocrine system through glucagon-like peptide 1 (GLP-1) and peptide YY (PYY) that are produced in enteroendocrine L-cells, even though the exact mechanisms involved in this communication are not fully understood [17].

The endocrine activities of the microbiome are very heterogeneous. Frequently, the metabolic products of the microbiome, such as short-chain fatty acids (SCFAs), histamine, monoamines, and γ -aminobutyric acid (GABA), act as second messenger molecules or neurotransmitters [18-20]. These molecules mediate gut-brain and endocrine communications and contribute to the micro-

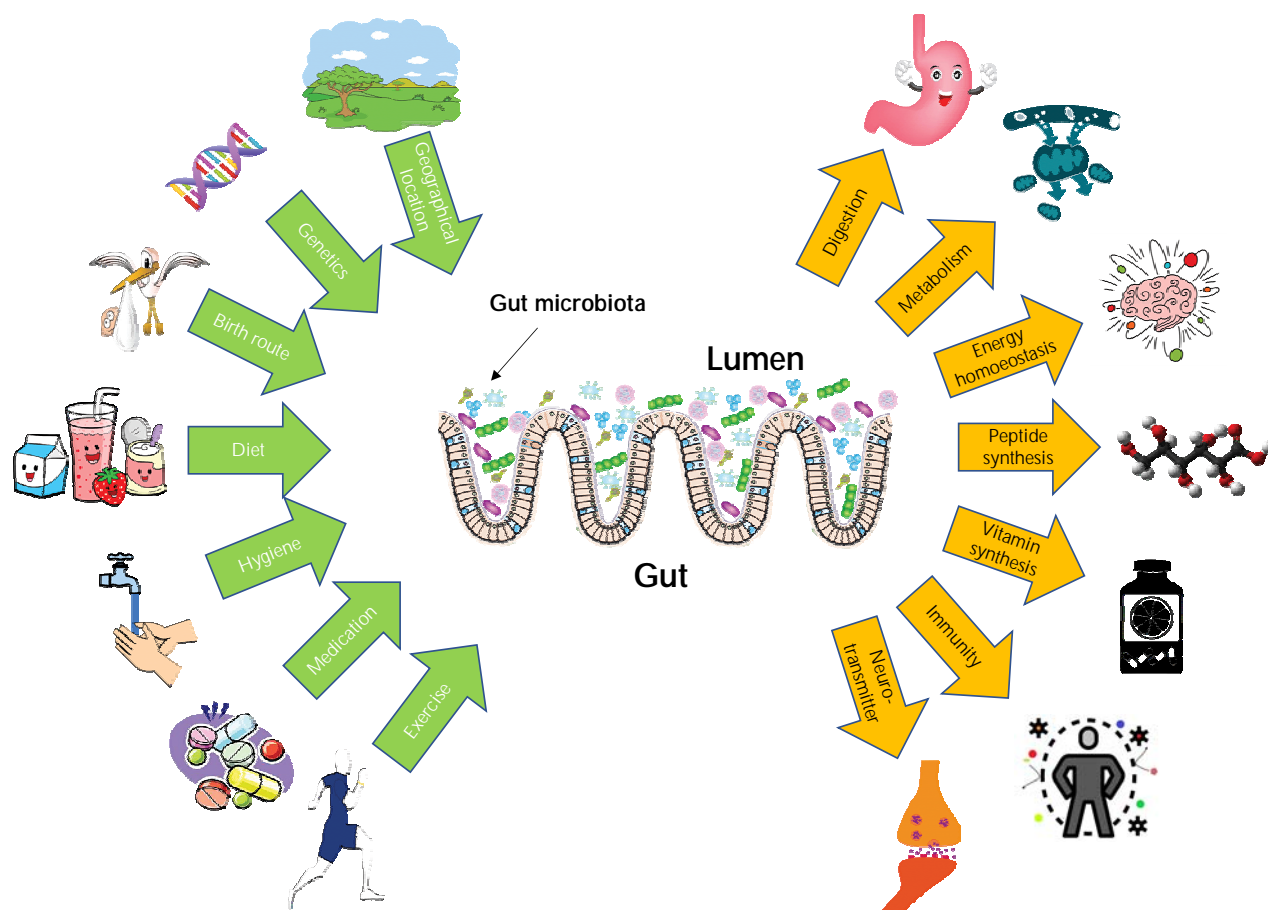


Figure 2. Factors that influence the human microbiome architecture and different functions performed by our microbiome. Genetics and mode of delivery at birth are the first predisposing factors impacting the gastrointestinal (GIT) microbiome. They work together with breast-feeding, hygiene, exposure to medication (antibiotics), dietary habits, and physical exercise to shape GIT microbiome. The diagram also sketches the different functions of the GIT microbiome. The microbiome helps to digest food and control host body energy homeostasis. Microbes secrete several useful peptides and are part of our body's largest immune system.

biome's systemic functions. Gnotobiotic rodent models and probiotic supplementation studies have highlighted the significance of the microbiome. The poor microbial composition in gnotobiotic rats is explicated by averted hypothalamic-pituitary-adrenal (HPA) responses to psychological or environmental stressors [21, 22]. This neuroendocrine containment within the microbiome influences behavior and learning aptitude through biosynthesis of microbial neurochemicals either directly or by bi-directional transfer of genetic information between host and microbiome [23].

SCFAs are the main metabolic products of microbiome fermentation. These molecules act as signaling ligands for G-protein-coupled receptors (GPRs) on enterocytes for chemosensation, appetite regulation, and peripheral nutrient disposal

[24]. Acetate, butyrate, and propionate are the main SCFAs that are produced by bacterial fermentation. Acetate and propionate enhance the expression of GPR-43 in immune cells to regulate immune-inflammatory responses [25]. Furthermore, these metabolites improve intestinal barrier function and display potent immunoregulatory properties through T cell proliferation and cytokine production [26].

4. Microbial dysbiosis and aberrant energy homeostasis: the prodrome of metabolic syndromes

Obesity is the consequence of genetic predisposition, chronic energy imbalance, and lack of

proper physical activity. Extensive diet restriction as well as lifestyle, surgical, and pharmacological interventions have been implemented to treat obesity. These efforts are targeted at losing weight, but they have demonstrated insufficient, unsafe, and inconsistent results in most of the cases [27-29]. In recent years, significant progress has been made in the understanding of the GIT microbiome, suggesting a crucial intermediate role in energy homeostasis.

The basic functions of the gut microbiome are to digest dietary fibers and provide energy to the host. Changes in GIT microbiome composition alter digestive efficiency and consequently host metabolism [30]. Several studies have demonstrated substantial changes in microbiome composition in obese subjects compared with healthy controls, and these changes were associated with ORMDs. Palleja *et al.* observed that weight loss interventions are associated with beneficial changes in bacterial diversity and improvements in metabolic functions [31]. Furthermore, fecal microbiome transplant from lean donors recapitulates the functional metabolic phenotype and improves insulin sensitivity in the obese recipients [32].

Consumption of energy-rich food or antibiotics can induce microbiome perturbations and promote the incidence of metabolic syndromes. For example, prolonged antibiotic treatment, or antibiotic consumption during early life may induce microbial dysbiosis and promote obesity and diabetes [33, 34]. Hicks *et al.* observed a positive correlation between microbial dysbiosis, antibiotic consumption, and metabolic syndrome [35]. The authors performed a retrospective analysis using major data pools, including USA population data, American Community Survey, Area Resource File data, and the Behavioral Risk Factor Surveillance System (BRFSS) data obtained from Centers for Disease Control and Prevention. They found that the obese portion of the American population has the highest antibiotic prescription rate.

While these studies suggest an association between microbiome and host energy homeostasis, most of them lack a mechanistic explanation of the impact of microbiome changes on host energy cycle. It is still unknown whether these microbiome changes are cause or effect of obesity. We present the most discussed microbiome-energy homeostasis pathways below.

Recent evidence suggests bi-directional communication, through neuroendocrine signals, between gut microbiota and host cell energy homeostasis [36]. Several high-impact studies link SCFA production in gut fermentation to host energy ho-

meostasis [37, 38]. Furthermore, microbiome composition is a key regulator of non-digestible carbohydrate and fiber digestion capacity. Cani *et al.* have discussed the microbiome's capacity to digest fibers, the production of SCFAs, and host energy homeostasis in symbiotic relationship [39]. In brief, SCFAs promote mitochondrial function, lipolysis, neoglucogenesis, beige adipogenesis, and release of GLP-1 and PYY. Conversely, these intestinal hormones influence food intake by modulating the activity of brain feeding centers. Furthermore, as revealed by germ-free mice experiments, microbial fermentation products stimulate the production of leptin in adipocytes that promote hepatic lipogenesis and surplus energy storage [40].

5. Exercise as a modulator of gut microbiome

There is compelling evidence to suggest that long-term exercise has a positive effect on energy homeostasis and plays a preventive role against various ORMDs. The most important physiological adaptations that occur in response to exercise are mitochondrial biogenesis, muscular hypertrophy and angiogenesis, cardiovascular fitness, better organized energy supply chain, and activation of the HPA axis. Exercise-induced activation of the HPA axis improves energy balance and better regulates immune-inflammatory responses [20]. Exercise promotes the cholinergic anti-inflammatory pathway and reduces tumor necrosis factor α (TNF- α) release. Lowder *et al.* reported that continuous moderate exercise is able to prevent influenza infection in rats by enhancing the production of anti-inflammatory cytokines (IL-10 and IL-4) and by lowering the secretion of pro-inflammatory cytokines (TNF- α , IL-1, interferon- γ) [41].

Since 2008, more than 50 research articles have been published on the impact of exercise on gut microbiome diversity. Effects of exercise on GIT health, though not fully explained, have emerged as a key interest in cancer and metabolic disease research. Endurance exercise may modulate GIT immune-inflammatory and redox responses, GIT permeability, motility, and stool transit time and consistency [42-44]. Hoffmann *et al.* reported that acute and continuing exercise regimes of varying intensities could increase antioxidant capacity, lymphocyte turnover, and expression of anti-inflammatory cytokine in the intestinal wall [43]. In the initial stages of intense exercise, GIT hypoxia and hypoperfusion may increase gut perme-

ability, endotoxemia and oxidative stress [45, 46]. However, these changes in inflammation and immune response are transient and subside quickly. In particular, these changes are associated with the GIT microbiome in one way or another. It is also observed that specifically designed exercise therapies mitigate GIT inflammatory diseases by preferentially modulating microbiome diversity and metabolic profile [47, 48]. It is suggested that exercise increases gut motility, which may increase the shedding of loosely bound microbes in the GIT epithelium. This effect promotes the growth of other commensals that participate in the development of healthy mucosal immunity and provide benefits in gut tissue and beyond [49].

Physical exercise has positive effects on GIT microbiome biodiversity, as observed in several animal and human studies [10, 50, 51]. Several authors suggest that these changes in microbiome diversity are associated with cardiorespiratory fitness and GIT microbial metabolic profile [49, 52, 53]. People with better cardiorespiratory fitness have better microbial diversity and chemotaxis activity with decreased lipopolysaccharide (LPS) biosynthesis [52]. This improvement in microbial diversity and cardiorespiratory fitness in exercising subjects may be attributed to higher abundance of butyrate-producing bacteria by the *Clostridiales*, *Erysipelotrichaceae*, *Lachnospiraceae*, and *Roseburia* families.

Microbiome involvement in metabolic diseases is perhaps most vividly portrayed in gnotobiotic rodents and probiotic/prebiotic supplementation studies. Allen *et al.* reported that microbiome transplanted from exercising mice to gnotobiotic mice improved bacterial diversity and metabolic profile and decreased colon inflammation [54]. These changes in microbiome, metabolites, and inflammation markers are linked to gut permeability during scenarios of physiological stress. Therefore, targeting the intestinal microbiota may provide a novel strategy to prevent increases in intestinal permeability and thus to cope with physiological stress [55]. Exercise started in early life is more effective in correcting microbiome (increasing Bacteroidetes to Firmicutes ratio) and in developing stable lean body mass [56]. An increase in Firmicutes to Bacteroidetes ratio, characterized by microbiome enrichment with Firmicutes or depletion of Bacteroidetes, is recognized as an obesogenic trait and is often found in obese children [57]. Furthermore, an increase in butyrate-producing bacteria influences the metabolic pathways involved in fat accumulation and prevents obesity. Estaki *et al.* reported an increased production of

SCFAs and reduced biosynthesis of LPS by gram-negative bacteria in exercising subjects, suggesting a protective role of exercise in immune-inflammatory inhibition of ORMDs [52]. Similarly, Hsu *et al.* observed that probiotic *Bacteroides fragilis* supplementation improved exercise tolerance and swimming performance in gnotobiotic mice [58]. The improvement in exercise tolerance was also associated with improvements in liver, muscle, brown fat weight, and the antioxidant enzyme production system [58]. Although the initial response to intense exercise regimes constitutes an increase in gut barrier permeability, continuing exercise for longer duration may improve GIT microbial diversity which attenuates production of the cytokines involved in tissue regeneration [54].

Clarke *et al.* reported conformational changes in microbiome diversity in high-performance rugby player's stool, particularly, a decrease in Firmicutes to Bacteroidetes ratio [59], which has implications for the pathogenesis of obesity and diabetes [8]. Data from the American Gut Project revealed that adopting moderate exercise (from never to daily) increases species diversity among Firmicutes phylum and promotes healthier gut environment [60]. Habitual moderate to intense physical activity improves microbial diversity and immune functions and reduces inflammatory diseases [61]. Petersen *et al.* performed shotgun metagenomics and metatranscriptomic analysis of competitive cyclists, and reported changes in microbial diversity and an upregulation in carbohydrate metabolism and energy production [62]. Similarly, Welly *et al.* reported that exercise-induced microbiome changes in rats were associated with low insulin resistance, adipose tissue inflammation, and better exercise tolerance [63].

In most of the published studies, exercise-induced microbiome changes are investigated at the holistic level, while very few studies describe specific taxonomic associations between exercise and GIT microbiome. However, a few rodent exercise studies have associated weight loss with an increase in the Bacteroidetes/Firmicutes ratio [64, 65]. Furthermore, investigators observed a positive correlation between *Lactobacillus* and *Bifidobacterium spp.* and serum leptin levels in exercising rats. Although we assume that an increase in the Bacteroidetes/Firmicutes ratio is beneficial for metabolic health, as has been observed in several studies [57, 59, 64-66], there are also contradictory findings, suggesting exercise-induced health benefits in association with increased Firmicutes diversity and depletion in Bacteroidetes populations [60, 67]. These differences in observations may be

attributed to different exercise interventions, dietary patterns, or different genetic background of the host. The members of the Firmicutes phylum are most responsive to exercise-associated microbiome changes. However, it is hard to identify specific bacteria genera that create beneficial health responses or that lead to systemic inflammation.

Most of the studies reviewed here reported that the changes in microbiome diversity are associated with improvements in health status. Perhaps, most of these studies describe dissimilar taxonomical changes in microbiome in response to exercise, and generally present more global changes at phylum level (e.g. changes in Bacteroidetes/Firmicutes ratio) or in terms of alpha and beta diversity (species richness and diversity). These inconsistent taxonomical observations may be due to different experimental settings, such as differences in subjects' age, sex, disease state, supplementation regime, and exercise pattern. Although, similar exercising patterns may have different results in different ages [56] or diet groups [64], Allen *et al.* reported that different exercising patterns (voluntary wheel running vs. forced treadmill running) may also affect microbiome differently [68]. Therefore, the only conclusive observations in almost all of these studies are that exercise increases microbial species' richness and evenness (alpha diversity), and results in distinct clustering patterns (beta diversity), and that there is no single microbial taxon or group of bacteria that can be associated with such variation, as reported elsewhere [9, 10, 58].

Furthermore, most of the reviewed studies recapitulate enhanced production of SCFA, particularly butyrate [53, 64, 69], which are produced by microbial fermentation of fibers and non-digestible starch. This discussion endorses a significant role of physical exercise in microbial metabolism and energy cycles. Perhaps, further mechanistic experimentation is required to elucidate host energy homeostasis in the framework of mode and duration of training that imprints positive effects on microbial community and its functional containment. The future goals of research on exercise-induced microbiome changes should include molecular evaluation of immune-inflammatory pathways or neuroendocrine mediators in gut and metabolic tissues.

6. Metabolic disorders in the view of exercise-induced microbial diversity

Patients with metabolic disorders are characterized by excessive release of pro-inflammatory

cytokines from bone marrow-derived monocytes, NF- κ B, and chemoattractant protein 1 that disturb the blood-brain barrier and induce neuroinflammatory responses [70, 71]. Inactive lifestyle enhances fatigability and acts as mediator of the development of obesity, hypertension, dyslipidemia, and insulin resistance [72]. In a sedentary rat model of metabolic syndrome, Feng *et al.* observed that exercise rectified immune-inflammatory markers of obesity and immune-inflammatory responses [73]. These changes in ORMD markers were coequally associated with an exercise-induced improvement in microbiome diversity.

Recently, Dalton *et al.* reviewed the effects of exercise on the gut-brain axis and microbiome diversity [74]. The authors proposed that exercise-induced improvements in mood and psychological disorders are linked with changes in microbial diversity [75]. Increase in microbiome diversity in response to continuous aerobic exercise has been shown to stabilize GIT barrier function and lower symptoms of irritable bowel syndrome [76]. Probiotic supplementation and aerobic exercise may improve the diversity and abundance of genera from the Firmicutes phylum [76]. This may be explained as a neuroendocrine link between the positive effects of exercise on GIT and brain to prevent or treat psychological distress through the microbiome-gut-brain axis [74]. This neuroendocrine containment within the microbiome influences behavior or learning aptitude through biosynthesis of microbial neurochemicals either directly or through bi-directional transfer of genetic information between host and the microbiome [23].

Allen *et al.* observed that at baseline, the fecal microbiome, microbiome metabolic profile, and cardiorespiratory fitness level of obese subjects were different from those of lean subjects, and exercise training was positively correlated with these parameters [77]. However, when the rats were returned to a sedentary lifestyle, these changes also reverted to baseline values. Furthermore, exercise-induced microbiome changes reduced fatigue and depression, and improved exercise tolerance, motivation, and cardiorespiratory fitness. Similarly, Feng *et al.* showed that pre-operative exercise minimizes postoperative weight-gain, neuroinflammation, and cognitive decline and corrects the microbiome in rats [78]. Carbajo-Pescador *et al.* studied the effects of exercise training on the functionality of gut microbiota, intestinal barrier integrity, and hepatic steatosis in high-fat diet rats [9]. Exercise lowered body weight, mitigated metabolic syndrome and hepatic steatosis, and improved intestinal barrier function.

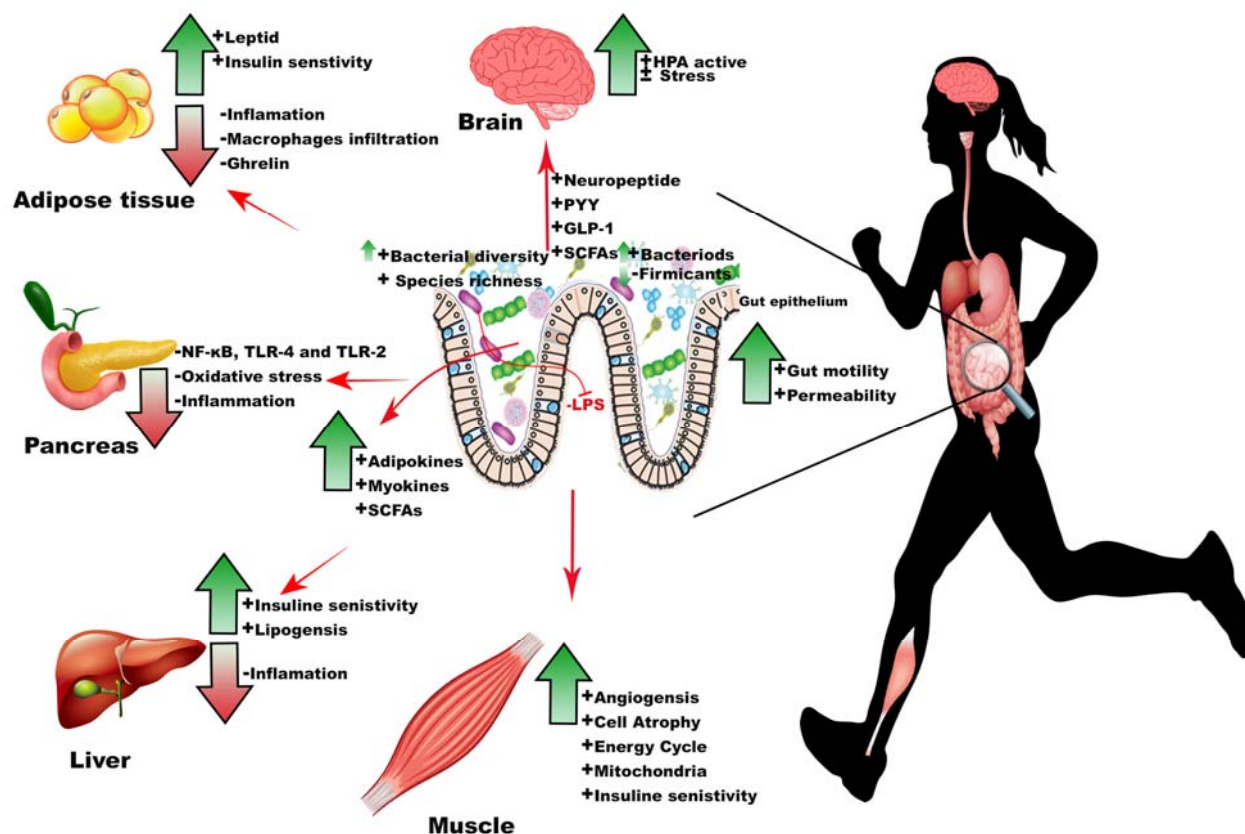


Figure 3. How does physical exercise influence the microbiome and how does the microbiome then impact on metabolism? Physical exercise changes GIT barrier permeability, GIT motility, and microbial diversity, which subsequently change the metabolic profile of liver, adipose tissue, and muscles. An improvement in gut barrier permeability prevents systemic release of lipopolysaccharides (LPS) and inflammation. Furthermore, exercise-induced changes in the microbiome activate the hypothalamus-pituitary-adrenal (HPA) axis and reduce different types of stress. Subsequent release of short-chain fatty acid (SCFAs) and activation of adipokines and chemokines prevent inflammation and autoimmune destruction of beta-cells, and improve energy balance and tissue metabolism.

These exercise-induced metabolic changes were associated with an increased population density of *Bacteroides*, *Flavobacterium*, and *Parabacteroides* genera. These microbial genera have been recognized previously for their protective role against obesity, liver steatosis, and anti-inflammatory capacities [79, 80]. Denou *et al.* reported that exercise could improve the *Bacteroidetes*/*Firmicutes* ratio, metabolic rate, and tricarboxylic acid pathway genes in high-fat diet fed obese mice [66]. In general, overabundance of phylum *Firmicutes* is associated with obesity, whereas increase in Phylum *Bacteroidetes* is associated with weight loss in obese individuals. Similarly, higher species richness and diversity of the GIT microbiome corresponds to improved metabolic markers and energy balance, whereas low bacterial diversity is corre-

lated with insulin resistance, obesity, and dyslipidemia (**Figure 3**) [81]. Since continuous physical exercise is known to prevent or treat these metabolic diseases, these findings may suggest that exercise-associated changes in the GIT microbiome may support the host's metabolic health. Furthermore, enhanced microbiome diversity in endurance performers can also improve antioxidant capacity, which is crucial in lowering insulin resistance and preventing ORMDs [58, 82].

Another plausible mechanism that may impart exercise-induced changes in the microbiome in ORMDs is the change in microbiome metabolic profile. The microbiota-produced SCFA activate the 5' AMP-activated protein kinase (AMPK) pathway to control lipids and glucose metabolism and act as a sensor for cellular energy status [83].

Table 1. Summary of exercise-induced gut microbiome changes observed in different metabolic syndrome conditions

Study format	Exercise format	Clinical features and major findings	Microbiome changes		Reference
			Phylogenetic	Microbiome diversity	
Physical exercise decreased weight gain in rats	Volunteer wheel running, aerobic	Exercise decreased food intake and weight gain, and increased cecal weight and size	Increased butyrate-producing bacterium SM7/11 and T2-87	Exercising rats had a different microbiome clustering pattern from controls	[67]
Exercise corrected toxin-induced microbial dysbiosis in mice	Volunteer wheel running, aerobic	--	93 exclusive taxa were present in both groups (67 in exercising and 26 in sedentary mice)	Exercise increased bacteria abundance and diversity	[45]
Diet restriction- or exercise-induced microbiome changes in rats	Volunteer wheel running, aerobic	Exercise and anorexia decreased body weight and leptin secretion and increased ghrelin secretion	Exercise and anorexia increased <i>Bacteroides</i> , <i>Prevotella</i> , <i>Lactobacillus</i> , and <i>Bifidobacterium</i> population sizes	Exercised rats had a different microbiome clustering pattern from control or diet restricted rats	[63]
Microbiome changes in rats subjected to diet vs. exercise	Volunteer wheel running, aerobic	Exercise more effectively reduced adiposity, adipose inflammation, insulin resistance, and LDL cholesterol	Exercise more effectively increased the <i>Streptococcaceae</i> family and reduced the <i>Rikenellaceae</i> family	Exercise had no impact on species richness or diversity, but had a relatively different clustering pattern	[61]
Exercise altered microbial composition and increased lean mass in juvenile rats	Volunteer wheel running, aerobic	Exercise lowered body weight in adult rats and increased body weight in juvenile rats compared with the sedentary counterpart groups	Juvenile runners had higher <i>Bacteroidetes</i> to <i>Firmicutes</i> ratio	Both adult and juvenile runners had less species richness and evenness than sedentary rats	[54]
Microbiomes of professional or amateur cyclists	Professional or amateur cyclists, aerobic cycling	--	<i>Bacteroides</i> , <i>Prevotella</i> , <i>Eubacterium</i> , <i>Ruminococcus</i> are the dominant bacteria in all cyclists	Microbiome split into three clusters dominated by either <i>Bacteroides</i> , <i>Prevotella</i> , or a mix of many microbes in all cyclists	[60]
Specific microbiome structure improved exercise performance	Exhaustive anaerobic swimming	Specific pathogen-free rats had more liver, muscle, and brown adipose tissue mass and serum antioxidants level	--	--	[56]
Active participation in athletics lowered anxiety and improved oral immunity	Professional soccer and hockey players, both aerobic and anaerobic training	Active athletes had higher cardiorespiratory fitness and oral immunity, and lower anxiety.	Active athletes had lower oral bacterial colony-forming units (CFU)/ml, particularly for <i>Streptococci</i>	--	[87]
Exercise altered microbiome and SCFA concentration	Both volunteer wheel and forced treadmill running	--	Volunteer exercise reduced <i>Turicibacter</i> spp.	Exercise changed microbiome diversity	[88]
Exercise altered the microbiome of obese and hypertensive rats	Controlled treadmill training at 12 to 20m.m-1 speed, aerobic	--	Increased population size of <i>Allobaculum</i> , <i>Pseudomonas</i> and <i>Lactobacillus</i>	Exercise increased alpha diversity and changed clustering pattern	[89]
Exercise improved postoperative rehabilitation	Treadmill training at 20m.m-1 speed, aerobic	Preoperative exercise prevents postoperative neuroinflammation and improves cognition in metabolic syndrome rats	Exercise improved microbiome alpha diversity	--	[76]
Exercise prevented weight gain and altered the microbiome in rats	Volunteer wheel running, aerobic	Exercise lowered body weight and insulin resistance in high-fat diet rats	Exercise decreased <i>Bacteroidetes</i> to <i>Firmicutes</i> ratio	Exercise increased alpha diversity and changed clustering pattern	[62]
Exercise changed the microbiome of diabetic and control mice	Treadmill running at 2 to 4m.m-1 speed, aerobic	--	Exercise decreased <i>Bacteroidetes</i> to <i>Firmicutes</i> ratio in both diabetic and control rats	--	[65]
Exercise changed microbiome composition and influenced cardiac function in mice	Treadmill running at 12-15 m.m-1, aerobic	Exercise inhibited decline in cardiac output and stroke volume in the mouse model for surgical myocardial infarction	Exercise increased <i>Butyricimonas</i> , <i>Prevotella</i> , and <i>Akkermansia</i> compared with the control	Exercise increase bacterial diversity. A positive association was observed between microbial diversity and cardiac function	[90]
Exercise corrected HFD-induced bone loss and microbial dysbiosis in rats	Volunteer wheel running, aerobic	Exercise prevented BWG, bone loss, bone marrow adiposity	Exercise decreased <i>Firmicutes</i> to <i>Bacteroidetes</i> ratio and increased <i>Bifidobacteriaceae</i> population	--	[91]
Exercise improved cardiorespiratory fitness in elderly women	Brisk walking, trunk muscles, aerobic	Exercise increased cardiorespiratory and energy expenditure	Exercise increased <i>Bacteroides</i> , decreased <i>Clostridium</i> populations	--	[92]
Exercise lowered BWG and insulin resistance in HFD rats	Treadmill running, both aerobic and anaerobic	Exercise lowered oxidative stress, inflammation, insulin resistance, and liver damage	HFD rats had a higher <i>Firmicutes</i> / <i>Bacteroidetes</i> ratio that was partially reverted by exercise	Exercise improved microbiome diversity and richness and corrected dysbiosis	[7]
Exercise had limited effect on HFD-induced dysbiosis in mice	Treadmill running, 20m.m-1, aerobic	Exercise failed to improve HFD-induced decrease in exercising capacity and aerobic power	Exercise decreased abundances of <i>Proteus</i> and <i>Vagococcus</i> genera	Exercise had no effect on microbiome alpha or beta diversity	[93]
Intense exercise improved glycemic index and adipose tissue loss in Zucker rats	Treadmill running, 10 to 18 m.m-1, aerobic	Exercise decreased inflammation and adiposity and improved glycemic control in obese rats	Exercise did not affect the abundance of major bacteria taxa	Exercise did not affect species richness or microbiome clustering pattern	[94]

Legend: BWG - body weight gain, CFU - colony forming unit, HFD - high-fat diet, LDL - low-density lipoprotein, SCFA - short-chain fatty acid. References: [7, 45, 54, 56, 60-63, 65, 67, 76, 87-94].

GIT microbiome is critical for sodium-dependent glucose transporter 1 (SGLT1) and G-protein-coupled receptor (GPR) expression on colon epithelial cells. However, it has been observed that antibiotic-induced GIT microbiota depletion in rats results in impaired muscle function in endurance running performance. Furthermore, decrease in microbial diversity is also associated with loss in muscle glycogen levels and insufficient metabolic performance because of SGLT1 and GPR depletion [84]. However, SCFA-regulated GPR stimulation controls size and function (expression of Foxp3 and IL-10) of the colonic T regulatory (Treg) and T helper (Th) cells in intestinal inflammation [85, 86]. Therefore, a decrease in the richness of bacteria species, as observed during antibiotic therapy, may suppress SCFA production, which eventually results in decreased Treg cell production and activation of pro-inflammatory and autoimmune pathways.

Furthermore, exercise increased fecal abundance of arabinose and carnosine metabolites, which are promising candidate for therapies to combat diabetes and obesity [87]. These microbes and metabolites prevent LPS influx, TLR-4-mediated NF- κ B activation, and inflammatory cytokine production. Eventually, downregulation of TLR-2, TLR-4, and the NF- κ B pathway resulting from exercise may improve insulin sensitivity [88]. In brief, exercise-induced changes in GIT microbiome and metabolome help to maintain intestine wall integrity and downregulate both local and systemic immune-inflammatory pathways that may contribute to the prevention of ORMDs [51].

As discussed in the above sections and shown in **Table 1**, most of the literature available provides only common diversity or phylum level taxonomic changes in microbiome in response to exercise, both in health and metabolic diseases. It is therefore reasonable to assume that exercise-induced changes in microbiome diversity could improve host metabolic profile by:

- Decreasing lipid production and storage in visceral organs

- Correcting chronic low-grade inflammation
- Decreasing insulin resistance

We have attempted to establish a link between exercise-induced phylogenetic changes in microbiome and metabolic disorders that associate microbiome changes with predictive immune-inflammatory pathways. Eventually, this may provide a mechanistic understanding of the beneficial effects of physical exercise in metabolic disorders through microbiome changes.

7. Conclusions

The observations reviewed here suggest a plausible microbiome basis for exercise-induced prevention of metabolic diseases, and provide an opportunity to explore novel avenues to prevent metabolic diseases in critically lethargic subjects who are incapable of voluntarily exercise to manage their weight. Although, most of the favorable health outcomes are independent of microbiome-host interactions, there is also evidence to indicate that some of the benefits may be mediated by exercise-induced adaptations in gut microorganisms.

Reciprocal associations between immune-inflammatory mediators and exercise-induced microbiome/metabolome corrections may engage several cellular energy sensors and trigger adipokine and myokine activation that monitor energy balance and tissue metabolism. Although abundant evidence is available on the role of the microbiome in metabolic disorders, no significant clinical findings are available to prevent or correct these disorders. Extensive work is needed on metagenomics beyond taxonomic diversity profiling. More mechanistic research is required to connect exercise with microbiome, body metabolism, and energy homeostasis.

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