



Exerkines in health, resilience and disease

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Abstract | The health benefits of exercise are well-recognized and are observed across multiple organ systems. These beneficial effects enhance overall resilience, healthspan and longevity. The molecular mechanisms that underlie the beneficial effects of exercise, however, remain poorly understood. Since the discovery in 2000 that muscle contraction releases IL-6, the number of exercise-associated signalling molecules that have been identified has multiplied. Exerkines are defined as signalling moieties released in response to acute and/or chronic exercise, which exert their effects through endocrine, paracrine and/or autocrine pathways. A multitude of organs, cells and tissues release these factors, including skeletal muscle (myokines), the heart (cardiokines), liver (hepatokines), white adipose tissue (adipokines), brown adipose tissue (baptokines) and neurons (neurokines). Exerkines have potential roles in improving cardiovascular, metabolic, immune and neurological health. As such, exerkines have potential for the treatment of cardiovascular disease, type 2 diabetes mellitus and obesity, and possibly in the facilitation of healthy ageing. This Review summarizes the importance and current state of exerkine research, prevailing challenges and future directions.

Resilience

Resilience is the ability of the body to resist, adapt to, recover or grow in response to stressors.

High-intensity interval training

High-intensity interval training is a form of exercise training characterized by bursts of high-intensity activity followed by less intense recovery periods.

Irrefutable evidence supports the importance of physical activity, exercise and cardiorespiratory fitness in the prevention and treatment of chronic diseases, such as cardiovascular disease, obesity, type 2 diabetes mellitus, cognitive decline and many cancers, while enhancing the immune system, healthspan, longevity and resilience¹ (FIG. 1). Conversely, physical inactivity poses a major public health threat, as it is associated with increased mortality² and a notable economic burden³. Moreover, the COVID-19 pandemic clearly reinforces the relevance of physical activity for health, due to the effects of COVID-19-related reductions in physical activity⁴ and increases in sedentary behaviour, especially due to COVID-19-related quarantine⁴. Moreover, physical inactivity is associated with increased risk of severe COVID-19 outcomes⁵.

Although the terms ‘exercise’ and ‘physical activity’ are commonly used interchangeably, exercise is typically regarded as intentional physical activity, such as aerobic training¹, resistance training¹ or high-intensity interval training^{6,7}. By contrast, physical activity encompasses exercise as well as usual occupational and/or domestic activity¹. Promoting physical activity remains a

critical intervention to reduce the incidence and prevalence of common metabolic diseases. In the USA, official guidelines for physical activity were first published in 1995 and recommended that every US adult should accumulate at least 30 min of moderate intensity physical activity on most, preferably all, days of the week⁸. Subsequently, these guidelines have evolved¹. In 2020, the World Health Organization stated that all adults should aim for 150–300 min of moderate intensity physical activity per week or 75–150 min of vigorous intensity physical activity per week or an equivalent combination of moderate intensity and vigorous intensity physical activity per week⁹. Despite these recommendations, objective data obtained with accelerometers of physical activity in the US population indicated poor adherence to recommended guidelines, with only 5% of US adults having more than 30 min of moderate intensity physical activity per day¹⁰.

In this Review, we focus on the potential role of exerkines in driving the established benefits of exercise, such as preventing and mitigating disease, promoting health and increasing resilience. The term exerkine was coined in 2016 (REF.¹¹), although the concept of humoral

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Key points

- Although the benefits of exercise in enhancing health and treating disease are well-acknowledged, the molecular mechanisms underlying exercise-associated benefits remain ill-defined and are actively being investigated.
- ‘Exerkines’ encompass a broad variety of signalling moieties released in response to acute and/or chronic exercise that exert their effects through endocrine, paracrine and/or autocrine pathways.
- Exerkines can come in many forms, such as hormones, metabolites, proteins and nucleic acids; interest is increasing in moving beyond singular changes of specific factors to profiling exerkine alterations using ‘omics’ platforms.
- There is burgeoning interest in the role of extracellular vesicles, which are membranous structures released from cells, in serving as important carriers of molecular signals and drivers of inter-organ crosstalk related to exercise.
- Multiple organ systems, including the cardiometabolic system, nervous system and immune system, produce exerkines and are influenced by exerkines, which probably contributes to the pleiotropic and variable response to exercise.
- Emerging research on exerkines suggests multiple promising avenues for translational research and therapeutic modulation to capture exercise-associated benefits; enhanced rigour in experimental design will facilitate comparison between studies.

Exerkines

Exerkines encompass a broad variety of signalling moieties that are released in response to acute and/or chronic exercise that exert their effects through endocrine, paracrine and/or autocrine pathways.

Acute exercise

Acute exercise is typically considered a single episode of exercise (often resistant or aerobic exercise) that is completed during one visit.

factors mediating the benefit of exercise has long been recognized. A prime example is lactic acid; its secretion from skeletal muscle was identified over 100 years ago¹². In 1961, Goldstein speculated about the existence of a non-insulin humoral factor that regulates the effect of exercise on skeletal muscle and liver glucose utilization¹³. For the purposes of this Review, we define an exerkine as a signalling moiety released in response to acute exercise and/or chronic exercise, exerting its effects through endocrine, paracrine and/or autocrine pathways.

As skeletal muscle comprises approximately one third of body mass and has an important role in exercise¹⁴, the effects of physical activity (FIG. 1) were initially attributed to blood-borne factors, particularly muscle-secreted

hormones (myokines)¹⁵. Of the myokines, IL-6 has been the most extensively studied since its discovery in 2000 (REF. 16). Subsequent exerkine work has broadened to include exercise-related humoral factors arising from the heart (cardiokines), liver (hepatokines), white adipose tissue (WAT; adipokines) and brown adipose tissue (BAT; batokines) and the nervous system (neurokines), with local autocrine effects (affecting the cell of origin) and paracrine effects (affecting adjacent cells) (TABLE 1). Exerkines are increasingly recognized to include a broad range of signalling moieties, including cytokines, nucleic acids (microRNA¹⁷, mRNA and mitochondrial DNA), lipids and metabolites, which are frequently driven by cell-specific extracellular vesicle secretion¹⁸.

Understanding the role of exerkines in the physiological and biological response to exercise is a principal objective of many investigations sponsored by the National Institutes of Health (NIH)¹⁹, given the demonstrated benefits of exercise in enhancing and prolonging human health across the lifespan. In 2020, the National Heart, Lung, and Blood Institute and the National Institute of Diabetes and Digestive and Kidney Diseases convened a virtual 2-day public workshop inviting 21 international experts to discuss ‘Exerkines in Health, Resilience, and Diseases’. The workshop executive summary was published online²⁰, laying the foundation for this article. In this Review, we summarize the importance and current state of exerkine research, the prevailing challenges and future directions.

**Exercise response variability
Potential role for exerkines**

The majority of exercise research has been limited to studies with genetically homogeneous animal models and/or small numbers of human participants. Moreover,

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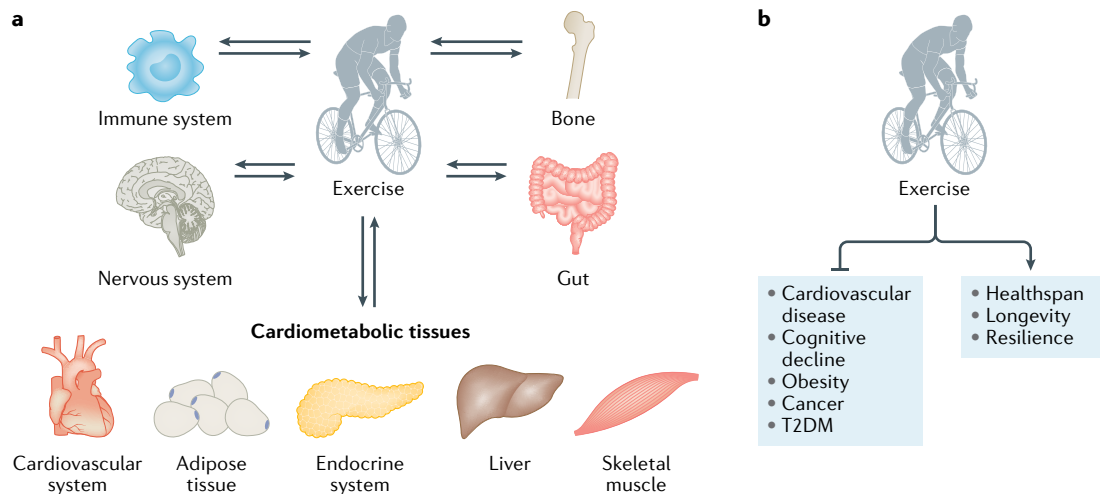


Fig. 1 | The systemic effects of exercise. **a** | Organs and tissues that can serve as source of exerkines and that are directly affected by exercise. **b** | Exercise results in profound health benefits, including reductions in the presence or severity of certain diseases, as well as increases in healthspan, longevity and resilience. T2DM, type 2 diabetes mellitus.

the physiological response to a structured exercise training stimulus remains highly variable in humans and animals owing to a multitude of external and internal factors. In terms of external factors, the context of exercise matters, as exercise timing relative to circadian rhythms²¹, fed–fasting status²² or post-exercise dietary composition²³ might influence metabolic outcomes. This variability is well-detailed in a study published in 2022, which included an atlas describing the time-dependent effects of exercise across multiple tissues after a single bout of treadmill exercise²⁴. Addressing these external factors will require careful consideration of the exercise exposure, controlling the exercise exposure’s environmental context and serial sampling of blood and tissue prior, during and after exercise. This level of rigor is needed to ‘reduce the noise’ and facilitate interpretation of the temporal signatures of circulating and tissue-specific exerkines. In terms of internal factors, genetics have a critical role in the response to exercise. The Health, Risk Factors, Exercise Training and Genetics (HERITAGE) family study involved 20 weeks of supervised aerobic exercise training in 481 sedentary, healthy, white adults from 98 two-generation families, and found that the maximal heritability estimate for the aerobic capacity response was roughly 47%²⁵. Furthermore, chronic exercise training elicited a ‘non-response’ in terms of improved aerobic capacity in ~20% of individuals²⁵. In addition, 7–15% of individuals demonstrated an ‘adverse response’ regarding alterations in systolic blood pressure, as well as in fasting levels of HDL cholesterol, triglycerides and insulin^{25,26}.

To drive the application of precision medicine to exercise, investigations into the mechanisms underlying the response variability to exercise are sorely needed. The contribution of exerkines to the variation in exercise response remains under active research and will be a key focus of the [Molecular Transducers of Physical Activity Consortium](#) (MoTrPAC; NCT03960827). This NIH-supported research consortium is designed to discover and broadly characterize the range of molecular

transducers that underlie the variable effects of exercise in humans and animals. For humans, MoTrPAC has several unique features: its size (2,280 estimated participants); its recruitment of sedentary participants who will undergo a 12-week programme of aerobic exercise, resistance exercise or no exercise (control), with a comparison group of highly active endurance exercise or resistance exercise participants; and its time course analysis of changes in tissue (muscle and adipose tissue) and plasma metabolites¹⁹. In animals (~800 studied), the unique feature of MoTrPAC will be its focus on detailed biospecimen analysis across multiple time points and organs, which cannot be easily replicated in humans, in young (6 months) and old (18 months) male and female rats after an acute (single session) bout of treadmill exercise, or after chronic treadmill exercise (8 weeks) versus non-exercised control rats¹⁹. Identification of an exerkine or a panel of exerkines, which capture the benefits of exercise, would have potential implications for improving the health of those unable to exercise, such as those with ageing-associated exercise intolerance.

Influence of exercise exposure

Exerkines are secreted in response to acute exercise, which is usually a single episode of either aerobic or resistance exercise. Chronic exercise is also associated with altered humoral factors, even in the resting state, suggesting that exerkine alterations can reflect the effects of chronic training²⁷.

The acute exerkine response is influenced by the type of exercise, duration of exercise, underlying fitness, fed–fasting status and sample timing after exercise. In a human model, the blood concentration of glucose typically remains stable during acute exercise, with the liver releasing glucose for brain and skeletal muscle usage²⁸. During exercise, skeletal muscle also uses lipid as fuel, which originates from the triglycerides stored in muscle and free fatty acids (FFAs) released from WAT²⁹. The classic exerkines released during acute exercise, as found in human and animal models, include IL-6, IL-8, IL-1

Chronic exercise

Chronic exercise is typically described as multiple exercise episodes (often resistant or aerobic exercise) performed over the course of weeks to months.

MicroRNA

MicroRNAs are non-protein-coding RNA molecules that are regulated in a transcriptional or post-transcriptional fashion to affect mRNA transcription and/or degradation.

Table 1 | Examples of paracrine and autocrine effects of exerkines

Exerkine ^a	Source tissue	Affected tissue
Autocrine effects^b		
12,13-diHOME	–	BAT ^{91,169}
Apelin	–	Muscle ^{94,96}
Adiponectin	–	WAT ¹⁵³
BDNF	–	Brain ¹⁷⁰ , muscle ¹⁷¹
FGF21	–	WAT ^{56,172}
HSP72	–	Muscle ^{107,173}
IL-6	–	Muscle ^{15,54,120}
IL-7	–	Muscle ¹⁰²
IL-15	–	Muscle ¹⁷⁴
Lactate	–	Muscle ¹⁷⁵
LIF	–	Muscle ¹⁰³
Musclin (also known as osteocrin)	–	Muscle ^{43,176} , bone ^{62,177} , brain ¹⁷⁸
Myostatin	–	Muscle ⁴⁴
Nitric oxide	–	Endothelium ¹⁷⁹
Reactive oxygen species	–	Muscle ¹⁸⁰
SPARC	–	Muscle ^{63,181}
SDC4	–	Muscle ¹⁰⁴
TGFβ1	–	Muscle ¹⁸²
Paracrine effects		
Adiponectin	Adipose tissue	Muscle ⁵⁸
Angiopietin 1	Vascular smooth muscle	Vasculature ⁴⁷
BAIBA	Muscle	WAT ⁵⁵ , bone ⁵⁹
BDNF	Muscle	Nerves ⁵³
Fractalkine	Muscle	Leukocytes ⁶⁶
FGF21	WAT	BAT ⁵⁶
IL-6	Muscle	WAT ^{15,121}
IL-7	Muscle	Bone ⁶⁰
IL-8	Muscle	Vasculature ⁴⁸
IL-13	Tissue-resident ILC2	Muscle ¹³⁶
IL-15	Muscle	WAT ⁵⁷
LIF	Nerves, immune cells	Muscle ^{183,184}
Musclin	Bone	Cartilage ⁶²
Myostatin	Muscle	Bone ⁶¹
SPARC	Muscle	Extracellular matrix ⁶³
SDC4	Endothelium	Muscle ¹⁸⁵
TGFβ1	Muscle	Extracellular matrix ⁶⁴
TGFβ2	Adipose tissue	Muscle, BAT ⁴²
VEGF	Muscle	Endothelium ^{45,46}

12,13-diHOME, 12,13-dihydroxy-9Z-octadecenoic acid; BAIBA, β-aminoisobutyric acid; BAT, brown adipose tissue; BDNF, brain-derived neurotrophic factor; FGF21, fibroblast growth factor 21; HSP72, heat shock protein 72; ILC2, type 2 innate lymphoid cells; LIF, leukaemia inhibitory factor; SDC4, syndecan 4; SPARC, secreted protein acidic and rich in cysteine; TGFβ1, transforming growth factor-β1; TGFβ2, transforming growth factor-β2; VEGF, vascular endothelial growth factor; WAT, white adipose tissue. ^aAlthough exerkine effects are commonly thought to be distant (endocrine) from the originating tissue, exerkines also exert local effects within the originating tissue (autocrine) and neighbouring tissues (paracrine). More details and relevant references are provided in Supplementary Table 1. ^bFor exerkines with autocrine effects, the source tissue is indicated as ‘–’, as the source tissue and affected tissue are the same.

receptor antagonist (IL-1RA) and IL-10. In a study in humans in which blood samples were collected before and after a marathon, plasma levels of several cytokines (IL-6, IL-1RA, IL-10 and tumour necrosis factor (TNF)) were higher than baseline levels when collected immediately after exercise, peaking 1–2 h after exercise and remaining elevated for ~4 h after exercise³⁰. Certainly, the type and intensity of exercise matters. As an example, the exerkine response to high-intensity interval training depends on exercise intensity; higher exercise intensity corresponds with higher plasma levels of IL-6, while IL-10 levels remain unchanged compared with the levels before exercise⁶. Supplementary Table 1 shows examples of singular exerkine alterations. Currently, exerkine research is evolving from measuring singular exerkine changes to characterizing metabolic profiles³¹, for which challenges in analysis and interpretation remain (TABLE 2).

Notably, the acute exerkine response does not necessarily parallel the chronic exerkine response (Supplementary Table 1). Typically, acute exercise exposure is associated with responses focused on the maintenance of metabolic homeostasis, with acute inflammation balanced by anti-inflammatory mediators³⁰ and accommodating shifts in fuel utilization. By contrast, chronic exercise exposure is associated with responses focused on long-term metabolic adaptations and decreased inflammation²⁷. However, when investigating chronic exercise exposure, the caveats of recent (24–72 hours prior to exercise) acute exercise or dietary composition, underlying fitness, fed–fasting status, circadian timing and training modality need to be considered. Moreover, the effects of exercise could also be influenced by alterations at the level of the exerkine receptor, in addition to alterations in plasma levels of exerkines. For example, in humans, chronic exercise training reduces plasma concentrations of IL-6; however, this effect could be partially mitigated by increased skeletal muscle mRNA expression of IL-6 receptor³².

Exerkines: technical considerations

Discovery techniques

Exerkine research is increasingly focused on measuring changes across a broad swathe of factors rather than singular change. Specifically, interest is increasing in ‘omics’ technology to capture exercise-related changes in lipids (lipidomics), metabolites (metabolomics), proteins (proteomics), gene expression (transcriptomics) and DNA alterations (epigenomics)³¹ (TABLE 2). A paper in 2020 studied humans across the spectrum of insulin sensitivity ($n = 36$, ranging from people with high insulin sensitivity to patients with diabetes mellitus) who performed an acute bout of treadmill-based exercise to reach peak oxygen uptake. This exercise exposure altered >50% of measured molecules, spanning platforms based on lipidomics, metabolomics, proteomics, transcriptomics and epigenomics³¹. TABLE 2 lists commonly used platforms for exerkine analysis, including their relative advantages and disadvantages. Mass spectrometry is often used for targeted and untargeted omics analysis, whereas immunoassays are commonly used for analysis of proteins and metabolites. Genetic analyses include RNA sequencing,

methylation sequencing (Methyl-seq) and assay for transposase-accessible chromatin with high-throughput sequencing (ATAC-seq)³³. Together, these platforms provide a rich profile of the molecular and epigenomic changes that occur in response to acute and chronic exercise.

Extracellular vesicles

In the exerkine field, interest is also intensifying in the role of extracellular vesicles as important carriers of molecular signals and drivers of inter-organ crosstalk related to exercise¹⁸. Extracellular vesicles are membranous structures that are released from almost all cell types, with cell-specific profiles. They vary in size, ranging from 150 nm to 1,000 nm, and carry an assortment of material, including proteins, nucleic acids and lipids^{18,34}. The content of extracellular vesicles reflects the unique and varied composition of the cells from which they are released. As an example of extracellular vesicles acting as an exerkine in humans, acute exercise increases plasma levels of various microRNAs after exercise, and chronic exercise increases various microRNAs in the resting state¹⁷, supporting the possibility of microRNAs exerting their endocrine effects via extracellular vesicle-based transport^{11,34}.

Extracellular vesicles can be routinely isolated and profiled from cell culture media. Studying the molecular cargo of plasma-derived extracellular vesicles, however, remains uniquely challenging. Critical to extracellular vesicle analysis is the careful consideration of pre-analytical steps, including proper collection and isolation. Isolation techniques include ultracentrifugation (using a differential density gradient), ultrafiltration, size exclusion chromatography, high-resolution mass spectrometry, capillary electrophoresis, asymmetric-flow field-flow fractionation and immunoaffinity capture³⁵. Moreover, contamination at the time of collection needs to be considered

as extracellular vesicles can arise from ex vivo platelet activation^{36,37}. A 2021 paper presented an optimized size-exclusion chromatography method for proteomic analysis of plasma-derived extracellular vesicles from platelet-poor plasma; this technique has greater precision than conventional extracellular vesicle techniques and demonstrates a distinct exosome protein cargo profile after acute exercise in humans³⁷.

Autocrine, paracrine and/or endocrine effects

Initially, exerkine research focused on changes in plasma levels of cytokines, especially before and after an acute exercise exposure³⁰. The classic exerkines are cytokines, of which IL-6 has been the most extensively studied since its identification as a myokine in 2000 (REF.¹⁶). Subsequently, the field evolved into examining the endocrine effects of exerkines, where molecules secreted from the source tissue, traditionally viewed as skeletal muscle, affect distant tissues¹⁵. Especially within the past 15 years, interest has been increasing in the local effects of exerkines, either on the secreting tissue (autocrine) or the adjacent environment (paracrine)³⁸ (TABLE 1), non-muscle exerkine sources³⁹ (Supplementary Table 1) and exerkine profiling rather than on singular exerkine alterations³¹ (TABLE 2).

A common perception among the general scientific community is the view of exerkines as a cytokine exerting its effects in an endocrine fashion, affecting tissues distant from the originating tissue. Exerkines are not merely cytokines, however, as hormones, neurotransmitters or metabolites associated with exercise, such as catecholamines⁴⁰, lactate⁴¹ or FFAs²⁹, can also serve as exerkines with endocrine signalling potential⁴².

From an autocrine standpoint, exerkines affect their origin cells by coupling local energy balance to tissue growth and metabolic homeostasis (TABLE 1). For example, in skeletal muscle, myocytes secrete factors such as lactate⁴¹, musclin⁴³ and myostatin⁴⁴ that couple exercise

Table 2 | Common platforms for exerkine measurement

Platform	Commonly measured exerkines	Advantages	Disadvantages	Refs
Untargeted mass spectrometry	Lipidomics; metabolomics; proteomics	Profiles large numbers of molecules (>1,000 for metabolites and lipids; over 5,000 proteins); fairly inexpensive	Relative quantification rather than absolute quantification; bias towards abundant molecules; throughput lower than targeted assays	31
Targeted mass spectrometry	Lipidomics; metabolomics; proteomics	Accurate and absolute quantification; fast	Fewer molecules than untargeted assays	31
Immunoassays	Proteins (including cytokines) and metabolites	Accurate and absolute quantification; measures low-abundance molecules	Fewer molecules than untargeted assays	31
RNA-seq	Transcripts; splicing isoforms	Comprehensive (>10,000 genes); accurate absolute quantification	Does not reflect protein levels or protein modification	31,33
Methyl-seq	DNA methylation	Measures stable epigenome changes	Expensive	31,33
ATAC-seq	Open chromatin	Measures chromatin epigenome changes; easy to perform	Does not reflect RNA or protein expression levels	31,33

ATAC-seq, assay for transposase-accessible chromatin using sequencing; Methyl-seq, methylation sequencing; RNA-seq, RNA sequencing.

to changes in mitochondrial biogenesis and myocyte substrate utilization.

Muscle and other highly metabolically active tissues can also secrete exerkinins that exert local (paracrine) effects³⁸. For example, muscle secretes vascular endothelial growth factor (VEGF)^{45,46}, angiopoietin 1 (REF.⁴⁷) and IL-8 (REF.⁴⁸) to regulate tissue angiogenesis, modulate blood flow and increase nutrient availability to support tissue growth^{27,49–52} (TABLE 1). Exercise-related paracrine effects are also observed in the nervous system⁵³, adipose tissue^{42,54–58}, bone^{59–61}, cartilage⁶², extracellular matrix^{63,64} and the immune system^{65,66}.

Tissue-specific exerkin relationships

The cardiovascular system

Physical activity reduces the risk of cardiometabolic disease and mortality. Although exercise mitigates traditional cardiovascular risk factors, such as obesity and dyslipidaemia, these benefits incompletely account for the effects of exercise on cardiometabolic health. Studies in both humans and animal models support a role for exerkinins potentially enhancing cardiometabolic health (FIG. 2; TABLE 3; Supplementary Table 1). Exerkinins could also oppose multiple mechanisms associated with cardiovascular disease, such as persistent systemic inflammation, dysregulated energy balance and fuel utilization.

Furthermore, the enhanced angiogenesis associated with certain exerkinins could mitigate ischaemia. Notably, exercise might improve endothelial function. As the vascular endothelium lies at the interface between blood and tissue, its wide-ranging distribution and strategic positioning supports its potential role as an initiator and recipient of exerkinin-related effects. For example, interplay between the endothelium and established exerkinins, such as nitric oxide⁵² and VEGF²⁷, has been shown to influence vascular tone, inflammation, regeneration and thrombosis, and has an important role in cardiovascular and overall resilience⁶⁷.

Contracting skeletal muscle produces many molecules that can enhance the cardiovascular system. Studies in humans^{15,45,68–74} and animal models^{15,43,49,75,76} have shown that the exerkinins angiopoietin 1 (REFS^{49,68,69}), fractalkine^{70,71}, fibroblast growth factor 21 (FGF21)^{72,73}, IL-6 (REF.¹⁵), IL-8 (REFS^{50,74}), myonectin⁴³, myonectin⁷⁵ and VEGF^{27,45,76} are generally increased with acute exercise; however, the exerkinin response to chronic training, as measured by assessing exerkinins in plasma during the resting state, can be quite variable and discrepant from the acute effects. As shown in TABLE 3 and Supplementary Table 1, examples include angiopoietin^{68,69}, FGF21 (REFS^{73,77}), fractalkine⁷⁰, IL-6 (REFS^{30,78}) and IL-8 (REFS^{50,74}).

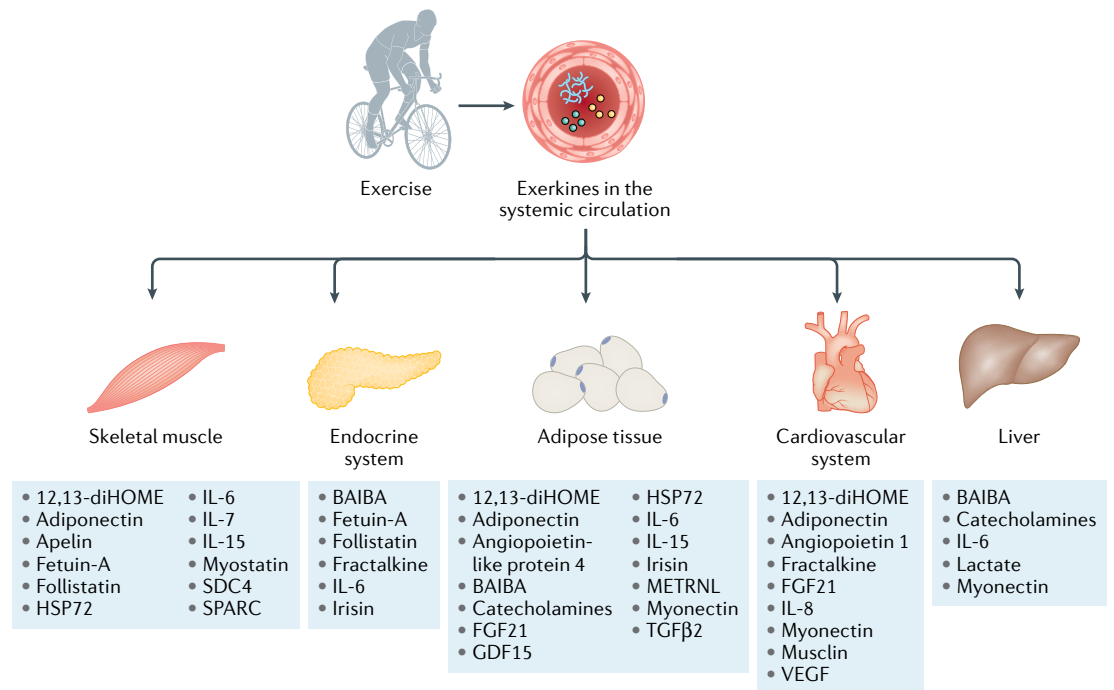


Fig. 2 | Examples of exerkinins that affect the cardiometabolic system. Exerkinins released after exercise into the systemic circulation (see TABLE 3 for tissue sources, detailed effects and relevant references), including proteins (blue lines), metabolites (yellow circles) and extracellular vesicles (green circles), affect the cardiometabolic system. The effects are wide-ranging and systemic. In the cardiovascular system, exerkinins enhance vascularization and angiogenesis, as well as improve blood pressure, endothelial function and overall fitness, resulting in cardioprotection. In adipose tissue, exerkinins increase fatty acid uptake, enhancing lipolysis, thermogenesis and glucose metabolism. In the liver, exerkinins enhance glucose metabolism and fatty acid uptake. In skeletal muscle, exerkinins enhance muscle formation, maintenance and repair, glucose uptake, lipid oxidation, mitochondrial biogenesis and muscle capillarization. In the pancreas, exerkinins enhance cell viability and influence insulin secretion. Commonly described exerkinins are noted. 12,13-diHOME, 12,13-dihydroxy-9Z-octadecenoic acid; BAIBA, β-aminoisobutyric acid; FGF21, fibroblast growth factor 21; GDF15, growth and differentiation factor 15; HSP72, heat shock protein 72; METRNL, meteorin-like; SDC4, syndecan 4; SPARC, secreted protein acidic and cysteine rich; TGFβ2, transforming growth factor-β2; VEGF, vascular endothelial growth factor.

Table 3 | Examples of exerkinins that affect the cardiometabolic system

Name	Species or model ^a	Main tissue of origin	Main target tissue	Effect ^b	Main biological action	Response to acute exercise bout ^c	Response to chronic training ^c	Refs
12,13-diHOME	H, A, C	BAT	BAT, skeletal muscle, heart	A, E	Increases fatty acid uptake	↑	↑	91,93, 169
Adiponectin	H, A, C	WAT	Many tissues, including liver, muscle, heart	A, P, E	Enhances glucose and lipid utilization	↑	↑,↔	58,153, 186
Angiopoietin I	H, A, C	Skeletal muscle	Vasculature	P	Enhances angiogenesis	↑,↔	↑,↓	47,68, 69
Angiopoietin-like protein 4	H, A	Liver,	WAT	E	Decreases lipoprotein lipase activity and enhance WAT lipolysis to increase plasma FFAs and triglycerides	↑	↑,↔	39,68, 79
Apelin	H, A, C	WAT, skeletal muscle	Skeletal muscle,	A, E	Enhances skeletal muscle mass and mitochondria	↑	↑,↔	94–96
BAIBA	H, A, C	Skeletal muscle	WAT, Liver, β-cells	E, P	Enhances 'browning' of white adipocytes and β-oxidation in liver; attenuates insulin secretion from β-cells	↑	↑	55,113, 187
Catecholamines	H	Adrenal	Skeletal muscle, WAT	E	Stimulates glycogenolysis; stimulates lipolysis of WAT	↑	↔	40
Fetuin-A	H	Liver	Skeletal muscle, pancreas	E	Impedes β-cell function; increases insulin resistance	↓,↔	↓	39,72, 97,101, 115
Fractalkine (also known as CX3CL1)	H, A, C	Skeletal muscle	Leukocytes, endothelium, myocytes, β-cells	P, E	Increases inflammatory, angiogenic, and chemotactic factors; regulates β-cell secretion	↑	↔	66,70, 71,114
FGF21	H, A	Many tissues, especially liver; also WAT	Many tissues, including heart and WAT	E, A, P	Augments fuel utilization (glucose, lipid); protects from apoptosis	↑	↔	56,72, 73,106, 172,188
Follistatin	H, A	Many tissues, especially liver	Skeletal muscle	E	Decreases serum levels of myostatin to enhance skeletal muscle growth; might affect glucose homeostasis	↑	↑	39,97, 98,100, 116,189
GDF15	H, A	Many sites	Many sites	E	Marker of stress response, promotes WAT lipolysis	↑	↑	80,190
HSP72	H, A	Many tissues, especially liver	Many sites	A, E	Maintains cellular homeostasis; protects cells from stress	↑	↑	107,173
IL-6	H, A, C	Primarily skeletal muscle	Many sites	A, P, E	Multiple effects: including enhancing WAT lipolysis; lipid oxidation; glucose homeostasis; anti-inflammatory response; skeletal muscle growth	↑	↓	15,16, 30,78,81, 119–121
IL-7	H, A, C	Skeletal muscle	Skeletal muscle, bone	A, P	Regulates skeletal muscle development	↑	↔	102,191
IL-8	H, A, C	Skeletal muscle	Endothelium	P	Regulates tissue angiogenesis and blood flow	↑,↔	↔	48,50, 74
IL-15	H, A, C	Many tissues, especially immune cells	Many tissues, especially immune cells	A, P, E	Regulates immune cell functioning and might reduce WAT mass; improves glucose homeostasis; promotes skeletal muscle growth	↑,↔	↔	57,99, 174
Irisin (also known as FNDC5)	H, A	Skeletal muscle	WAT, bone, β-cells, brain	E	Benefits primarily in animal models: increases fatty acid uptake; beiging of WAT; improves insulin secretion	↑	↓,↔	84,117, 167
Lactate	H	Skeletal muscle	Many tissues, including central nervous system	A, E	Provides substrate for hepatic gluconeogenesis	↑	↔	41,175

Table 3 (cont.) | Examples of exerkinins that affect the cardiometabolic system

Name	Species or model ^a	Main tissue of origin	Main target tissue	Effect ^b	Main biological action	Response to acute exercise bout ^c	Response to chronic training ^c	Refs
METRNL	H, A, C	Many tissues, including WAT and skeletal muscle	Immune cells	E	Increases energy expenditure; improves glucose tolerance	↑	↑	134,192, 193
Myonectin (CTRP15)	H, A	Skeletal muscle	Liver, WAT, heart	E	Promotes fatty acid uptake; might be cardioprotective	↑,↔	↔	27,75, 82,194
Musclin (also known as osteocrin)	H, A, C	Skeletal muscle, bone, brain	Skeletal muscle, heart, vasculature, cartilage, brain	A, P, E	Regulates mitochondrial biogenesis and might exacerbate insulin resistance	↑	↓	27,43, 176,195
Myostatin (also known as GDF8)	H, A, C	Many tissues, especially skeletal muscle and WAT	Many tissues, especially skeletal muscle and bone	A, P, E	Blunts skeletal muscle growth and glucose uptake	↑,↔	↔	44,73, 99,189
SPARC	H, A, C	Many tissues	Many tissues	A, P, E	Regulates cell function and tissue remodelling	↑,↔	↔	63,181
SDC4	H	Many tissues	Immune system	A, P, E	Involved in cell–extracellular matrix crosstalk, inflammation and skeletal muscle growth	↑	↑	104,185, 196,197
TGFβ1	H, A, C	Many tissues	Many tissues especially immune cells	A, P, E	Chemotactic factor for immune cells; affects skeletal muscle growth	↑	↑	64,65, 182,198, 199
TGFβ2	H, A, C	Adipose tissue	Many tissues, especially muscle and immune cells	P, E	Promotes glucose and fatty acid metabolism; reduces inflammation	↑	↑	42
VEGF	H, A, C	Many tissues, especially skeletal muscle	Vascular endothelium	P, E	Promotes angiogenesis and exercise-induced neurogenesis	↑,↔	↑	27,45, 46

12,13-diHOME, 12,13-dihydroxy-9Z-octadecenoic acid; BAIBA, β-aminoisobutyric acid; BAT, brown adipose tissue; CTRP15, complement C1q tumour necrosis factor-related protein 15; CX3CL1, chemokine (C-X3-C motif) ligand 1; FFAs, free fatty acids; FGF21, fibroblast growth factor 21; FNDC5, fibronectin type III domain containing 5; GDF8, growth and differentiation factor 8; GDF15, growth and differentiation factor 15; HSP72, heat shock protein 72; METRNL, meteorin-like; SDC4, syndecan 4; SPARC, secreted protein acidic and cysteine rich; TGFβ1, transforming growth factor-β1; TGFβ2, transforming growth factor-β2; VEGF, vascular endothelial growth factor; WAT, white adipose tissue. ^aRelevant species or models are human (H), animal (A) or cell (C). ^bEffects are autocrine (A), paracrine (P) or endocrine (E). ^cArrows indicate: ↑, plasma levels increase; ↓, plasma levels decrease; ↔, plasma levels remain the same.

Adipose tissue

Exercise facilitates WAT lipolysis to provide FFA for utilization as fuel²⁹. Although this lipolysis was typically attributed to adrenaline release⁴⁰, acute exercise in humans also releases additional molecules⁷⁹, such as growth and differentiation factor 15 (GDF15)⁸⁰ and IL-6 (REF⁸¹), which also affect lipolysis (TABLE 3; Supplementary Table 1). Lipolysis is not the only avenue by which exerkinins can affect adipose tissue mass. For example, in myonectin knockout mice, WAT lipolysis is unaffected; however, dietary lipid clearance is impaired compared with lipid clearance in wild-type mice, resulting in increased WAT mass and decreased liver steatosis⁸².

A potential effect of exercise on WAT is ‘browning’, where WAT increases mitochondrial content, metabolic rate and heat production. WAT browning might have metabolic importance, as individuals with PET-CT-defined BAT had a decreased prevalence of cardiometabolic disease, particularly if they had overweight or obesity⁸³. In a mouse model, fibronectin type III domain containing 5 is cleaved in the muscle cell and secreted as irisin, which induces WAT browning to increase energy expenditure and consequently reduce

obesity⁸⁴. Transgenic mice that overexpress muscle peroxisome proliferator-activated receptor-γ coactivator 1-α (PGC1α) have higher circulating levels of irisin and increased WAT browning than control mice⁸⁴. Hence, the initial excitement regarding irisin as an exerkinin, as exercise generally increases muscle PGC1α expression in both animal models⁸⁵ and humans⁸⁶.

As the irisin findings and exercise-induced browning of WAT concepts were re-evaluated in humans, the initial excitement was subsequently tempered. Although acute exercise in humans generally increases plasma levels of irisin⁸⁷, the effect of chronic exercise training remains highly variable. A meta-analysis of several randomized controlled trials even showed lower levels of irisin after training than before training⁸⁸. Trained athletes have lower BAT activity and no difference in WAT browning markers compared with lean, sedentary men⁸⁹; this observation is supported by a study in humans of chronic exercise training, which did not find any browning of WAT (as assessed by biopsy)⁹⁰. Thus, whether exercise can brown WAT in humans, especially through an irisin-mediated pathway, remains controversial^{55,83,84,90}.

Adipose tissue can also secrete exerkinins. A prime example is 12,13-dihydroxy-9Z-octadecenoic acid (12,13-diHOME), which is secreted from BAT and increases skeletal muscle oxidative capacity⁹¹. In humans, circulating levels of 12,13-diHOME are inversely associated with adipose tissue mass, fasting blood levels of insulin and blood levels of triacylglycerol⁹². A 2021 study showed that BAT transplantation in mice increased plasma levels of 12,13-diHOME and improved cardiac haemodynamics⁹³. These findings suggest that a sustained increase in plasma levels of 12,13-diHOME preserves cardiac function and remodelling and increases cardiac haemodynamics through a direct effect on the cardiomyocyte. These findings were reinforced by observations in humans, in whom the presence of cardiovascular disease is associated with decreased plasma levels of 12,13-diHOME⁹³.

Interestingly, skeletal muscle can influence the adipose tissue response to exercise via lactate secretion. The prototypical example is transforming growth factor- β 2 (TGF β 2)⁴². In a mouse model, specific lactate exposure *in vitro* and *in vivo* increased adipocyte expression of TGF β 2 (REF.⁴²). Furthermore, the same study found that in a mouse model of chronic exercise, increased adipocyte levels of TGF β 2 expression and secretion were associated with improvements in glucose metabolism, lipid oxidation and a possible reduction of adipose tissue inflammation. Parallel findings were also observed in humans undertaking chronic exercise, albeit to a less pronounced degree than the animal model observations⁴². Nevertheless, these findings demonstrate the possibility of lactate mediating tissue-to-tissue communication during exercise.

Skeletal muscle

Exerkinins originating from multiple tissues have demonstrated the capacity to improve skeletal muscle function and growth (TABLE 3; Supplementary Table 1). Apelin is an example of a myokine affecting muscle function. In both humans and animal models, exercise increases muscle mRNA levels of apelin⁹⁴ and possibly serum levels of apelin^{95,96}. In an animal model, skeletal muscle^{95,96} served as a source of apelin secretion, which improved skeletal muscle function in the setting of ageing, supporting the potential of apelin as a therapeutic to combat age-related sarcopenia^{94–96}. Specifically in old mice, increased apelin exposure (by daily injection or skeletal muscle overexpression) stimulated muscle mitochondrial biogenesis, muscle protein synthesis and enhancement of muscle stem cells to stimulate muscle regeneration⁹⁶. 12,13-diHOME is an example of a batokine with muscle effects. In both humans and mouse models, exercise facilitates BAT secretion of 12,13-diHOME, which enhances skeletal muscle FFA uptake and oxidation⁹¹. The hepatokines follistatin and fetuin-A also affect muscle function. For example, in both humans^{97–99} and mouse models¹⁰⁰, acute exercise^{97,100} and chronic exercise⁹⁸ increase liver-secreted follistatin, which has been reported to antagonize the effects of myostatin⁹⁹. Decreased function of myostatin enhances skeletal muscle growth and improves whole-body glycaemic control^{44,99}. Furthermore,

fetuin-A worsens peripheral insulin resistance by reducing insulin signalling and glucose transporter type 4 trafficking³⁹. Although acute exercise in humans does not alter plasma levels of fetuin-A⁹⁷, chronic exercise might decrease plasma levels of fetuin-A^{72,101}. Additional exerkinins involved in muscle growth and development include the following: IL-7 (REF.¹⁰²), IL-15 (REF.⁹⁹), follistatin⁹⁷, leukaemia inhibitory factor¹⁰³, syndecan 4 (REF.¹⁰⁴) and myostatin^{44,73}.

The liver and the gut

Exercise reduces hepatic steatosis independently of weight loss¹⁰⁵. The liver is recognized as a source for many circulating proteins, with ~2,500 liver-secreted proteins identified using modern liquid chromatography and mass spectroscopy technologies³⁹. Not surprisingly, the liver is the source of many acute exercise-responsive cytokines (Supplementary Table 1). These exerkinins affect glucose and/or lipid metabolism (for example, angiopoietin-like protein 4 in humans and animal models^{39,79}), browning of WAT (FGF21 in a mouse model¹⁰⁶), lipolysis (FGF21 in humans and a mouse model⁷⁷) and the maintenance of cellular homeostasis (heat shock protein 72 in humans¹⁰⁷).

Exercise also alters the gut microbiome¹⁰⁸. Chronic exercise in humans and animal models alters the composition and functional capacity of the gut microbiota, independently of diet; these exercise-dependent changes in the microbiota might be independent of weight while being contingent on exercise intensity, modality and sustainment¹⁰⁸. In humans, chronic exercise altered the gut microbiome to increase availability of short chain fatty acids, particularly butyrate¹⁰⁹. Once these participants ceased training, exercise-induced changes in the microbiota were largely reversed when re-measured after a 6-week sedentary period¹⁰⁹. The mechanisms by which exercise might alter the gut microbiome remain numerous, including altering the gene expression of intraepithelial lymphocytes for a more favourable inflammatory profile¹¹⁰, influencing blood flow in the gut¹¹¹ or changing bile acid excretion¹¹².

The endocrine system

As exercise has established benefits in improving dysglycaemia, this section focuses specifically on exerkinins affecting glucose homeostasis (TABLE 3; Supplementary Table 1). In humans, circulating levels of β -aminoisobutyric acid (BAIBA) increased with chronic training⁵⁵ and inversely correlated with insulin resistance⁵⁵. In wild-type mice, BAIBA treatment reduced insulin resistance and suppressed inflammation⁵⁵. In another study using C2C12 mouse myocytes and a wild-type mouse model (palmitate or high-fat diet exposure), BAIBA treatment attenuated insulin resistance, reduced inflammation and increased fatty acid oxidation through AMP-activated protein kinase (AMPK) and a AMPK–PPAR δ -dependent pathway in skeletal muscle¹¹³. Limited human data show that an acute exercise bout increases plasma and muscle expression levels of fractalkine (encoded by *CX3CL1*)⁷⁰, which is a chemokine that favourably regulates glucose-stimulated insulin secretion by enhancing

β -cell function¹¹⁴. Chronic exercise in humans also reduces circulating levels of fetulin-A^{72,101}. Fetulin-A has been shown to impair β -cell sensing by reducing glucose-stimulated insulin secretion¹¹⁵.

In humans, both acute exercise⁹⁷ and chronic exercise⁹⁸ increase circulating levels of follistatin. The extent to which follistatin might improve glycaemic measures remains controversial. After bariatric surgery, improvements in HbA_{1c} have been observed in the setting of reduced levels of follistatin¹¹⁶. Furthermore, inactivating hepatic follistatin in a mouse model improved WAT sensitivity and reduced hepatic glucose production¹¹⁶. In vitro, irisin prevented excessive lipogenesis of mouse islets under glucolipotoxic conditions, resulting in improved insulin secretion, inhibition of apoptosis and restored β -cell function-related gene expression¹¹⁷.

The myokine IL-6 is also associated with favourable alterations in glucose homeostasis. In humans, IL-6 infusion delays gastric emptying and lowers postprandial glucose levels¹¹⁸. In rodents, increasing IL-6 by exercise or by IL-6 injection increases the production of glucagon-like peptide 1 by intestinal L cells and pancreatic α -cells to enhance glucose-stimulated insulin secretion. These benefits of IL-6 in enhancing insulin secretion were seen across multiple rodent models of T2DM¹¹⁹. In healthy humans, IL-6 infusion to levels similar to those seen with strenuous exercise enhances insulin-stimulated glucose-uptake but does not alter whole-body lipolysis or lipid oxidation¹²⁰. However, another study of IL-6 infusion into humans found that IL-6 stimulates lipolysis and lipid oxidation^{54,121}. Further research into these seemingly conflicting findings is warranted to establish the effect of exerkinins on glucose metabolism.

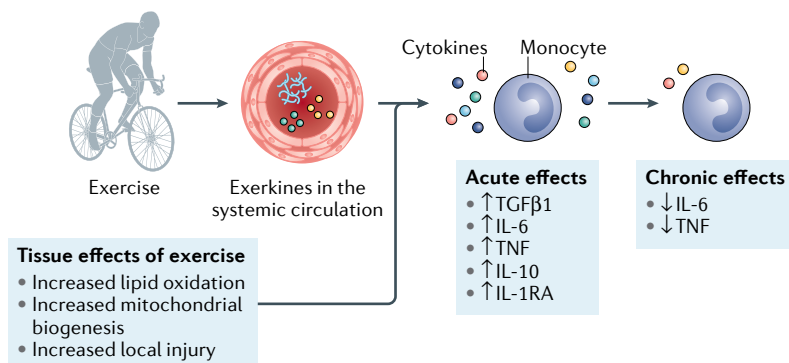


Fig. 3 | Effects of exercise on the immune system. Exercise induces lipid oxidation, mitochondrial biogenesis and local injury, which stimulates exerkinine release into the circulation to influence the immune system. See Supplementary Table 1 for detailed effects and relevant references. These include proteins (blue lines), metabolites (yellow circles) and extracellular vesicles (green circles), which have a multitude of effects on the immune system (generically represented by a monocyte). Acutely, exercise increases cytokines such as circulating levels of transforming growth factor β 1 (TGF β 1) and IL-6 relative to the resting state. This change results in acute inflammation, characterized by increases in tumour necrosis factor (TNF) and IL-6. Once the acute exercise-induced effects have diminished, an increase in anti-inflammatory cytokines (such as IL-10 and IL-1 receptor antagonist (IL-1RA)) occurs in response to the acute inflammatory response. Chronic training is associated with a reduction in systemic and tissue inflammation, as characterized by lower circulating levels of TNF and IL-6 in the resting state, relative to sedentary individuals. Reduced insulin resistance and tumour growth has been attributed to the effects of chronic training on decreasing systemic and/or tissue inflammation.

The immune system

The broad effects of exercise on immune function implicate mobilization and altered function of cytokines and immune cells, such as neutrophils, leukocytes and natural killer cells (FIG. 3; Supplementary Table 1)^{30,122,123}. The effects of chronic exercise on the immune system might depend on intensity, with immune enhancement by moderate exercise and possible impairment by strenuous exercise¹²⁴. As shown in humans, an acute bout of exercise might be initially pro-inflammatory, but subsequently this effect is offset by an anti-inflammatory response^{30,124}. The exercise-induced acute increase in circulating levels of IL-6 increases plasma levels of anti-inflammatory cytokines, such as IL-1RA and IL-10 (REF.¹²⁵). IL-1RA inhibits IL-1 β signal transduction¹²⁶, whereas IL-10 inhibits production of pro-inflammatory cytokines, such as TNF¹²⁷. In healthy humans, one bout of exercise or an IL-6 infusion blunted the increase in circulating levels of TNF induced by infusion of lipopolysaccharide¹²⁸. Thus, an acute bout of exercise induces anti-inflammatory effects that might in part be mediated by IL-6, possibly in conjunction with other known anti-inflammatory factors, such as adrenaline and cortisol¹²². As a pleiotropic factor, the effect of IL-6 on metabolism and inflammation remains context-dependent. Although IL-6 is transiently increased after acute exercise, the baseline (or ‘resting’) circulating levels of IL-6 are lower in exercise-trained individuals than in untrained individuals⁷⁸. Future studies focusing on tissue and cell type-specific effects as well as different exercise regimens will help delineate the temporal and spatial requirement of IL-6 in mediating exercise benefits.

An emerging frontier in exercise biology involves exerkinine-induced immune effects in increasing resilience to cancer or as co-adjuvant to cancer therapy. The anticancer effects of exercise might not be limited to its effect on body weight. A meta-analysis pooled data from 12 prospective cohorts with self-reported physical activity and found that increased physical activity levels are associated with decreased risk of incident cancer across multiple types; many of these associations remained even after adjusting for BMI¹²⁹. Acute exercise creates a unique exerkinine milieu that lasts several hours after exercise cessation, which provides a temporal window for immune function stimulation³¹. For this reason, exercise could potentially serve as a co-adjuvant treatment for cancer therapy. In tumour-bearing mouse models (across five tumour models), mice that undertook voluntary wheel running had a reduction of more than 60% in tumour incidence and tumour growth compared with sedentary mice. Further analysis of these mouse models showed that adrenaline and IL-6 induced natural killer cell mobilization, redistribution and tumour infiltration to inhibit tumour growth¹²³. Another mouse model found that exercise metabolites such as lactate and possibly tricarboxylic acid intermediates enhance the antitumour effector profile of CD8⁺ lymphocytes¹³⁰. Of note, exercise is associated with enhanced secreted protein acidic and rich in cysteine (SPARC) secretion in humans and animal models¹³¹; this extracellular protein regulates cell function and tissue remodelling,

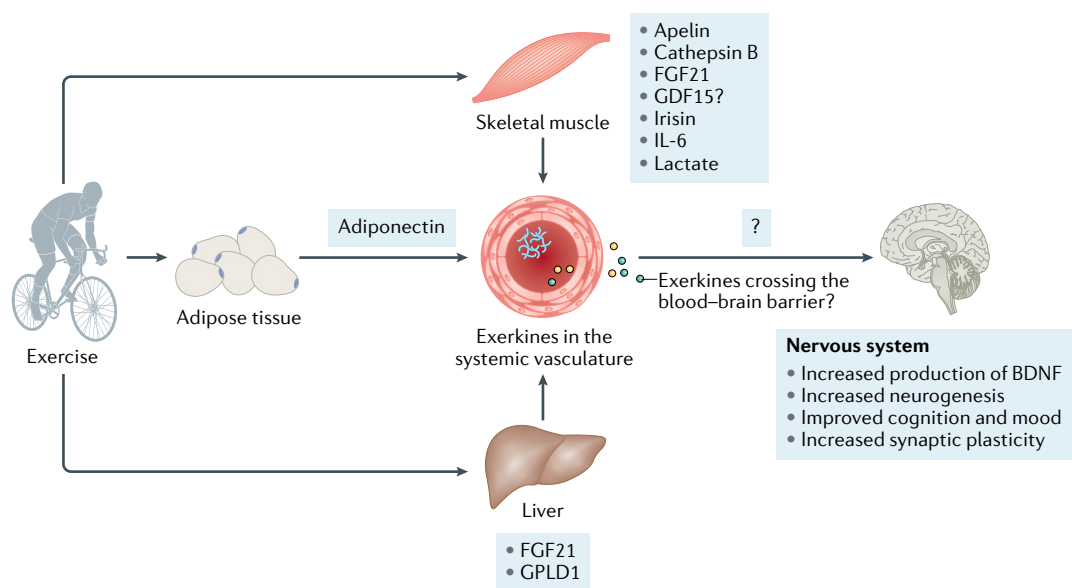


Fig. 4 | Effects of exercise on the nervous system. Exercise stimulates the production of exerkines from tissues, such as skeletal muscle, adipose tissue or the liver, to affect the nervous system. See Supplementary Table 1 for detailed effects and relevant references. These exerkines are released into the circulation and include proteins (blue lines), metabolites (yellow circles) and extracellular vesicles (green circles), which have a multitude of purported effects on the nervous system. These effects include increasing production of brain-derived neurotrophic factor (BDNF), enhancing neurogenesis (even in adults), cognition, mood and synaptic plasticity. The extent to which exerkines cross the blood–brain barrier to exert their effects remains unknown, symbolized by the question mark. There is uncertainty with GDF15, as symbolized by the question mark, as pharmacological GDF15 inhibits appetite and reduces activity, whereas physiological induction of GDF15 by exercise does not²⁰⁰. Commonly described exerkines are noted. FGF21, fibroblast growth factors 21; GDF15, growth and differentiation factor 15; GPLD1, glycosylphosphatidylinositol-specific phospholipase D1.

while inhibiting proliferation and promoting apoptosis of a mouse colon cancer cell line¹³².

Crosstalk exists between skeletal muscle and the immune system. Contemporary views towards skeletal muscle now consider muscle as an immunoregulatory organ that especially affects lymphocyte and neutrophil trafficking and inflammation. During acute exercise, immune cells are mobilized by muscle secretion of exerkines such as fractalkine to enhance regeneration (in humans)¹³³ or meteorin-like (METRNL) to increase beige adipose tissue thermogenesis (animal model)¹³⁴. As previously noted, one bout of exercise or an IL-6 infusion in humans blunts the increase in circulating levels of TNF that are induced by lipopolysaccharide infusion¹²⁸.

IL-13 is an important main T helper 2 (T_H2) cell cytokine that mediates the anti-inflammatory polarization of resident macrophages in WAT¹³⁵. IL-13 is also an exerkine, increasing in the circulation after exercise training in humans and mice¹³⁶. Mice lacking IL-13 show reduced running capacity and do not show certain beneficial effects of exercise training, such as improvements in glucose tolerance and endurance running capacity¹³⁶. Unlike IL-6, IL-13 is produced by type 2 innate lymphoid cells (ILC2s) in skeletal muscle. As IL-13-deficient mice show defective muscle fatty acid utilization after acute exercise and fail to show increased muscle mitochondrial biogenesis after chronic exercise, the ILC2 to IL-13 axis might have an important role in the metabolic adaptation to exercise training¹³⁶. Interestingly, ILC2 and T_H2 cell cytokines also control beige adipocyte recruitment in

rodents¹³⁷, suggesting that ILC2 to T_H2 signalling might partially mediate exercise-induced beiging of WAT. The stimulants within skeletal muscle or WAT that activate ILC2s during exercise remain to be identified.

The nervous system

Exercise is a promising non-pharmacological strategy to maintain and improve brain function¹³⁸. FIGURE 4 presents an overview of the purported effects of exercise on the nervous system (Supplementary Table 1). Of note, evidence for the benefits of exercise on cognition remains variable, probably owing to the lack of randomized controlled trials throughout the lifespan with standardized exercise interventions and comparable methods for cognitive assessment^{139–142}. The effects of exercise on the brain are most apparent in the hippocampus, a part of the brain involved in learning and memory¹⁴³. In older adults (aged 55–80 years), participation in an aerobic walking programme increased hippocampal volume and improved memory¹⁴⁴. Moreover, accumulating evidence suggests that physical activity, as shown in preclinical, observational and interventional studies in humans, can prevent or delay the onset of neurodegenerative conditions¹³⁸. In humans, acute exercise increases plasma levels of brain-derived neurotrophic factor (BDNF)⁷, whereas chronic exercise training has been shown to not alter¹⁴⁰, increase or even decrease plasma levels of BDNF^{140,145}. In rodents, chronic exercise upregulates BDNF in the hippocampus, which is essential for adult hippocampal neurogenesis and neural plasticity¹⁴⁶. Chronic exercise in rodents also enhances

hippocampal synaptic plasticity, adult neurogenesis and neurotrophin levels, as well as memory function¹⁴⁷. In addition, voluntary wheel running in rodents increases the number of new hippocampal neurons, enhances morphological maturation, such as dendritic branching and spine density, and alters the circuitry of adult-born neurons¹⁴⁷.

There is increasing recognition that peripheral factors might trigger the effects of exercise on the brain. In the past 20 years, researchers have begun to test the hypothesis that metabolites, peptides and proteins released from liver, adipocytes, blood cells (particularly platelets) and muscle might influence the central nervous system^{138,148,149}. Since 2020, several studies have transferred plasma from exercised animals into sedentary animals, with subsequent improvements in cognitive function, supporting the presence of a transferable factor in improving cognitive function^{148,149}. Evidence is now accumulating that factors released from non-neuronal tissue^{148–152} and delivered via the vasculature to the brain might have important roles in synaptic plasticity, memory function and mood regulation¹⁵⁰. Adiponectin is an adipocyte-secreted protein that seems to have neuroprotective effects¹⁵¹, in addition to its insulin-sensitizing, anti-inflammatory and anti-atherogenic effects^{151,153}. In mice, adiponectin was demonstrated to pass through the blood–brain barrier and was associated with increased neurogenesis and reduced depression-like behaviours¹⁵¹. Interestingly, discrepancies can exist between the plasma levels of adiponectin and levels in the cerebrospinal fluid. For example, in humans, acute exercise increases plasma levels of adiponectin but decreases cerebrospinal fluid levels of adiponectin¹⁵⁴. The mechanisms underlying the beneficial effects of exercise on brain structure and function remain an active area of investigation.

Muscle–brain crosstalk. Myokines seem to have an important role in hippocampal neurogenesis (animal model) and neurotrophin levels (animal model), and enhanced cognition and mood (animal model and humans) (Supplementary Table 1). For instance, in mice, increasing intrinsic irisin expression in neurons or increasing plasma levels of irisin elevates hippocampal *Bdnf* gene expression¹⁵². Irisin administration in a mouse model of Alzheimer disease improves synaptic plasticity and memory function^{155,156}. In both animal models and humans, plasma levels of cathepsin B, a lysosomal thiol proteinase produced by muscle, is positively associated with hippocampus-dependent memory^{145,157}. Studies in humans have shown that acute exercise does not clearly increase plasma levels of cathepsin B⁷, although chronic exercise does increase plasma levels of cathepsin B^{145,157}. These same studies also showed that acute exercise increases plasma levels of BDNF⁷, whereas chronic exercise does not increase BDNF^{140,145} suggesting that investigation of local neuronal effects remains warranted.

Liver–brain crosstalk. The liver secretes factors that are important for brain function. Kynurenine is a metabolite of the amino acid L-tryptophan and is

primarily synthesized in the liver. In both mice and humans, chronic aerobic training increases muscle expression of kynurenine aminotransferase, which facilitates conversion of kynurenine into kynurenic acid, a metabolite that is unable to cross the blood–brain barrier. This shift in kynurenine metabolism is able to protect the brain from stress-induced depression¹⁵⁰.

Exchanging plasma from exercising aged mice to sedentary aged mice enhances adult hippocampal neurogenesis and memory function¹⁴⁸. Upon further investigation, plasma proteomic analysis led to the identification of a novel hepatokine, glycosylphosphatidylinositol-specific phospholipase D1 (GPLD1), which increases after exercise and correlates with improved cognitive function in aged mice. These findings are supported in humans, as concentrations of GPLD1 in blood were higher in active, healthy older adults ($n=20$, >66 years old) than in their sedentary counterparts¹⁴⁸. Investigations into the underlying mechanisms indicate that GPLD1 does not cross the blood–brain barrier¹⁴⁸. Hence, the benefit of GPLD1 (REF.¹⁴⁸) and other exerkinins on brain structure and function might relate to its peripheral effects, such as the complement signalling cascade^{149,158}, or coagulation¹⁵⁹. Additional exercise studies are needed to better appreciate the exercise–liver–brain axis.

Bone

Exercise, especially resistance exercise, increases bone mineral density¹⁶⁰. Multiple mechanisms exist, although mechanical loading is considered a major factor¹⁶¹. Noted exercise-associated bone-derived factors affecting bone formation include TGF β 1 (REF.¹⁶²) and sclerostin¹⁶³. Sclerostin inhibits bone formation^{163,164} and blood levels of sclerostin are lower in highly active humans than in sedentary humans¹⁶³. Emerging data demonstrate that crosstalk exists between bone and muscle, probably mediated by secretory factors¹⁶⁵. Noted myokines affecting the bone include apelin¹⁶⁶, myostatin⁶¹, irisin¹⁶⁷, IL-6 (REF.¹⁶⁸), IL-7 (REF.⁶⁰) and BAIBA⁵⁹ (Supplementary Table 1).

Gaps and future opportunities

Gaps in exerkin science

Contentious questions remain that temper the enthusiasm regarding exerkinins (BOX 1). These controversies include the lack of consistency between the acute and chronic exercise response, discrepancies between humans and animal models of exercise and interpretation challenges due to variability in outcomes and sampling. These knowledge gaps set the stage for future opportunities in exerkin research.

Despite the acceleration in exerkin-related research since the identification of IL-6 as a myokine in 2000 (REF.¹⁶), much remains to be done in the scientific areas of research, technology and therapeutic interventions (BOX 1). Specifically, a critical need exists to move beyond the ‘skeletal muscle-centric’ view of exerkinins and focus more on their roles in inter-organ communication, tissue regeneration, immune regulation, metabolic adaptation, cardiovascular fitness, psychological health and overall health across the lifespan. The vasculature and endothelium are emerging as a probable central facilitator, which enables systemic exerkinins to exert their

Box 1 | Contentious questions regarding exerkinetics

Overenthusiasm about the potential of exerkinetics needs to be tempered, as controversies remain.

Inconsistency between the acute and chronic exerkinetic response

As demonstrated in Supplementary Table 1, the acute exerkinetic response does not necessarily parallel and could even contradict the chronic exercise response, as exemplified by brain-derived neurotrophic factor (BDNF), IL-6, irisin and myostatin. Hence, the question arises as to whether the discrepancy reflects the difference in mechanism between acute versus chronic perturbation, or more pragmatically, remains unrelated to the observed exercise-induced changes.

Inconsistency between studies in animals and humans

As demonstrated in Supplementary Table 1, the exerkinetic response in animals does not necessarily parallel the exerkinetic response in humans, as exemplified by angiotensin, BDNF, follistatin, IL-7, IL-8, IL-10, irisin, leukaemia inhibitory factor, myostatin, myostatin and secreted protein acidic and rich in cysteine (SPARC). Hence, the question arises as to whether the excitement generated from discoveries in animal models will translate to specific human populations or more pragmatically, whether animal findings remain unrelated to observed changes in humans.

Variability of outcomes and sampling, which hinder interpretation

The literature is replete with studies showing variable, if not conflicting, benefits from exercise, commonly attributed to differences in selected populations, exercise type, exercise intensity and exercise duration^{141,142,201}. However, even when an exercise exposure is fixed, as exemplified by a clinical trial setting, the cardiovascular response can be ostensibly disassociated from the metabolic response^{25,26,202}. Hence, the question arises as to whether these discrepancies can be explained by differences in the exerkinetic response. If alterations in exerkinetics might explain the observed findings, even more questions arise regarding sampling relative to the timing of exercise exposure (during versus after), sampled tissue (soft tissue versus blood) and context (fasted or fed²², or circadian rhythm^{21,24}). Hence, the question arises as to the translational relevance of the exerkinetic literature to date, given the inconsistency of observed exercise effects and highly variable sampling protocols.

specific effects within the various local environments, as well as being a direct target and source of exerkinetics. Understanding the system-wide effects of exercise and the myriad of exercise-related improvements is essential for understanding resilience. Such knowledge will provide new translational research opportunities to develop novel, targeted interventions that increase physiological reserve, maintain and/or enhance resilience and thus promote healthy ageing, as well as interventions that prevent and treat comorbidities and chronic disease.

Many more exerkinetics, and their sources, targets and mechanisms, remain to be discovered. In 2016, the MoTrPAC project was launched to uncover novel exerkinetics through deep omics profiling of biomaterial from humans and rodents before and after acute exercise as well as chronic exercise training. As potential candidate transducers are uncovered, follow-up mechanistic studies will be required to delineate their function. In addition to molecular discoveries, substantial work remains to decipher the dosage and type of exercise needed to elicit positive health outcomes. Intervention studies are needed to investigate the effect of different types of exercise on resilience to various conditions, with guidance available from the NIH in designing resilience-based studies. To address these knowledge gaps, detailed studies are needed to identify and validate the exerkinetic responses after exercise (acute, chronic or intermittent) exposure, including a detailed post-exercise response. Certainly, the demographics and

phenotypes of the population will matter, as the response in healthy participants can vary greatly from that in participants with comorbidities. Further work in these areas will advance our understanding of ageing, health and disease prevention.

Opportunities for new technologies

Technology gaps also remain in exerkinetic research. One emerging area is the use of wearable technologies and devices to capture quantitative and dynamic phenotypes over long periods of time in healthy individuals as well as in those with mild or severe diseases. For instance, wearable technology could provide valuable information regarding physical activity levels and exercise capacity during and following COVID-19 illness and recovery, adding to the description of post-acute sequelae of SARS-CoV-2 infection, which is under active investigation. Currently, non-invasive and minimally invasive devices have enabled the monitoring of many behavioural and physiological phenotypes, including heart rate and electrical activity in the heart (ECG), body temperature, physical activity and sedentary behaviour, peripheral blood oxygen saturation and blood concentrations of glucose. These devices enable the real-time monitoring of the effects of exercise in natural and controlled settings at an unprecedented level. Moreover, wearable technologies can be scaled to the analysis of over a million people and can enable 'citizen science' whereby individuals with devices can readily participate in studies. These measurements, when combined with molecular measurements such as exerkinetics, have the potential to greatly improve our understanding of exercise adaptations in large cohorts with deep phenotyping across a broad age range.

Although wearable devices are powerful, multiple challenges remain in data interpretation and analysis. These challenges include device accuracy as well as device standardization and a lack of readily available high-resolution data from the manufacturers. In addition, many wearable devices do not characterize the environmental context associated with data capture. For example, if a device does not sense any physical activity, one explanation could be the lack of movement by the wearer while another explanation could be device removal. A remaining challenge entails approval and regulation from entities such as the FDA before widespread use of wearables as therapeutic interventions.

In addition to wearables, the technology, analysis and approach for exerkinetic discovery needs to be further developed. Deep omics profiling (transcriptome, metabolome, proteome and lipidome) of human and animal-based exerkinetics is occurring in MoTrPAC; this effort is expected to reveal novel molecules and mechanisms involved in exercise by providing a more comprehensive view of the multi-omics landscape of exerkinetics to the research community. Extracellular vesicle analysis, especially of exosomes, will be a critical component of MoTrPAC's analysis owing to burgeoning interest in the role of extracellular vesicles as carriers of molecular signals and drivers of inter-organ crosstalk.

Exosomes

Exosomes are a type of extracellular vesicle released by parent cells, which contain RNAs, proteins and lipids, to facilitate crosstalk between tissues.

Data reporting and data sharing

To promote comparison between studies and enhance translation, a crucial need exists for the establishment of community-wide standards for data reporting and data sharing. As an example, capturing physical activity and body composition (for example, adipose tissue mass) in the electronic health record will facilitate electronic health record data mining to examine clinical outcomes outside a structured clinical study. Cross-study comparisons of exercise studies will be facilitated by setting standards for a minimum metadata set, for consistent documentation and of covariates, such as time of day, diet or exercise exposure. Advances in computational modelling will accelerate our understanding of the physiological process of exercise and exerkin effects. Establishing a uniform knowledge base remains a critical next step in driving this process.

Exerkines as therapeutics

The health benefits of exercise are well documented. However, not all individuals are able to benefit from exercise owing to physical limitations, such as paraplegia, or imposed limitations, such as quarantining during the current COVID-19 pandemic. Moreover, in humans, metabolic non-response or even adverse responses to exercise have been described²⁶. As the role of exerkines and their biological effects are increasingly clarified, exerkines could potentially be harnessed to mimic the benefits of exercise in individuals who are limited in their exercise capacity or to counterbalance a metabolic

non-response or adverse response to exercise. Although this ‘exercise in a pill’ is currently wishful thinking, it remains a tantalizing goal for future research directions.

Conclusions

Although exercise exerts many beneficial effects across multiple organ systems, our understanding of the mechanisms driving the benefits of exercise and the variability in these benefits remains rudimentary. Much of the initial exerkine research has been focused on skeletal muscle; however, contemporary research is now rapidly expanding to include non-skeletal muscle-based sources and targets for exerkines that contribute to maintaining and restoring health. Exerkines are increasingly recognized as critical mediators of exercise-related changes and health benefits, particularly in their role in inter-organ and systemic communication and coordination. Yet, much remains to be done. To improve translation of results, the heterogeneity across studies needs to be minimized by reducing exposure variability and using standardized, consistent outcome measures. Large-scale, structured studies will be key resources in providing a structured environment to pursue future exerkine-related questions. In summary, exerkines are a highly promising direction for future research initiatives, with high potential as biomarkers to predict outcomes, facilitate personalized exercise programmes to improve health, reduce disease and promote resilience across the lifespan.

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Author contributions

The authors contributed equally to all aspects of the article.

Competing interests

L.S.C. has received a Dexcom investigator initiated grant (product only). C.J.L. is a consultant for PALhealth on their Personalized Activity Intelligence applications. A.P. is on the advisory board of Roche Diagnostics. M.P.S. is a cofounder and scientific advisory board member of Personalis, SensOmics, Qbio, January, Filtricine, Protos, NiMo, Mirvie, and an advisor for Genapsys. The other authors declare no competing interests.

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