Suppressive effects of exercise-conditioned serum on cancer cells: A narrative review of the influence of exercise mode, volume, and intensity

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Highlights:

- Exercise-conditioned serum has inhibitory effects on cancer cell growth and promotes apoptosis in different cancer cell lines.
- A single bout of moderate to vigorous intensity aerobic training may have suppressive effects on cancer cells and elicit substantial increases in circulating factors.
- Training volume may play a role in inhibiting cancer cells and altering circulating factors, regardless of the training mode used, when examining short-term and chronic training interventions.
- Future research should be directed to targeted investigations into the effects of exercise-conditioned serum on cancer cells, based on specific training modes, volume, and intensity stratifying by treatments or disease stage.

Review

Suppressive effects of exercise-conditioned serum on cancer cells: A narrative review of the

influence of exercise mode, volume, and intensity

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Running head: Suppressive exercise-effects on cancer cells

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Abstract

Cancer is a major cause of morbidity and mortality worldwide, and the incidence is increasing,

highlighting the need for effective strategies to treat this disease. Exercise has emerged as

fundamental therapeutic medicine in the management of cancer, associated with a lower risk of

recurrence and increased survival. Several avenues of research demonstrate reduction in growth,

proliferation, and increased apoptosis of cancer cells, including breast, prostate, colorectal, and

lung cancer, when cultured by serum collected after exercise in vitro (i.e., the cultivation of cancer

cell lines in an experimental setting, which simplifies the biological system and provides mechanistic insight into cell responses). The underlying mechanisms of exercise-induced cancer suppressive effects may be attributed to the alteration in circulating factors, such as skeletal muscle-induced cytokines (i.e., myokines) and hormones. However, exercise-induced tumor suppressive effects and detailed information about training interventions are not well investigated, constraining more precise application of exercise medicine within clinical oncology. To date, it remains unclear what role different training modes (i.e., resistance and aerobic training) as well as volume and intensity have on exercise-conditioned serum and its effects on cancer cells. Nevertheless, the available evidence is that a single bout of aerobic training at moderate to vigorous intensity has cancer suppressive effects, while for chronic training interventions, exercise volume appears to be an influential candidate driving cancer inhibitory effects regardless of training mode. Insights for future research investigating training modes, volume and intensity are provided to further our understanding of the effects of exercise-conditioned serum on cancer cells.

Keywords: Cancer cells; High intensity interval training; Moderate intensity continuous training; Resistance training; Myokines

1. Introduction

The importance of exercise for promoting general wellbeing, maintaining good health, and even as a form of medical treatment has been an increasingly popular topic of research over

the past decades. In recognition of the effects of exercise, the World Health Organization (WHO) has provided exercise recommendations specifically for the management of noncommunicable diseases (NCDs), including cardiovascular disease, chronic respiratory disease, type 2 diabetes mellitus (T2DM), and cancer. The WHO expert panel recommends at least 150–300 or 75–150 min of moderate to vigorous aerobic activity per week respectively, along with strengthening exercises at moderate to high intensity twice per week, to prevent the onset of NCDs. Among these diseases, cancer represents a significant contributor to both morbidity and mortality worldwide, with breast, lung, colorectal, and prostate cancer being the leading causes of cancer-

related deaths (10 million deaths in 2020).⁵ Unfortunately, the incidence of cancer is increasing worldwide, highlighting the need for effective strategies to prevent and treat this disease.⁵

Exercise has emerged as a novel and fundamental therapeutic intervention in the management of cancer, as supported by multiple studies.^{6–9} Indeed, a robust body of evidence now exists on the safety and effectiveness of exercise as medicine, either during or post-cancer treatments, to improve various health-related cancer outcomes, such as fatigue, quality of life, cardiorespiratory capacity, neuromuscular strength, physical function, body weight, and body composition (both fat mass (FM) and lean mass (LM)), as well as the ability to alleviate symptoms of anxiety and depression. 10-14 Furthermore, exercise is associated with a lower risk for the development of a range of cancers as well as reduced recurrence and improved survival in patients with cancer. 15-¹⁷ The underlying mechanisms are not yet fully understood; however, over the past two decades, several avenues of preclinical and clinical research have investigated the effects of acute and chronic exercise-conditioned serum on different cancer cell lines in vitro (e.g., breast, prostate, and colon). Briefly, this novel methodology involving direct application of the serum obtained from human subjects/patients before and after exercise to a cultured-controlled environment with cancer cell lines (in vitro experiments) revealed that exercise-conditioned serum had inhibitory effects on cancer cell growth (i.e., proliferation and metastatic capacity) and survival (i.e., viability), as well as increased cell death (i.e., apoptosis and necrosis). 18-22 This methodology may partially provide the underlying biological reasons for the association between exercise and reduced recurrence or improved survival in patients with cancer, as well as reveal some of the potential molecular candidates. The underlying reasons come from the fact that previous in vitro studies pointed out how altered circulating factors (e.g., insulin-like growth factor-1 (IGF-1) in obese subjects) may drive cancer cell growth, which was subsequently confirmed in mechanistic

experiments in co-culture systems and, lastly, validated in human subjects.^{23,24} Accordingly, *in vitro* experiments can be employed for therapeutic interventions in the translational continuum from basic discovery to clinical outcomes.

It has been postulated that exercise alters the concentrations of hormones (e.g., insulin, IGF-1), adipokines (e.g., leptin, adiponectin), and cytokines (e.g., tumor necrosis factor alpha (TNF-α)),^{25–28} which are considered to be factors associated with tumorigenesis.^{29–31} Notably, among them, myokines (e.g., interleukin 6 (IL-6), secreted protein rich in cysteine (SPARC)), which are cytokines released into the systemic circulation in response to muscular contractions, have been demonstrated to potentially suppress cancer cell growth, proliferation, and increase apoptosis in different cancer cell lines in several *in vitro* studies.^{32–41} These findings are encouraging, and myokines may be an attractive area for future investigations.

Given the novelty of exercise as medicine, experimental studies have been conducted in both healthy individuals and cancer patients with the aim of collecting blood following both acute and chronic training interventions and culturing it *in vitro* with cancer cells. It has been reported that exercise-conditioned serum suppresses cancer cell growth, proliferation as well as increases apoptosis in breast, prostate, colon and lung cancer. A2-47 Notably, the magnitude of such effects on circulating factors (e.g., myokines) and cancer cells have varied in blood collected immediately after exercise (i.e., acute effects) compared to that taken at rest (i.e., chronic effects) to avoid the arousal effects of exercise. A2,44,45,48 When it comes to the type of exercise undertaken, two distinct modes have been commonly used: resistance training (RT) and aerobic training (AT). The purpose of RT is to elicit muscle hypertrophy and improve muscular strength, whereas AT is to improve cardiorespiratory fitness and metabolic health. AT can be performed as moderate intensity continuous training (MICT) or high intensity interval training (HIIT). Although these training modes have different physiological pathways, both have been commonly used in exercise oncology.

However, it remains unclear which type of training mode (i.e., RT vs. AT), along with associated volume and intensity, is more effective in altering circulating factors (i.e., hormones, cytokines, and myokines), which then have suppressive effects on cancer cells. Thus, the aim of the current review was to critically examine which training mode, volume, and intensity have been adopted in different populations to alter blood contents, which in turn have suppressive effects on cancer cells. To the best of our knowledge, no reviews have been

conducted to critically examine such training parameters. Additionally, we present directions for future research that may provide critical insight into the practice of exercise medicine for patients with cancer.

2. Methods

Empirical research studies and review journal articles were retrieved from electronic databases of MEDLINE, PubMed, Scopus, and CINHAL databases. Of note, our search strategy did not exclude previously published review articles (like many systematic reviews do) because our goal was to conduct a narrative review on a substantially under-researched topic. Thus, review articles served as a useful source for any articles that were not found in our initial searches. The search

strategy combined specific terms with the words: "cancer", "serum", and "exercise" to ensure relevant articles were extracted. The search terms included: "cancer OR tumor OR neoplasm AND exercise OR physical activity OR resistance training OR aerobic training AND serum AND growth OR proliferation OR viability OR apoptosis". Articles were deemed relevant after scanning the title and abstract and where subsequent access to the full text was available from the relevant publishers. The reference lists of each study were also checked to ensure no further articles were omitted from the search process. All searches were conducted between February and April 2023.

As a result, only studies in breast, prostate, colorectal, and lung cancer conducted in subjects with and without cancer were found. Such studies examined the effects of the exercise-conditioned serum on cancer cells and are discussed in the following sections followed by directions for future research. It should be noted that all cancer patients in the reviewed studies were either undergoing primary cancer treatments (e.g., chemotherapy, hormone therapy, radiotherapy) or were in the survivorship phase, during which exercise was applied as adjuvant therapy.

3. Breast cancer

Breast cancer is the most diagnosed cancer worldwide;^{5,54} thus, several studies have been conducted to examine the effects of circulating factors (e.g., myokines) on breast cancer cells in preclinical studies, and with promising results.^{39–41,55–57} Recently, studies have been conducted in both healthy individuals and patients with breast cancer. However, and

somewhat surprisingly, there remains a lack of studies investigating the effects of exercise-conditioned serum on breast cancer cells.

3.1. Studies in subjects without breast cancer

The effects of exercise-conditioned serum on breast cancer cells have been investigated in subjects without cancer. For example, in regard to the acute effects (i.e., single bout of exercise) of exercise-conditioned serum, Dethlefsen et al. 42 examined MICT performed at 55% maximal oxygen uptake (VO_{2max}) for 2 h on a cycle ergometer in 7 healthy women (Table 1). Serum acquired 1 to 2 h post exercise resulted in a significant decrease in the viability (i.e., the ability of cancer cells to survive, grow, and maintain cellular function) of breast cancer cell lines MCF-7 by 10%–19% and MDA-MB-231 by 13%–14% (p < 0.05). Thus, preliminary evidence shows that an acute bout of exercise at 55% VO_{2max}, which corresponds to approximately 70% maximum heart rate (HR_{max}), 58 may

suppress breast cancer cell growth. Similarly, Baldelli et al.⁵⁹ investigated the effects of a high intensity endurance test to exhaustion (i.e., up to 90% maximal power (P_{max})) in 12 healthy women before and after a 9-week HIIT protocol performed 3-4 days a week. They reported that exerciseconditioned serum taken before commencing the HIIT protocol (i.e., acute) inhibited MDA-MB-231 cell proliferation by 12.0%–24.9% in the untrained state (with serum acquired immediately after to 24 h after the endurance test to exhaustion) (p < 0.05). Interestingly, suppressive cancer cell proliferation effects were greater when applying the serum (collected at the same time points mentioned above) after the 9-week HIIT protocol (i.e., MDA-MB-231 cell proliferation decreased by 15.7%–35.3%; p < 0.05). Regarding the circulating factors, the authors examined only levels of creatine kinase, finding a transient increase immediately after exercise. The high intensity (i.e., 90% P_{max}) appears to mediate the inhibitory response of breast cancer cells after the incremental test. Additionally, suppressive cancer effects were even greater after a 9-week HIIT protocol, highlighting how the weekly training volume selected increased the capacity of the exerciseconditioned serum to inhibit breast cancer cell proliferation. This supports the contention that exercise training interventions not only have inhibitory effects due to acute exercise bouts but also that the systemic changes that occur with chronic training have tumor suppressive effects.

Barnard et al.²² investigated a short-term diet and exercise program in 38 overweight and obese post-menopausal women (Table 2). Participants received a low-fat and high-fiber diet in addition to 30–60 min MICT (i.e., treadmill) at 70%–85% HR_{max} 4–5 days a week and 40–

60 min < 70% HR_{max} 1–2 days a week for 2 weeks. Resting serum levels of IGF-1 (-19%), IGF-binding protein-1 (IGFBP-1) (+32%), insulin (-29%), and oestradiol (-37% to -34%) significantly improved after two weeks (p < 0.05). When examining the effects on breast cancer cell lines, the exercise-conditioned serum significantly reduced *in vitro* MCF-7 cell growth by 6.6%, ZR-75-1 by 9.9%, and T-47D by 18.5%, as well as increased apoptosis of the breast cancer cell lines MCF-7 by 23%, ZR-75-1 by 20%, and T-47D by 30% (p < 0.05). Interestingly, blood was collected in a resting state, showing that even a short-term training program had positive effects on hormones and cancer cell growth. This may be related to the high volume of exercise used (i.e., >5 days per week), even though it is unknown which component (i.e., diet or exercise) contributed most to these effects.

3.2. Studies in patients with breast cancer

The first acute study in patients with breast cancer was reported by Dethlefsen and colleagues, ⁴³ who examined the effects of a single bout consisting of a 30-min warmup, RT for 60 min, and 30 min of HIIT on a stationary cycle ergometer at 80%–85% HR_{max} in 20 overweight breast cancer patients undergoing chemotherapy half-way through a 6-week supervised training intervention (Table 3). Significant increases in IL-6 (+110%), IL-8 (+20%), TNF- α (+13%), lactate (+500%), epinephrine (+190%), and norepinephrine (+120%) were found in the serum collected immediately after the acute session which, in turn, resulted in significant decreases in MCF-7 and MDA-MB-231 cell viability, by 9.2% and 9.4%, respectively (p < 0.05). This was in accordance with another study led by the same research group using the same cohort where MCF-7 and MDA-MB-231 cell viability significantly decreased, by 11% and 9%, respectively (p < 0.05). ⁴² Notably, the mixed approach (i.e., RT plus HIIT) resulted in significant positive alterations in blood content taken immediately after the acute exercise session. However, it is unclear whether changes were induced by the training volume (i.e., 2 h) and/or intensity (i.e., >70% HR_{max}) or by different training stimuli (i.e., RT vs. HIIT).

In contrast, results differed when Dethlefsen et al.⁴³ investigated the chronic effects of a 6-month training intervention once per week compared to a control group in 74 breast cancer survivors (Table 4). The training intervention comprised RT, including 3 sets of 8–10 repetitions for major muscle groups (e.g., leg press, knee extension, chest press, *etc.*) at 70%–90% 1 repetition maximum (RM), plus HIIT based on work bouts of 30 s to 6 min at 80%–90% HR_{max} on a stationary cycle ergometer. The control group received 3 health evaluation

consultations. Exercise resulted in a significant decrease in resting levels of IL-6 (Intervention = -37%; Control = -20.7%), TNF- α (Intervention = -21.1%), and low-density lipoprotein/high-density lipoprotein (LDL/HDL) (Intervention = -18.2%; Control = -11.4%) after 6 months (p < 0.05), while no changes were observed in IL-8, IL-10, insulin, leptin, or glucose levels. When examining the effects of exercise-conditioned serum on MCF-7 and MDA-MB-231 breast cancer cell lines, no substantial effects were found on cancer cell viability. However, the low volume of training intervention (i.e., 1 day per week) may have been a factor limiting potential adaptations, as highlighted by lack of significant changes in most of the circulating factors (i.e., IL-8, IL-10, insulin, leptin). This was compounded by a training session attendance of only 66%, which is unlikely to be sufficient for maximizing adaptations when training volume is already low.

In summary, it appears that exercise can positively impact circulating factors and that it suppresses cancer cell growth in breast cancer cells. In regard to the acute sessions, preliminary findings show

that MICT, HIIT, and mixed (RT plus HIIT) approaches suppressed cancer cell viability and proliferation, with volume up to 2 h and an intensity ranging from moderate to high (i.e., >50% VO_{2max}, 50%–90% P_{max}, 85% HR_{max}). ^{42,43,59} In both short-term and chronic training interventions (i.e., 11 days and 9 weeks, respectively), adopting 30–60 min of MICT or HIIT, at >70% HR_{max}, at least 3 days per week suppressed cancer cell growth and proliferation, ^{22,59} while a 90-min moderate to high RT plus HIIT (i.e., 70%–90% 1RM and 80%–90% HR_{max}) for 6 months, once per week, did not inhibit cancer cell viability, although the potential limitations mentioned above should be considered. ⁴³

4. Prostate cancer

Prostate cancer is the most diagnosed cancer among men,⁶⁰ and a number of preclinical studies exploring the effects of circulating factors (e.g., myokines) on cancer cells have shown their suppressive role in various prostate cancer cell lines ^{33,34,61-64} This provides the rationale for examining such potential effects in human subjects and the effects of exercise-conditioned serum on prostate cancer cells.

4.1. Studies in subjects without prostate cancer

Similar to breast cancer, several studies have been conducted to assess whether exercise-conditioned serum from subjects without cancer suppresses prostate cancer cell lines. For the acute effects, Rundqvist et al.²⁰ determined that a single bout of MICT in 10 healthy male subjects cycling for 20 min at 50% of VO_{2max} and for another 40 min at 65% of VO_{2max} significantly decreased the proliferation and growth of prostate cancer cell line LNCaP by 31% (p < 0.05), while there was no substantial change in apoptosis (Table 1). The exercise-conditioned serum acquired 2 h post exercise revealed a decrease in epidermal growth factor (EGF) (–18%) and an increase in IGFBP-1 (+35%) compared to the baseline. Moreover, Hwang et al.⁶⁵ replicated the training protocol by Rundqvist et al.²⁰ in 12 young and 10 old male subjects, reporting that SPARC and oncostatin M (OSM) significantly increased after exercise (p < 0.05). However, when determining the effects on the cell lines LNCaP and PC3, researchers found that only LNCaP metabolic activity in the older group significantly decreased (p < 0.05), while LNCaP and PC3 cell numbers were not affected by the exercise-conditioned serum. Such results were in contrast to previous

findings, 20,42 meaning that it needs to be further investigated if a single bout of MICT at moderate intensity (i.e., 50%–65% VO_{2max}) is sufficient stimulus for cancer suppressive effects. Additionally, Baldelli et al.⁵⁹ utilized a high intensity endurance test to exhaustion in 18 healthy male subjects and reported a reduction in LNCaP proliferation by 13.8%–22.8% (p < 0.05), while exercise-conditioned serum taken after a 9-week HIIT program resulted in a reduction in LNCaP by 14.0%–27.2% (p < 0.05), which was in line with findings on breast cancer cell lines previously found by the same research group.⁵⁹

Tymchuk and colleagues²¹ investigated the effects of MICT plus a low-fat, high-fiber diet in 13 overweight subjects compared to a control group (i.e., 7 subjects) for 11 days (short-term); additionally, serum was also taken from 8 men who had been compliant with the aforementioned diet and exercise regimen for 14 years (long-term) (Table 2). The MICT consisted of 30-60 min of walking on a treadmill with intensity between 70% and 85% HR_{max} (4–5 days a week) and 40– 60 min with intensity <70% HR_{max} (1–2 days a week). When LNCaP was cultured with resting serum taken after the last training session in the short-term group, cancer cell growth was significantly reduced by 30% (p < 0.05), while no changes were observed in the control group. Additionally, a further 15% growth reduction was found when culturing LNCaP cells with serum taken from men in the long-term group (p < 0.05). Similarly, Ngo et al.⁶⁶ collected blood from 14 obese subjects before and after an 11-day MICT and low-fat diet protocol (short-term) as well as from 8 men who had followed a similar long-term intervention (i.e., 14 years). They reported a significant decrease in IGF-1 (-20%; p < 0.05) and insulin, and an increase in IGFBP-1 (+53%; p< 0.05) following the 11-day intervention. LNCaP cancer cell growth was significantly reduced by 30% in vitro, with increased apoptosis (p < 0.05) and cell necrosis with short-term training serum compared to baseline serum. Interestingly, an additional increase in apoptosis and reduction in cell growth by 14% was found when culturing LNCaP cells with serum obtained from the longterm intervention (p < 0.05), which corroborates findings from the previous investigation.²¹

Furthermore, Barnard et al. ¹⁸ compared the exercise and diet protocol adopted by Tymchuk et al. ²¹ (i.e., MICT plus low-fat, high-fiber diet for 11 days) *vs.* exercise only in 20 healthy men who had participated for at least 10 years in a University adult fitness program *vs.* 14 obese subjects (control group). The exercise only group differed in their training regimen, as subjects performed flexibility activities followed by calisthenics and swimming laps for 1 h for 5 days a week. However, it should

be noted that in the exercise only group, training volume and intensity were not reported, which impeded comparisons between groups to be drawn. Importantly, blood was collected at 1 time point only, after the training interventions. Both intervention groups had significantly lower insulin and IGF-1 levels and higher IGFBP-1 (p < 0.05), which was also significantly higher in the exercise and diet group compared to the exercise only group (p < 0.05). In addition, the exercise and diet intervention resulted in greater effects on the number of apoptotic cells compared to exercise alone (p < 0.05), while both interventions were approximately equal in their suppressive effects on LNCaP cell growth compared to the control group. Similarly, Leung et al.⁶⁷ examined 12 overweight men who participated for at least 10 years in the aforementioned flexibility, calisthenics, and swim training compared to a control group of 10 men who were obese with sedentary lifestyle and poor dietary habits with blood taken at rest. Insulin and IGF-1 were significantly lower in the intervention group, while IGFBP-1 was significantly higher in the intervention group compared to the controls (p < 0.05). LNCaP cell growth was significantly reduced by 27% and apoptosis increased by 371% in the intervention compared to the control group (p < 0.05). However, as mentioned above, training parameters were not reported. Collectively, it can be inferred that a high volume of AT or mixed training over a relatively short period was effective in altering hormones and inhibiting cancer cell growth, which is in line with previous breast cancer research,²² and, furthermore cancer suppressive effects were greater when individuals were compliant with a long-term training program (i.e., >10 years). However, it is still to be determined to what extent diet intervention impacted tumor suppressive effects. 18,21,66

4.2. Studies in patients with prostate cancer

The first acute intervention study conducted in patients with prostate cancer was done by our group. We investigated the effects of a single bout of HIIT that comprised 6 sets of 4 min at an intensity between 70% and 85% HR_{max} (rating of perceived exertion (RPE) 7–8 on the 1–10 scale) with 2 min of active recovery (50%–65% HR_{max} ; RPE 5–6) in 9 obese patients with advanced metastatic castrate resistance prostate cancer (mCRPC) who had completed at least 12 weeks of exercise in the INTERVAL-GAP4 trial (Table 3). ^{48,68} We reported a significant increase (p < 0.05) in circulatory SPARC (+19.9%), OSM (+44.8%), IL-6 (+10.2%), and IL-15 (+7.8%) as well as a trend towards an increase in decorin and irisin immediately after the single bout of exercise compared to pre-exercise and returning to baseline level after 30 min of rest. With the serum

acquired immediately after the HIIT, total cell growth of the prostate cancer cell line DU145 was reduced by 9.7%-16.9% (p<0.05), while with the serum acquired 30 min after, reduction in total cancer cell growth was between 4.9% and 8.8% compared to baseline serum (p<0.05). Interestingly, it appears that serum collected immediately after exercise had greater effects on total cell growth compared to serum collected 30 min after.

In regard to chronic training interventions, Kang et al.⁶⁹ investigated the effects of a 12-week HIIT vs. control in 52 overweight prostate cancer patients on active surveillance. The training intervention comprised 5–8 sets of 2 min at high intensity (i.e., 85%–95% VO_{2max}) and 2 min of active recovery (i.e., 40% VO_{2max}) treadmill exercise for approximately 30–40 min, 3 days per week (Table 4). Significant reductions in prostate specific antigen were found in the HIIT group compared to controls (p < 0.05), and testosterone increased only in the intervention group. Furthermore, when examining the effects on LNCaP, cancer cell growth was suppressed by –5.1% in HIIT compared to the control condition.

Subsequently, we investigated the effects of combined RT and MICT for 12 weeks in 10 obese patients on androgen deprivation therapy (ADT). 46 RT ranged from 1 to 4 sets of 6–12 RM under supervision 3 days a week, while the MICT was self-directed daily (RPE 3–8 on the 1–10 scale) to achieve 300 min per week of exercise. Resting levels of myokines SPARC and OSM increased (p < 0.05), while IGFBP-3 levels decreased (p < 0.05) in resting serum from pre- to post-intervention; no meaningful changes were observed in decorin, IGF-1, or IGFBP-3/IGF-1 ratio. When the exercise-conditioned serum after 12 weeks was applied *in vitro* to the prostate cancer cell line DU145, a significant decrease of 22.5% was observed in total cancer cell growth rate, and the mean Cell Index (i.e., the estimated record of cell morphology or cell adhesion) decreased by 21.3% (p < 0.05) after incubating the cells with collected serum for 72 h. This was the first study to demonstrate an increase in resting myokine levels with suppressive effects on cancer cells in patients with cancer after a longer-term training intervention (i.e., 12 weeks). This may result from the high training volume selected (i.e., 300 min per week) and the intensity for RT and MICT, which led to significant improvements in body weight, FM and LM percentage, and muscle strength (p < 0.05).

A similar approach was undertaken when we examined the effects of a 6-month intervention compared to controls in 25 overweight and obese patients with advanced mCRPC.⁴⁵ The

intervention group performed RT (6–12 RM, 2–5 sets) plus HIIT (6 × 60 s at RPE 8) 2 days per week, and 30–40 min of MICT (RPE 6) 1 day per week. The control group was only advised of the current American College of Sports Medicine exercise guidelines. In line with the previous study, resting levels of myokines SPARC and OSM were significantly increased (p < 0.05) compared to controls, while no substantial differences were found in decorin, IGF-1, or IGFBP-3. When comparing the effects of serum on the prostate cancer cell DU145, cell growth was significantly decreased in the training group compared to controls (by 20%; p < 0.05). This underlines how chronic mixed-methods training with appropriate volume and intensity may alter resting myokine levels and can provide systemic and persistent changes which, in turn, create a less favorable environment for tumorigenesis; however, this finding was in contrast with that of Dethlefsen et al. Additionally, and worth mentioning, despite the disease load and extensive and ongoing cancer treatments received by these patients, both acute and chronic training (as well as different training modes) showed positive effects on circulating blood factors and cancer cell suppression.

Cumulatively, the current literature on prostate cancer appears promising, with significant and clinically meaningful alterations of circulating factors and tumor suppressive effects. Regarding the acute bout of exercise, investigations examining HIIT and MICT from moderate to high intensity (i.e., 50%-65% VO_{2max}, 50%-90% P_{max}, 70%-85% HR_{max}) for 60 min resulted in suppressive effects on cancer cell growth and proliferation. ^{20,48,59,65} In short-term training (i.e., 11 days), a 30- to 60-min MICT at 70%-85% VO_{2max} at least 4 days a week reduced cancer cell growth and increased apoptosis, with systemic changes induced by long-term training (i.e., 10 years) having cancer suppressive effects. ^{18,21,66,67} Lastly, HIIT alone and RT coupled with HIIT or MICT showed inhibitory effects on cancer cell growth and proliferation in chronic interventions when performed at a moderate to high intensity (i.e., 6–12 RM, RPE > 3, 85%–95% VO_{2max}) for at least 9 weeks, 3 days a week. ^{45,46,59,69}

5. Colorectal cancer

Colorectal cancer is the third most diagnosed cancer and the third leading cause of cancer death.^{70,71} To the best of our knowledge, very few preclinical studies have been conducted to date and even fewer involving human subjects.⁷² Although the evidence is sparse, we have summarized

the current literature pertaining to the effects of exercise-conditioned serum on colorectal cancer cells, which is crucial given the incidence and detrimental effects of this cancer.

5.1. Studies in subjects without colorectal cancer

Orange and colleagues⁷³ recruited 16 obese, sedentary males to undertake 60 min of moderate intensity interval training (MIIT) comprising 6×5 -min bouts of cycling at 60% HR_{reserve} (Table 1). Serum measured immediately after the single bout of exercise revealed a significant increase in IL-6 (+24.6%; p < 0.05) from pre to post, while no changes were found in IL-8, TNF- α , OSM, and SPARC. When exercise-conditioned serum was applied to LoVo cells (i.e., colorectal cancer cell lines) *in vitro*, proliferation was significantly reduced by 4.2% when compared to serum in the resting state (p < 0.05). Additionally, a significant decrease by 5.7% in cell proliferation was observed when examining the exercise-conditioned serum as opposed to serum acquired from the participants following a non-exercise control condition (p < 0.05). Thus, although only IL-6 was significantly altered, serum taken immediately after exercise reduced colon cancer cell proliferation. These findings are similar to those observed in other studies following acute exercise in people with and without cancer. 20,44,48,59 It should be noted that this is the first study using MIIT in which it appears to be a sufficient stimulus, possibly because the population was obese with a mean age of 60 years.

5.2. Studies in patients with colorectal cancer

Similarly, Devin et al.⁴⁴ examined both the effects of a single bout of exercise and short-term training in 10 overweight and obese male colorectal cancer survivors. The single bout of exercise (i.e., HIIT) was based on 4×4 -min bouts of cycling at 85%–95% HR_{max}, with 3 min of active recovery (total work 38 min) (Table 3). Significant increases were observed in IL-6, IL-8, TNF- α , and insulin levels (by 44.8%, 24.7%, 15.2%, and 38.8%, respectively) in serum acquired immediately after the acute exercise session (p < 0.05), with circulating factors returning to baseline levels 2 h post exercise. When applying the exercise-conditioned serum *in vitro*, a significant reduction was found in cell number for the colon cancer cell lines CaCo-2 (effect size (ES) = -1.7 to -1.1; p < 0.05) and LoVo (ES = -1.2 to -0.8; p < 0.05), which is similar to the results of investigations in other cancer populations. ^{43,48,74} Thus, HIIT performed at high intensity (i.e., 85%–95% HR_{max}) altered circulating factors and cancer cells, emphasizing that intensity is likely to be a potential moderator in promoting those alterations.

The short-term training program repeated the same HIIT undertaken 3 times weekly over a 4-week period. When examining the serum obtained at rest, there was no significant change in levels of IL-6, IL-8, TNF-α, insulin, glucose, or IGF-1, or in the viability of the cell lines CaCo-2 and LoVo compared to baseline⁴⁴ (Table 4). These results are in line with those of previous findings, where lack of change in blood contents translates to no apparent suppressive effects on cancer cells.⁴³ Our assumption is that the exercise volume selected (i.e., 38 min, 3 sessions per week, for 4 weeks) was insufficient to drive substantial alterations in blood collected at rest, as previously mentioned.⁴³

Cumulatively, there is insufficient evidence to draw meaningful conclusions about exercise mode, volume, and intensity for this type of cancer population. Preliminary evidence shows that acute MIIT and HIIT performed at approximately moderate to high intensity (i.e., 60% HR_{reserve}, 85%–95% HR_{max}) for at least 40 min resulted in a reduction in cell number and proliferation and alterations in circulating factors to a modest extent only ^{44,73} while a 4-week HIIT program performed 3 days per week (i.e., short-term) at 85%–95% HR_{max} did not elicit substantial modifications on serum content as well as cancer cell number.⁴⁴

6. Lung cancer

Lung cancer is the leading cause of cancer death worldwide.⁷⁵ As a result, compelling actions are required to mitigate the burden. However, the cancer load and co-morbidities that accompany a diagnosis of lung cancer make it difficult for experimental studies to be undertaken in this patient group.⁷⁶ Although the number of preclinical studies in this cancer type is increasing ^{32,35,37,77,78}, to date only 1 study has been conducted in humans involving lung cancer cells.

6.1. Studies in subjects without lung cancer

Kurgan et al.⁴⁷ recruited 23 male university students to perform a single 1-min high-intensity (i.e., 90% of maximum workload) bout on a stationary cycle ergometer followed by 1 min of active recovery, which was repeated 6 times (Table 1). As previously reported, in this study IL-6, IL-1 β , IL-1 α , TNF- α significantly increased 5 min after the acute exercise session (p < 0.05) and returned to baseline after 1 h.⁷⁹ The exercise-conditioned serum collected after exercise at 5 min, 1 h, and 24 h resulted in a reduction of lung cancer A549 cell proliferation by 84% to 91% and in survival rates by 21.5% to 35.8% (p < 0.05). Similarly, lung cancer cell lines H460 and H1299 showed a

decrease in survival rates ranging from 33.5% to 41.9% when cells were cultured with post-exercise serum (p < 0.05).

Thus, a brief HIIT protocol at high intensity (i.e., 90%_{max} workload) resulted in suppressive effects on lung cancer cell proliferation and survival rates, similar to those with and without cancer. ^{20,44,48,59,73} However, additional studies, particularly in lung cancer patients, are required for definitive conclusions to be drawn.

7. Directions for future research

Although preliminary, current studies in healthy, obese, and cancer patients and survivors indicate that acute training sessions adopting HIIT, MIIT, MICT, and mixed training (i.e., RT plus HIIT or MICT) at moderate to high intensity for at least 40 min confer an inhibitory effect on cancer cells and alter circulating factors. ^{20,42–44,47,48,59,73} In regard to short-term and chronic training interventions, HIIT and MICT (alone or coupled with RT performed for at least 30 min, 3 days a week) result in an alteration in resting circulating factors and provide a tumor suppressive effect. ^{18,21,22,43–46,59,66,67,69} However, future research is necessary to expand our understanding of the anti-cancer role of exercise (Fig. 1). The following are potential avenues of investigation:

(a) Further research should be focused on patients with cancer or a history of cancer (i.e., survivors) and, subsequently, stratifying by treatments (e.g., chemotherapy) or by disease stage. Furthermore, it should be noted that only studies in breast, prostate, colorectal, and lung cancer have been conducted to date, which represents a limitation of the current review and the state of knowledge more generally. The underlying reason is that cancer alters several physical and physiologic components. Thus, studies conducted on healthy subjects showing modifications in circulating factors and cancer cell behaviors should be interpreted with caution, even though this was the logical first line of investigation. In support of this, it should be acknowledged that cancer treatments (e.g., ADT for patients with prostate cancer) or stages (e.g., localized prostate cancer vs. mCRPC) may affect or impair a patient's ability to adapt with the morphological (e.g., muscle hypertrophy) and metabolic (e.g., hormonal adaptation) changes induced by exercise. Thus, their responses may differ from those without cancer and, from a practical point of view, this may lead to more targeted and tailored training programs. In line with this, we may expand our knowledge

about the timing of exercise in patients with cancer by exposing cancer cells to serum collected before, during, and after various cancer treatments *in vitro*. Furthermore, although it has not been discussed in detail, it was shown that elevated shear pressure caused by exercise in the circulatory system could induce apoptosis and necrosis of cancer cells in a dynamic-experimental setting using a microfluid system. ⁸² As such, evaluating the effects of shear stress induced by different exercise modes, volume and intensity on cancer cells in larger clinical trials would provide novel insight regarding targeted exercise prescriptions to reduce recurrence and metastasis of cancer. ⁸²

- (b) To deepen our knowledge of cancer suppressive mechanisms induced by exercise, it is necessary to design exercise trials with experimental models that include appropriate properties of cancer cell lines, co-culture systems, and parallel studies involving animal models and humans.⁸³ For instance, cancer cells in a co-culture system with cells known to promote cancer growth (e.g., adipocytes and immune cells) exposed to exercise-conditioned serum would provide biological insight about the impact of exercise on the interaction between cancer cells and cells around the tumor sites. Additionally, animal models could be employed to establish the *in vivo* causation of exercise-induced candidate factors, followed by validation in humans. Alternatively, the most optimal scenario for translating these findings to humans would involve window of opportunity trials.⁸⁴ Further, although it may be difficult to connect exercise mode, volume, and intensity in animal models and clinical trials, well-designed parallel animal and human studies will provide greater understanding of physical, physiological, and biological effects of exercise on specific cancer groups by validating preclinical outcomes in humans and *vice versa*.⁸³
- (c) In regard to training modes and chronic adaptations, most studies adopted a mixed mode approach incorporating both RT and AT.^{43,45,46} This is likely due to the combination of training modes resulting in greater benefits over a wider range of outcomes for those with and without cancer (e.g., FM, LM, muscle strength, cardiorespiratory capacity, *etc.*). On the other hand, studies examining an acute exercise bout mainly used AT in the form of HIIT, MIIT, or MICT.^{42,44,48,73} Although results are promising, investigations comparing different training modes (i.e., RT *vs.* AT), including also alternative RT training methods (e.g., cluster set training),⁸⁵ may elucidate whether different training stimuli result in different adaptations in both the acute and chronic training settings.

- (d) The effects of acute exercise sessions appear to be primarily dictated by intensity. ^{20,42–44,48,59,65,73} In this case, blood taken immediately after a single bout of exercise has been shown to have a higher content of circulating factors. However, it is still unknown whether there is a threshold that drives modifications to occur. Furthermore, it remains unclear whether a person's training status (i.e., trained *vs.* untrained) may play a role in eliciting exercise-induced factors after a single exercise session. Future investigations should explore single bouts of exercise conducted at different training intensities and with training statuses to determine whether a minimal threshold exists for impacting cancer cells.
- (e) For short-term and chronic interventions, current findings show that a high volume of combined resistance and aerobic exercise (i.e., >3 days a week) is necessary to alter circulating factors. ^{21,22,45–47,66,67,69,86} Future studies should take into account that 8–12 weeks of RT (2 sets of 8–15 repetitions performed at a moderate intensity at least 2 times per week) and AT (30–60 min performed at a moderate to vigorous intensity at least 3 times per week) are necessary to drive physical adaptations and, subsequently, changes in resting circulating factors (e.g., myokines). ⁸⁷ However, future investigations should explore what amount of training volume, frequency, and duration is required to induce such alterations. Additionally, whether more volume results in greater cancer suppressive effects is yet to be determined.
- (f) In regard to circulating factors, future investigations should examine myokines in depth, owing to their potential direct tumor suppressive effects observed to date (Fig. 2).⁸⁸ In particular, determining whether or not different training modes (i.e., RT *vs.* AT), volume and intensity elicit different responses in myokine expression in patients with cancer or survivors is of utmost importance to clearly elucidate potential underlying mechanisms and, therefore, implement practical applications in exercise oncology.⁸⁹
- (g) Lastly, it is unclear whether changes in body weight and body composition as a result of exercise training, disease, and treatments are related to different expressions of resting blood content. Our assumption is that an association may exist between body weight, body composition, and circulating factors (e.g., myokines). This comes from the fact that myokines are produced by skeletal muscles and a "higher" volume of this endocrine organ may translate to higher production;⁹⁰ however, this is only speculative. Thus, further research may explore to what extent alterations in body weight and body composition (in chronic interventions) relate to changes in

circulating factors, and whether patients with higher skeletal muscle mass would have a higher myokine response after an acute bout of exercise.

8. Conclusion

The potential anti-cancer and tumor suppressive effects of exercise have garnered increased attention in the last two decades. Several avenues of research demonstrate that exerciseconditioned serum has inhibitory effects on cancer cell growth and promotes apoptosis in different cancer cell lines. However, the precise role played by different training modes, volume, and intensity on exercise-conditioned serum and its effects on cancer cells remains unclear. The limited number of studies available in a relatively new area of exercise oncology precludes definitive conclusions to be drawn. However, the findings to date show that a single moderate to vigorous intensity bout of AT may have suppressive effects on cancer cells and elicit substantial increases in circulating factors. When examining short-term and chronic training interventions, it appears that training volume may play a role in inhibiting cancer cells and altering circulating factors, regardless of the training mode used. Future research should be directed to targeted investigations into the effects of exercise-conditioned serum on cancer cells based on specific training modes (i.e., RT vs. AT), volume, and intensity, to better understand the underlying mechanisms to induce suppressive cancer effects and alterations in circulating factors while considering muscle and fat mass and changes in body composition due to disease and treatments as well as exercise medicine and diet therapy intervention.

Data availability statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Authors' contributions

FB conceived the study design, searched studies in databases, elaborated the results and drafted the manuscript; DT, DG, CB, JSK, RN edited and revised the manuscript. All authors have read and approved the final version of the manuscript, and agree with the order of presentation of the authors.

Competing interests

The authors declare that they have no competing interests.

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Table 1. Summary of acute exercise studies in subjects without cancer.

Cance r	Subj ects	Trainin g interven tion	Blood samplin g	Circulatin g factors	Exercise effects (pre vs. post)	Setti ng and canc er cell line	Effects on cancer cells
Breast	Healt hy fema le (age = 24, BMI = 23) ⁴²	INT (n = 7): single bout of MICT (at 55% VO _{2max} on stationa ry cycle ergomet er) for 2	Pre: resting; post: 1 and 2 h after exercis e	IL-6 Lactate Epinephri ne Norepine phrine	↑* (2.4-fold) ↑* (1.5-fold) ↑*	Seru m exer cise in vitro on breas t canc er cell	Pre vs. post serum 1 h after: MCF-7 cell viabilit y ↓* by -10%; MDA- MB- 231 cell

	h under supervi sion		MCF -7 and MD A- MB- 231	viabilit y ↓* by -14%; Pre vs. post serum 2 h after: MCF-7 cell viabilit y ↓* by -19%; MDA- MB- 231 cell viabilit y ↓* by -13% at 48 h Pre vs. post serum immedi ately
Healt hy fema le (age = 21, BMI = 21) ⁵⁹	INT (n = 12): HIEC (increm Pre: 2 h ental fasting; test post: from immedi 50% to ately 90% after, 4 Pmax up h and to 24 h exhaust after ion on exercis stationa e ry cycle ergomet er)	Immediately after: ↑; 4 and 24 h: N/C	Seru m exer cise in vitro on breas t canc er cell MD A-MB-231	after: MDA- MB- 231 cell prolifer ation ↓* by − 12.1%; Pre vs. post serum 4 h after: MDA- MB- 231 cell prolifer ation ↓* by − 24.0%; Pre vs. post serum

	Healt hy male (age = 25, BW = 58– 82 kg) ²⁰	INT (n = 10): single bout of MICT (20 min at 50% VO _{2max} , 40 min at 65% VO _{2max} on stationa ry cycle ergomet er)	Pre: resting; post: 2 h after exercis e	IGFPB-1 EGF Cortisol	↑ (+35%) ↓ (-18%) N/C	Seru m exer cise in vitro on prost ate canc er cell LNC aP	24 h after: MDA- MB- 231 cell prolifer ation ↓* by - 24.9% Pre vs. post: LNCaP cell growth ↓* by 31% at 96 h; LNCaP prolifer ation ↓* at 48 h; LNCaP apoptos is N/C at 24 and 48
Prostate	Healt hy male (you ng grou p: age = 28, BMI = 23; old grou p: age = 63, BMI = 25) ⁶⁵	INT (n = 20): single bout of MICT (20 min at 50% VO _{2max} , 40 min at 65% VO _{2max} on stationa ry cycle ergomet er)	Pre: fasting; post: immedi ately after exercis e	IL-6 IL-15 Irisin OSM SPARC	NA NA NA ↑* ↑* (young); ↑ (old)	Seru m exer cise in vitro on prost ate canc er cell LNC aP and PC3	h Pre vs. post young: LNCaP metabol ic activity N/C; PC3 metabol ic activity N/C; LNCaP cell number N/C; PC3 cell

number N/C; *Pre vs.*

```
post
                                                                           old:
                                                                           LNCaP
                                                                           metabol
                                                                           ic
                                                                           activity
                                                                           ↓*; PC3
                                                                           metabol
                                                                           ic
                                                                           activity
                                                                           N/C;
                                                                           LNCaP
                                                                           cell
                                                                           number
                                                                           N/C;
                                                                           PC3
                                                                           cell
                                                                           number
                                                                           N/C at
                                                                           24 and
                                                                           48 h
                                                                           Pre vs.
                                                                           post
                                                                           serum
       INT (n
                                                                           immedi
       = 18):
                                                                   Seru
                                                                           ately
       HIEC
                                                                   m
                                                                           after:
                  Pre: 2 h
       (increm
                                                                           LNCaP
                                                                   exer
       ental
                  fasting;
Healt
                                                                   cise
                                                                           cell
       test
                  post:
                                                                           prolifer
hy
                                                                   in
                  immedi
        from
male
                                                                   vitro
                                                                           ation ↓*
       50% to
                  ately
(age
                                                                           by -
                                                                   on
       90%
                                         Immediately after: ↑; 4
                  after, 4
= 21,
                                                                           13.8%;
                            CK
                                                                   prost
       P<sub>max</sub> up
                  h and
                                         and 24 h: N/C
BMI
                                                                   ate
                                                                           Pre vs.
                  24 h
       to
                                                                           post
                                                                   canc
                  after
       exhaust
(22)^{59}
                                                                   er
                                                                           serum 4
                  exercis
       ion on
                                                                   cell
                                                                           h after:
       stationa
                                                                   LNC
                                                                           LNCaP
       ry cycle
                                                                   aP
                                                                           cell
       ergomet
                                                                           prolifer
       er)
                                                                           ation ↓*
                                                                           by-
                                                                           21.8%;
```

Pre vs.

Healt hy male Lung $= 23$): before single exercis $= 23$): before single exercis $= 24$ h: N/C $= 2$	Colore ctal	Healt hy male (age = 60, BMI = 30) ⁷³	INT (n = 8): single bout of MIIT (6 sets × 5 min at 60% HR _{reserv} e, 2.5 min active recover y, on stationa ry cycle ergomet er) for 60 min; CON (n = 8): rest	Pre: before exercis e (NA); post: immediately after	IL-6 IL-8 IL-10 TNFα Irisin OSM SPARC	INT vs. CON and Pre vs. post †* (+24.6%) N/C NA N/C NA N/C N/C N/C	Seru m exer cise in vitro on colo n canc er cell LoV o	post serum 24 h after: LNCaP cell prolifer ation ↓* by - 22.8% Pre vs. post INT: LoVo cell prolifer ation ↓* by - 4.2%; Pre vs. post CON: LoVo cell prolifer ation ↑* by 5.4%; INT vs. CON: LoVo cell prolifer ation ↑* by 5.4%; INT vs. CON: LoVo cell prolifer ation ↑* by 5.4%; INT vs. CON: LoVo cell prolifer ation ↓* by 5.4%; INT vs. CON: LoVo cell prolifer ation ↓* by - 5.7% at 48 h
	Lung	hy male (age = 22,	= 23): single bout of HIIT (6	before exercis e (NA); post: 5	IL-1β	24 h: N/C 5 min after: ↑*; 1 and 24 h: N/C	m exer cise in	post serum 5 min after:

= 24) ⁴⁷	90% max workloa d, and 1 min active recover y on stationa ry cycle ergomet er)	h after exercis e	IL-10	5 min after: ↑; 1 and 24 h: N/C 5 min after: ↑*; 1 and 24 h: N/C	lung canc er cell A54 9, H46 0, and H12 99	prolifer ation ↓* by 90%; Pre vs. post serum 1 h after: A549 cell prolifer ation ↓* by 91%; Pre vs. post serum 24 h after: A549 cell prolifer ation ↓* by 84%; Pre vs. Post serum 5 min after: A549 survival rates ↓* by 21.5%;
3						rates ↓* by

post

```
serum 1
h after:
A549
survival
rates ↓*
by
33.9%;
H460
survival
rates ↓*
by
38.4%;
H1299
survival
rates ↓*
by
41.9%;
Pre vs.
post
serum
24 h
after:
A549
survival
rate
survival
rates ↓*
by
35.8%;
H460
survival
rates ↓*
by
39.3%;
H1299
survival
rates ↓*
by
37.7%
at 7
days
```

Note: \uparrow = increase; \downarrow = decrease.

```
* p < 0.05.
```

Abbreviations: BMI = body mass index; BW = bodyweight; CK = creatine kinase; CON = control; HIEC = high intensity endurance cycling; HIIT = high intensity interval training; HR = heart rate; INT = intervention; MICT = moderate intensity continuous training; MIIT = moderate intensity interval training; NA = not available; N/C = no changes; P_{max} = maximal power; VO_{2max} = maximal oxygen uptake.

Table 2. Summary of chronic exercise studies in subjects without cancer.

Cance Subjects	Training interventi on Blood sampl	Circulatin g factors	Exercise effects	Settin g and Effects cancer on cancer cell cells line
Overwei ght and obese post-menopau sal Breast female (age = 51–79, BMI = 30 to 32, $n = 28$ or HT) ²²	HR _{max} on treadmill 14–5 overning fasting post: 0 overning fasting post: 0 overning fasting post: 0 overning fasting	ight	Pre vs. post ↑ * (+32%) ↓ * (-19%) ↓ * (-29%) ↓ * (-37 to -34%)	Pre vs. post: MCF-7 cell growth Serum ↓* by exerci 6.6%; se in ZR-75-1 vitro cell on growth breast ↓* by cancer 9.9%; T- cell 47D cell MCF- growth 7, T- ↓* by 47D 18.5%; and MCF-7 ZR- apoptosis 75-1 ↑* by 23%; ZR- 75-1 apoptosis ↑* by 20%; T-

				47D apoptosis †* by 30% at 48 h
to 90% P _{max} up to exhaustion on stationary cycle ergometer) after the last session	C Pre: 2 h n fasting; post: n immediat ely after, 24 h and 24 h after exercise	Immediately after: \(\tau\); 4 and 24 h: N/C	exerci se in vitro on breast cancer cell MDA -MB- 231	serum 4 h after: MDA- MB-231 cell proliferati on ↓* by -30.6%; Pre vs. post serum 24 h after: MDA- MB-231 cell proliferati on ↓* by -35.3%
Prosta Overwei = 13): te ght male MICT (age = (30–60) min at	Pre: 12 h fasting; post: 12 h Testostero fasting ne	Pre vs. post in INT and INT vs. CON ↓*	exerci se <i>in</i>	-

59, BMI = 29) ²¹	70% to 85% HR _{max} on treadmill 4–5 days/wee k; 40–60 min < 70% HR _{max} 1– 2 days/wee k) for 11 days; INT2 (n = 8): MICT (60 min, daily) + low fat high fiber diet for 14 years;	Cholester ol LDL HDL Triglyceri des	↓* N/C ↓*	te cance r cell LNCa P	growth ↓* by 30%; Pre vs. post INT2: LNCaP cell growth ↓ * by additional 15%; Pre vs. post CON: LNCaP cell growth N/C at 48 h
	CON (<i>n</i> = 7)				
			Pre vs. post INT1		Pre vs.
	INT1 (<i>n</i> = 14):	IGF-1	↓* (<i>-</i> 20%)		post INT1:
	MICT (60 min,	IGFPB-1	†* (+53%)	Serum	LNCaP
	daily) +	IGFPB-3	N/C	exerci se <i>in</i>	growth
Obese male (age = 60, BMI = 38) ⁶⁶	low fat high fiber diet for diet for fasting; 11 days; post: 12 h INT2 (n fasting) = 8): MICT (60 min, daily) + low fat high fiber	Insulin	↓	te cancer cell LNCa P	↓* by 30%; LNCaP apoptosis ↑*; LNCaP necrosis ↑; Pre vs. post INT2: LNCaP

```
diet for
                                                                             cell
         14 years
                                                                             growth
                                                                             ↓* by
                                                                             additional
                                                                             14%;
                                                                             LNCaP
                                                                             apoptosis
                                                                             ↑* at 48 h
         INT1 (n
                                        INT1 vs. INT2 vs. CON
         = 8):
                              IGFPB-1
                                        ↑ *
         MICT
         (30-60)
                              IGF-1
         min at
         70%-
                              Insulin
         85%
                                        INT1 vs. INT2
         HR<sub>max</sub> on
         treadmill
         4–5
Healthy
                                                                             INT1 and
         days/wee
(INT1:
age = 55, k; 40–60
                                                                             INT2 vs.
                                                                      Serum
                                                                             CON:
         min <
BMI =
                                                                      exerci
                                                                             LNCaP
         70%
21.5;
                                                                      se in
                                                                             cell
         HR_{max}\ 1-
INT 2:
                                                                      vitro
                                                                             growth
age = 62, 2
                   Post:
                                                                      on
         days/wee overnight
BMI =
                                                                      prosta
                                                                             LNCaP
26.5) and k) + low
                   fasting
                                                                      te
                                                                      cancer †*; INT1
         fat high
obese
         fiber diet
male
                                                                      cell
                                                                             vs. INT2:
         for 11
                                                                      LNCa
(CON:
                              IGFPB-1 ↑*
                                                                             LNCaP
age = 60, days
                                                                      P
                                                                             apoptosis
         following
BMI =
                                                                             ↑* at 48 h
         a regimen
38)^{18}
         for 14
         years;
         INT2 (n
         = 12):
         continuou
         s and
         strenuous
         calistheni
         c and
         swimmin
```

```
g laps for
          50 min 5
          days/wee
          k
          following
          a regimen
          for 10
          years;
          CON (n
          = 14)
          INT (n =
                                         INT vs. CON
          12):
                              IGFPB-1
          continuou
          s and
                                                                              INT vs.
                              IGF-1
Healthy
                                                                       Serum
          strenuous
                                                                              CON:
(INT:
                                                                       exerci
                                                                              LNCaP
          calistheni
age = 60,
                                                                       se in
          c and
                                                                              cell
BMI =
                                                                       vitro
                                                                              growth
          swimmin
26) and
overweig g laps for
                                                                       on
                                                                              ↓* by
                                                                       prosta
                    overnight
          50 min 5
                                                                              27%;
ht male
                    fasting
                                                                       te
          days/wee
                                                                              LNCaP
(CON:
                                                                       cancer
                              Insulin
                                                                              apoptosis
         k
age = 62,
                                                                       cell
          following
                                                                              ↑* by
BMI =
                                                                       LNCa
                                                                              371% at
          a regimen
31)^{67}
                                                                       P
          for 10
                                                                              48 h
          years;
          CON (n)
          = 10)
          INT(n =
                                                                              Pre vs.
                                         Pre vs. post
          18): HIIT
                                                                       Serum post
                                                                       exerci serum
          (3-4)
         days/wee Pre: 2 h
                                                                              immediat
                                                                       se in
                    fasting;
         k) for 9
                                                                              ely after:
Healthy
                                                                       vitro
                    post:
          weeks
                                                                              LNCaP
male
                                                                       on
         and HIEC immediat
                                         Immediately after: ↑; 4 and 24 prosta cell
(age =
21, BMI (incremen ely after, CK
                                                                              proliferati
                                                                       te
                                         h: N/C
                    4 h and
                                                                       cancer on ↓* by
          tal test
=22)^{59}
         from 50% 24 h after
                                                                              -14.0\%;
                                                                       cell
                    exercise
          to 90%
                                                                       LNCa Pre vs.
          Pmax up
                                                                              post
                                                                       P
                                                                              serum 4 h
          to
          exhaustio
                                                                              after:
```

n on		LNCaP
stationary		cell
cycle		proliferati
ergometer		on ↓* by
) after the		-22.9%;
last		Pre vs.
session		post
		serum 24
		h after:
		LNCaP
		cell
		proliferati
	C	on ↓* by
		-27.2%

Note: \uparrow = increase; \downarrow = decrease.

Abbreviations: BMI = body mass index; CON = control; HDL = high-density lipoprotein; HIIT = high intensity interval training; <math>HR = heart rate; HT = hormone therapy; INT = intervention; LDL = low-density lipoprotein; MICT = moderate intensity continuous training; N/C = no changes; $P_{max} = maximal$ power.

Table 3. Summary of acute exercise studies in patients with cancer.

Cancer	Subjects	Training intervent ion	Blood sampling	•	Exercise effects (pre vs. post)	Setting and cancer cell line	Effects on cancer cells
	Breast	INT (<i>n</i> =				Serum	Pre vs.
	cancer	20):	Pre: 2 h	IL-6	^* (+110%)	exercis	post:
Breast	patients	single	fasting;	IL 0	(111070)	e in	MCF-7
	undergoin	bout of	post:	IL-8	↑* (+20%)	vitro	cell
	g	warmup	immediat	II 10	N/C	on	viability
	chemother	for 30		IL-10	N/C	breast	↓* by –

^{*} *p* < 0.05.

apy (age = 49, BMI =	min + RT for	ely after exercise	TNFα Insulin	↑* (+13%) N/C	cancer	9.2%; MDA-
$(26)^{43}$	60 min +					MB-231 cell
	HIIT (at 80% to		Lactate	^* (+500%)	and MDA-	viability
	85% HR _{max}		Epinephrine	^* (+190%)	MB- 231	↓* by – 9.4% at 48 h
	stationar y cycle ergomete					
	r) for 30 min		X	c .		
	under		Norepineph	^* (+120%)		
	supervisi		rine			
	on					
	(whole			40		
	session					
	>70%					
	HR_{max})					
	INT (<i>n</i> =					
	20):					
	single		Epinephrine	↑*		
	bout of					
	warmup				Serum	Pre vs.
	for 30	10			exercis	Post:
Breast	min +				e in	MCF-7
cancer	RT for	Dua			vitro	cell
patients	60 min +	Pre:			on	viability
undergoin	HIIT (at	resting;			breast	↓* by –
g	>80%	post: immediat			cancer	11%; MDA-
chemother	HR_{max}	ely after	Norepineph	^ *	cell	MB-231
apy (age =	on	exercise	rine	I	MCF-7	cell
49, BMI =		0.1010150			and	viability
$(26)^{42}$	y cycle				MDA-	↓* by –
	ergomete				MB-	9% at 48
	r) for 30				231	h
	min					
	under supervisi					
	on					
	OII					

				Pre vs.
	INT (n = 9): single bout of HIIT (6	SPARC	Immediately after: ↑* (+19.9%);	post serum immediat ely after:
	sets × 4 min at 70%– 85%	OSM	30 min: N/C Immediately after: ↑* (+11.5%); 30 min: N/C	DU145 total cell growth \$\d\ \psi\$ by -
Metastatic castrate- resistant prostate	HR _{max} or RPE 7— Pre: 2 h 8, 2 min fasting; active post:	IL-6	Immediately after: ↑* (+10.2%); 30 min: N/C	Serum 9.67% exercis and – e <i>in</i> 16.93% <i>vitro</i> at 48 and
Prostate cancer patients (age = 67, BMI =	recovery immediat at 50%— ely and 65%— 30 min HR _{max} or after RPE 5–6 exercise	IL-15	Immediately after: ↑* (+7.8%); 30 min: N/C	on 72 h; Pre prostat vs. post e serum 30 cancer min cell after:
30) ⁴⁸	on stationar y cycle	Decorin	Immediately after: \(\pm\); 30 min: N/C Immediately after: \(\pm\); 30	DU145 DU145 total cell growth
	ergomete r) for 40 min under supervisi on	Irisin IGF-1	min: N/C Immediately after: ↑; 30 min: N/C	↓* by – 4.91%, – 6.91% and – 8.82% at 24, 48, and 72 h
Colorectal cancer	INT (n = 10): single Pre: bout of fasting HIIT (4 state;	IL-6	Immediately after: ↑* (+44.8%); 2 h: N/C	Serum post exercis serum e in immediat vitro ely after:
Colorec survivors tal (age = 67, BMI = 28) ⁴⁴	sets × 4 post: min at immediat 85%- ely and 2 95% h after	IL-8	Immediately after: ↑* (+24.7%); 2 h: N/C	colorec CaCo-2 tal cell cancer number cell \pmoderacter (ES
	HR _{max} , 3 exercise min active recovery	TNFα	Immediately after: \uparrow * (+15.2%); 2 h: N/C	CaCo- range = - 2 and 1.7 to - LoVo 1.1); LoVo

```
cell
   on
stationar
                                                                 number
y cycle
                                                                  ↓* (ES
ergomete
                                                                range = -
r) for 40
                                                                 1.2 to -
  min
                                                                   0.8);
                                                                 CaCo-2
                                                                   and
                                                                  LoVo
                                                                apoptosis
                                 Immediately after: ↑*
                                                                N/C; Pre
                                       (+38.8%);
                     Insulin
                                                                 vs. post
                                        2 h:↓*
                                                                 serum 2
                                                                 h after:
                                                                 CaCo-2
                                                                   and
                                                                  LoVo
                                                                   cell
                                                                 number
                                                                  N/C at
                                                                 24, 48,
                                                                 and 72 h
```

Note: \uparrow = increase; \downarrow = decrease.

Abbreviations: BMI = body mass index; HIIT = high intensity interval training; HR = heart rate; INT = intervention; N/C = no changes; RPE = rating of perceived exertion; RT = resistance training.

^{*} p < 0.05.

 Table 4. Summary of chronic exercise studies in patients with cancer.

Cancer	Subjects	Training interventio	Blood samplin g	Circulatin g factors	Exercise effects	Setting and cancer cell line	s on cancer
Breast	Breast cancer survivors (age = 47, BMI = 24) ⁴³	and lower back extension) + HIIT (30	Pre: 8 h overnig ht fasting; post: 8 h overnig ht fasting	IL-6 IL-8 IL-10 TNFa Insulin Leptin Glucose	INT vs. CON and Pre vs. post ↓* (INT = -37%; CON = - 20.7%) N/C N/C ↓* (INT = -21.1%) N/C N/C N/C 11.4%)	Serum exercise in vitro on breast cancer cell MCF-7 and MDA- MB- 231	INT vs. CON and Pre vs. post: MCF- 7 and MDA- MB- 231 cell viabili ty N/C at 48 h

		every 2–3 months; CON (<i>n</i> = 37): 3 health evaluation consultatio				
		INT (<i>n</i> =			Pre vs. post INT	
		26): HIIT (5–8 sets ×		PSA	↓ *	
	surveillan	2 min at 85%–95% VO _{2max} , 2 min active recovery Pre: 12		Testostero ne	1	
				PSA	Pre vs. post CON	INT Plasma vs.
Prostate					N/C	exercise CON: in vitro LNCa on P cell prostate growt cancer h \pm* cell by - LNCaP 5.1% at 48 h
		at 40% VO _{2max} on fast treadmill) por for 40 min 3 fadays/week for 12 weeks under supervisio n; CON (n = 26)	st: 12 h sting	Testostero ne	N/C	
		INT (<i>n</i> =			Pre vs. post	Pre vs.
	Prostate cancer patients on ADT (age = 73, BMI = 33) ⁴⁶	groups of h after the upper exercise and lower e body)	h	Irisin	NA	<i>post</i> : DU14
				Decorin	N/C	Serum 5 total
				IL-6	NA	<i>in vitro</i> growt
			sting;	IL-15	NA	on h \psi*; prostate DU14 cancer 5
				SPARC	↑	
				OSM	↑ *	cell mean DU145 cell
				Myostatin	NA	index
		under supervisio		IGF-1	N/C	↓* by _

	n, 3		IGFBP-3	*	21.3%
	days/week				;
	+ MICT				DU14
	(at RPE 3-				5
	8) self-				averag
	directed				e
	daily for				growt
	12 weeks,				h rate
	2				↓* by
	consultatio				_
	ns with		IGF-		22.5%
	dietician		1/IGFBP-	N/C	at 72 h
	to induce			IV/C	
	caloric		3		
	deficit of				
	500-1000				
	kcal per		4	10	
	week + 40				
	gr whey				
	protein				
	supplemen				
	t		2/		
	INT (<i>n</i> =			INT vs. CON	INT
	13): RT			INI VS. CON	VS.
	(6–12 RM		Irisin	NA	CON:
	and 1–5	Pre: 10 h	Danamin	N/C	DU14
N			Decorin	N/C	5 total
Metastati			IL-6	NA	cell
C				374	Serum growt
castrate- resistant	muscle		IL-15	NA	exercise h \psi*
prostate		overnig ht	SPARC	^ *	<i>in vitro</i> by –
cancer	the upper	fasting;			on 20% at
patients	and lower	nost 18	OSM	*	prostate 0–72
patients $(age = 72)$	body) +		Myostatin	NA	cancer h;
to 76,	HIIT (6	exercis			cell DU14
BMI =	$sets \times 60 s$	e	IGF-1	N/C	DU145 5
$28-31)^{45}$	at RPE 8)	·	IGFBP-3	N/C	adjust
26–31)	under		נ-זמיוטו	11/0	ed
	supervisio		IGF- 1/IGFBP-		mean
	n 2			N/C	cell
	days/week		3	-	index
	; MICT				↓* at

min at RPE 6, cycling or walking) for 40 min under supervisio n 1 day/week for 6 months; CON (n = 12): self- directed exercise ACSM			0-24, 24-48, 48-72 h
guidelines INT ($n = 10$): HIIT ($4 \text{ sets} \times 4$ min at $85\%-95\%$ Pre: HR _{max} , 3 resting min active and recovery fasting; on post: $8MI = 30\%$	IL-6 IL-8 TNFα Insulin Glucose	Pre vs. post N/C N/C N/C N/C N/C N/C	Pre vs. Serum post: exercise CaCo- in vitro 2 and colorect LoVo al cell cancer numbe cell r N/C
ergometer) fasting for 40 min state 3 days/week for 4 weeks	IGF-1	N/C	CaCo-2 at 24, and 48, LoVo and 72 h

Note: \uparrow = increase; \downarrow = decrease.

^{*} p < 0.05.

Abbreviations: ACSM = American College of Sports Medicine; ADT = androgen deprivation therapy; BMI = body mass index; CON = control; HDL = high-density lipoprotein; HIIT = high intensity interval training; HR = heart rate; INT = intervention; LDL = low-density lipoprotein; MICT = moderate intensity continuous training; N/C = no changes; PSA = prostate specific antigen; RM = repetition maximum; RPE = rating of perceived exertion; RT = resistance training; VO_{2max} = maximal oxygen uptake.

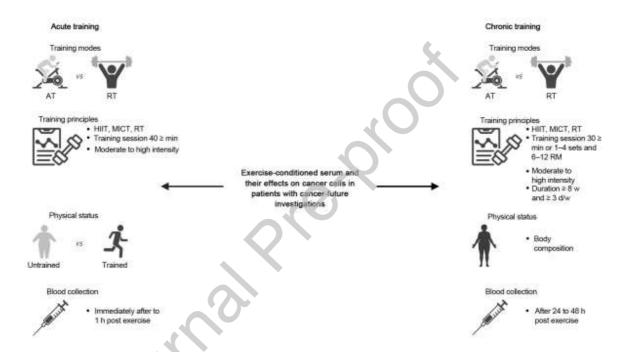


Fig. 1. Directions for future research for acute and chronic training interventions. Created with BioRender.com. AT = aerobic training; HIIT = high intensity interval training; MICT = moderate intensity continuous training; RM = repetition maximum; RT = resistance training; d/w = day per week; w = weeks.

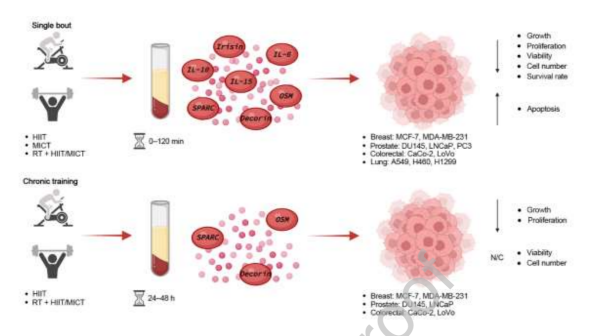


Fig. 2. Exercise-induced myokines in single bout and chronic training interventions and their effects on cancer cell lines. Created with BioRender.com. To be considered chronic, only training interventions > 4 weeks have been included. $\uparrow =$ increase; $\downarrow =$ decrease. HIIT = high intensity interval training; MICT = moderate intensity continuous training; N/C = no changes; RT = resistance training.

Graphical Abstract

