Breast Cancer Research and Treatment (2018) 169:93–103 https://doi.org/10.1007/s10549-018-4663-8

#### **CLINICAL TRIAL**



# Highly favorable physiological responses to concurrent resistance and high-intensity interval training during chemotherapy: the OptiTrain breast cancer trial

Sara Mijwel<sup>1,2</sup> • Malin Backman<sup>2,8</sup> • Kate A. Bolam<sup>2,3</sup> • Emil Olofsson<sup>4</sup> • Jessica Norrbom<sup>1</sup> • Jonas Bergh<sup>5,6</sup> • Carl Johan Sundberg<sup>1,7</sup> • Yvonne Wengström<sup>2,8</sup> • Helene Rundqvist<sup>4</sup>

Received: 4 January 2018 / Accepted: 5 January 2018 / Published online: 18 January 2018 © The Author(s) 2018. This article is an open access publication

#### **Abstract**

**Background** Advanced therapeutic strategies are often accompanied by significant adverse effects, which warrant equally progressive countermeasures. Physical exercise has proven an effective intervention to improve physical function and reduce fatigue in patients undergoing chemotherapy. Effects of high-intensity interval training (HIIT) in this population are not well established although HIIT has proven effective in other clinical populations. The aim of the OptiTrain trial was to examine the effects of concurrent resistance and high-intensity interval training (RT-HIIT) or concurrent moderate-intensity aerobic and high-intensity interval training (AT-HIIT), to usual care (UC) on pain sensitivity and physiological outcomes in patients with breast cancer during chemotherapy.

**Methods** Two hundred and forty women were randomized to 16 weeks of RT-HIIT, AT-HIIT, or UC. Outcomes: cardiorespiratory fitness, muscle strength, body mass, hemoglobin levels, and pressure-pain threshold.

**Results** Pre- to post-intervention, RT-HIIT (ES = 0.41) and AT-HIIT (ES = 0.42) prevented the reduced cardiorespiratory fitness found with UC. Handgrip strength (surgery side: RT-HIIT vs. UC: ES = 0.41, RT-HIIT vs. AT-HIIT: ES = 0.28; non-surgery side: RT-HIIT vs. UC: ES = 0.35, RT-HIIT vs. AT-HIIT: ES = 0.22) and lower-limb muscle strength (RT-HIIT vs. UC: ES = 0.66, RT-HIIT vs. AT-HIIT: ES = 0.23) were significantly improved in the RT-HIIT. Increases in body mass were smaller in RT-HIIT (ES = -0.16) and AT-HIIT (ES = -0.16) versus UC. RT-HIIT reported higher pressure-pain thresholds than UC (trapezius: ES = 0.46, gluteus: ES = 0.53) and AT-HIIT (trapezius: ES = 0.30).

**Conclusion** Sixteen weeks of RT-HIIT significantly improved muscle strength and reduced pain sensitivity. Both exercise programs were well tolerated and were equally efficient in preventing increases in body mass and in preventing declines in cardiorespiratory fitness. These results highlight the importance of implementing a combination of resistance and high-intensity interval training during chemotherapy for women with breast cancer.

Keywords Breast cancer · High-intensity interval training · Chemotherapy · Pressure-pain threshold

- Sara Mijwel sara.mijwel@ki.se
- Department of Physiology and Pharmacology, Karolinska Institutet, Stockholm, Sweden
- Department of Neurobiology, Care Sciences and Society, Karolinska Institutet, Stockholm, Sweden
- School of Human Movement and Nutrition Sciences, The University of Queensland, Brisbane, Australia
- Department of Cell and Molecular Biology, Karolinska Institutet, Stockholm, Sweden

- Department of Oncology and Pathology Cancer Center Karolinska, Karolinska Institutet, Stockholm, Sweden
- <sup>6</sup> Cancer Theme, Karolinska University Hospital, Stockholm, Sweden
- Department of Learning, Informatics, Management and Ethics, Karolinska Institutet, Stockholm, Sweden
- Division of Nursing, Karolinska University Hospital, Stockholm, Sweden

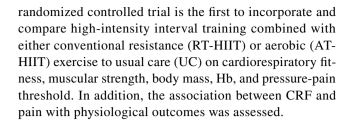


#### Introduction

The five-year relative breast cancer survival rate is continuously improving and approaching 90% in many countries [1]. Effective adjuvant therapies have the potential to substantially reduce recurrence and mortality but presents with significant side effects and quality of life consequences. The most commonly reported symptom during and after treatment for breast cancer is cancer-related fatigue (CRF), a multifactorial distress which together with treatment-induced pain contributes to physiological dysfunction [2]. The marked deterioration of the individual's physical capacity after a cancer diagnosis is due to direct effects of chemotherapy on physical function as well as general reductions of activity levels and is associated with decreased hemoglobin levels (Hb) [3], reduced cardiorespiratory fitness, and declines in muscle strength [4].

Exercise interventions for patients undergoing curative breast cancer therapy have been well tolerated and have shown positive effects on physical function and CRF [4]. Previous findings from the OptiTrain trial [5] showed favorable effects of exercise training on CRF and self-reported pain; however, whether muscle strength and/or cardiorespiratory fitness is associated with CRF and/or pain remains to be elucidated. Moreover, patients with breast cancer usually exhibit hypersensitivity to pressure pain compared with healthy control subjects [6]. Chronic aerobic exercise training in healthy individuals has shown positive effects on pain tolerance but not on pressure-pain threshold (PPT) [7]. However, the hypoalgesic effect of acute exercise is well established [8].

Despite the increasing use of taxane-based treatment in addition to anthracycline-based treatment, and that taxanerelated toxicities differ from those of taxane-free therapies [9], few studies have evaluated whether the exercise response differs depending on the type of chemotherapy [10]. Evidence-based exercise regimens with identified benefits are highly warranted by health care professionals and patients. In healthy individuals and in several pathological conditions, high-intensity interval training (HIIT) provides significant, time-efficient improvements in cardiorespiratory fitness [11] and pilot studies present HIIT as a safe training strategy also in patients with breast cancer [12]. In addition, HIIT may induce beneficial neuromuscular adaptations [13] and anti-inflammatory effects [14], both proposed as mechanisms contributing to fatigue [15]. The physiological outcomes of HIIT and whether it best combines with aerobic or resistance exercise during chemotherapy are currently unknown. While aerobic exercise provides established QoL improvements, recent studies emphasize incorporation of resistance training to address loss of muscle strength and function [4]. This in-clinic,



#### **Materials and methods**

#### **Participants**

Between March 2013 and July 2016, two hundred and forty patients at the Karolinska University Hospital (Stockholm, Sweden) were eligible for and accepted participation in the OptiTrain study (NCT02522260) [16]. The flow of participants through the trial has been reported elsewhere [5]. Inclusion criteria were Swedish-speaking women, 18-70 years old, breast cancer stage I-IIIa, scheduled to undergo chemotherapy consisting of anthracyclines, taxanes, or a combination of the two. Exclusion criteria were cardiac pathologies (assessed by routine electrocardiogram and a questionnaire), major psychiatric disorders, or other concurrent malignant diseases. Ethical approval was obtained from the Regional Ethical Review Board in Stockholm (Dnr 2012/1347-31/1, 2012/1347-31/2, 2013/632-32, 2014/408-32). All participants gave written informed consent prior to enrollment.

#### Randomization and blinding

The participants were randomly allocated by the Clinical Studies Unit at Radiumhemmet, Karolinska University Hospital (Stockholm, Sweden) to either RT-HIIT, AT-HIIT, or UC at a 1:1:1 ratio using a computer-generated assignment program blinded to the research team. Participants, exercise-supervisors, and outcome-assessors were not masked to group allocation. The first measurement took place 1 week prior to participants' second chemotherapy session and intervention groups (RT-HIIT and AT-HIIT) commenced the exercise training 3 days after the second chemotherapy session. Due to the limited time to perform ECG and baseline measurements before the first chemotherapy session, the intervention was initiated after the second chemotherapy session.

#### **Exercise training intervention**

Exercise groups trained twice per week for 16 weeks at the Karolinska University Hospital rehabilitation center. All sessions were supervised by an exercise physiologist or oncology nurse to ensure safety, correct technique, progression,



and encourage adherence to exercise protocols. The program was extended for participants with delays in chemotherapy (RT-HIIT, n = 8, range: 15–35 days; AT-HIIT, n = 5, range 13–32 days). The details of the training protocols have been published elsewhere [5, 16]. In brief, the RT-HIIT group completed both resistance and high-intensity interval exercise during each session. Participants performed 2-3 sets of 8–12 repetitions at an intensity of 80% of the patients' estimated 1-repetition maximum. To ensure progressive overload, loads were adjusted throughout the training period when participants were able to perform more than 12 repetitions. The AT-HIIT group commenced with 20 min of moderate-intensity continuous aerobic exercise at a rating of perceived exertion (RPE) of 13–15 on the Borg scale [17]. Both RT-HIIT and AT-HIIT concluded with  $3 \times 3$  min bouts of HIIT at an RPE of 16-18 interspersed with ~ 1 min of recovery.

#### **Outcomes**

The primary outcome in the OptiTrain trial is cancer-related fatigue (CRF) [16], and findings from both the unidimensional European Organization for Research and Treatment for Cancer Quality of Life Questionnaire (EORTC-QLQ-C30), and multidimensional Piper Fatigue Scale have previously been published [5]. In the present study, we report findings on physiological outcomes including muscle strength, cardiorespiratory fitness, pressure-pain threshold, hemoglobin levels, and body mass. Moreover, assessments of associations between changes in physiological outcomes and changes in self-reported cancer-related fatigue as well as pain as assessed by the EORTC-QLQ-C30 were performed. (The EORTC-QLQ-C30 tool is sensitive to change in patients receiving chemotherapy [18]).

Handgrip strength was assessed by a hydraulic hand dynamometer (JAMAR, SAEHAN Corporation, Changwon, S. Korea), and lower-limb muscle strength by isometric mid-thigh pull (Baseline leg dynamometer, Fabrication Enterprises Inc., White Plains, NY, USA). Cardiorespiratory fitness, measured as predicted peak oxygen uptake (VO<sub>2peak</sub>), was assessed by the Astrand-Rhyming submaximal cycle test [19], which has been validated for this population [20]. Hb was measured in venous blood (Clinical Studies Unit, Karolinska University Hospital, Stockholm, Sweden). Pressurepain threshold (PPT) was measured bilaterally on the middle trapezius and gluteus muscles with an electronic algometer (Somedic Sales AB, Hörby, Sweden). No analgesics were taken 24 h prior to measurement. The applied pressure was at a rate of approximately 50 kPa/s by a 1 cm<sup>2</sup> probe. The mean PPT from the bilateral measurements at each anatomical site was used for analysis. All outcomes were measured at baseline and at 16 weeks.

#### **Activity measures**

At baseline, objective activity patterns were assessed by an accelerometer (model GT3X ActiGraph® Corp, Pensacola, Florida, USA) and analyzed using validated wear-time specifications and cut-offs for adults [21]. Calculation of attendance and adherence to the exercise regimen has been described elsewhere [5].

#### **Statistics**

With fatigue as the primary outcome measure in the Opti-Train trial, a sample size of 65 patients per group was required, based on an effect size of 0.53, using a two-sided ANOVA test with a power of 80% at 5% significance level. We aimed to recruit 80 participants into each group to account for an expected 20% attrition rate. Variables were visually checked for normality through QQ-plots and histograms and were found to be normally distributed. Baseline medical and demographic characteristics of each group were summarized using descriptive statistics. Exact  $\chi^2$  tests were used for categorical variables. For between-group analyzes, we used analysis of covariance, adjusted for baseline values. Repeated-measures ANOVA was used to test for within-group differences. Standardized effect sizes (ES) were calculated and interpreted as described previously [5, 22]. An intention-to-treat approach was used, and missing data (9%) were imputed using the expectation maximization method after being determined to be "missing completely at random" [23]. The expectation-maximization algorithm is based on group change and the individual baseline score. For the outcome predicted VO<sub>2peak</sub>, participants receiving betaadrenergic blocking agents (n = 2), or not reaching the target heart rate of 120–170 beats per min (n = 2), were excluded from the analysis. For all outcomes, a subgroup analysis was performed, and data were stratified into patients receiving taxanes (TAX) or not receiving taxanes (non-TAX). Post hoc comparisons were conducted with Bonferroni correction for three pairwise comparisons between RT-HIIT, AT-HIIT, and UC. Pearson's coefficient of correlation (r) was used to evaluate the association between changes in physiological outcomes and changes in self-reported CRF and pain. All tests were two-tailed and a p value of < 0.05 was required for significance.

#### Results

All groups were balanced at baseline (Table 1) [5], and no significant baseline differences were found between participants who dropped out versus participants who completed the study. Attendance and adherence to the training program have previously been reported [5]. There was no difference



Table 1 Participant characteristics at baseline

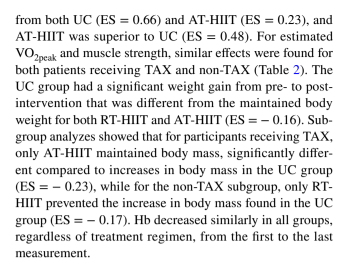
	RT-HIIT $n = 74$	AT-HIIT $n = 72$	UC n = 60	
	Mean ± SD	Mean ± SD	Mean ± SD	
Age (years)	$52.7 \pm 10.3$	$54.4 \pm 10.3$	$52.6 \pm 10.2$	
Body mass (kg)	$68.7 \pm 11.3$	$67.7 \pm 13.0$	$69.1 \pm 11.0$	
Height	$165.7 \pm 6.7$	$165.3 \pm 6.6$	$166.4 \pm 7.0$	
SED (% of daily wear time)	$63.7 \pm 7.7$	$65.6 \pm 6.2$	$66.6 \pm 7.2$	
MVPA (% of daily wear time)	$9.6 \pm 2.8$	$8.3 \pm 2.8$	$8.5 \pm 4.3$	
	n (%)	n (%)	n (%)	
Married or partnered	60.6	59.7	69.5	
University completed	67.6	64.7	66.0	
Current smokers	4.3	5.9	5.2	
Employed	74.6	86.8	79.7	
Postmenopausal	51.4	63.9	61.7	
Tumor profile				
Triple negative	14.9	11.0	16.7	
HER2+, ER+/-	21.6	30.2	20.0	
HER2-, ER+	62.2	58.9	61.6	
HER2-, ER-	1.4	0.0	1.7	
Anthracycline-based therapy	40.6	37.0	41.7	
Taxane-based therapy	59.4	63.0	58.3	

SD standard deviation, RT-HIIT resistance and high-intensity interval training, AT-HIIT moderate-intensity aerobic and high-intensity interval training, UC usual care, MVPA objectively measured moderate- to vigorous intensity physical activity, SED objectively measured sedentary behavior

in attendance to exercise sessions for participants receiving TAX compared to those receiving non-TAX. No adverse events prompting medical attention occurred during exercise sessions.

### Cardiorespiratory fitness, muscle strength, body mass, and Hb

Changes in cardiorespiratory fitness, muscle strength, body mass, and Hb are shown in Fig. 1. Over the intervention, a significant decline in predicted  $VO_{2peak}$  was found for UC that was significantly different compared to unchanged levels for both RT-HIIT (ES = 0.41) and AT-HIIT (ES = 0.42). Participants in the UC group had significant handgrip strength losses, while significant handgrip muscle strength gains were found for the RT-HIIT group. Correspondingly, RT-HIIT was superior to both UC and AT-HIIT for handgrip strength (surgery side: RT-HIIT vs. UC: ES = 0.41, RT-HIIT vs. AT-HIIT: ES = 0.28; non-surgery side: RT-HIIT vs. UC: ES = 0.35, RT-HIIT vs. AT-HIIT: ES = 0.22). Both RT-HIIT and AT-HIIT significantly improved lowerlimb muscle strength. The improved isometric mid-thigh pull strength test for RT-HIIT was significantly different



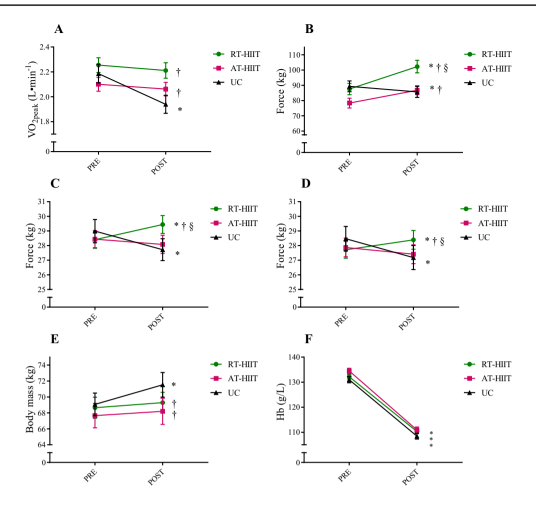
#### Pressure-pain threshold

Changes in PPT are shown in Fig. 2 and results from the subgroup analysis are shown in Table 2. Over the intervention, PPT measurements showed significant reductions for UC at both trapezius and gluteus muscles while RT-HIIT showed an increased PPT at the trapezius muscle, different when compared to UC (ES = 0.46) and AT-HIIT (ES = 0.30). PPT at the gluteus muscle favored RT-HIIT (ES = 0.53) compared to UC. For patients receiving non-TAX treatment, the UC group displayed significant reductions in PPT for both trapezius and gluteus muscles. For trapezius muscle, PPT for RT-HIIT was superior to both UC (ES = 1.17) and AT-HIIT (ES = 0.45), although AT-HIIT was still significantly different compared to UC (ES = 0.45). Significant differences were also found for PPT at the gluteus muscle favoring both RT-HIIT (ES = 1.02) and AT-HIIT (ES = 0.61) compared to UC. Patients receiving TAX treatment did not change in PPT and no between-group differences were found.

## Associations between physiological outcomes and self-reported cancer-related fatigue (CRF) and pain

Associations between changes in physiological outcomes and changes in self-reported cancer-related fatigue and pain are shown in Table 3. A weak inverse correlation was found between change in self-reported CRF and change in lower-limb strength (r = -0.28, p < 0.001), and between change in self-reported CRF and change in PPT at the gluteus muscle (r = -0.24, p = 0.001). No significant associations were found between change in self-reported fatigue and change in handgrip strength (surgery side r = -0.12, p = 0.09; non-surgery side r = -0.13, p = 0.06), or change in self-reported fatigue and change in cardiorespiratory fitness (r = -0.10, p = 0.16). Similar associations were found for change in self-reported CRF measured by the Piper Fatigue





**Fig. 1** Effects of concurrent resistance and high-intensity interval training (RT-HIIT) and moderate-intensity aerobic and high-intensity interval training (AT-HIIT) versus usual care (UC) on physiological outcomes:  $\mathbf{a}$  estimated  $VO_{2peak}, \mathbf{b}$  isometric mid-thigh pull,  $\mathbf{c}$  handgrip strength surgery side,  $\mathbf{d}$  handgrip strength non-surgery side,

**e** body mass, and **f** hemoglobin levels. \*p < 0.05 at post versus pre measurement; †p < 0.05 compared to UC; \*p < 0.05 between RT-HIIT and AT-HIIT. Data is presented as mean and standard error of the mean. No statistically significant differences were found at baseline between groups

Scale and change in physiological outcome measures (data not shown). Moreover, we found a significant correlation between changes in lower-limb muscle strength and changes in PPT at trapezius (r = 0.17, p = 0.03) and gluteus muscles (r = 0.16, p = 0.04), as well as between change in handgrip strength (non-surgery side) and change in PPT trapezius (r = 0.17, p = 0.02). Self-reported CRF was significantly associated with self-reported pain (r = 0.37, p < 0.001); however, no associations were found between physiological outcomes and self-reported pain.

#### **Discussion**

This is the first randomized controlled trial to include highintensity interval training (HIIT) in two different exercise regimens in patients with breast cancer during chemotherapy. Results from this trial provide important updates in our understanding of potential management strategies for treatment-induced fatigue and physical dysfunction in this relatively large subgroup of cancer patients. This is also the first trial to investigate the ability of exercise training to prevent chemotherapy-induced hyperalgesia. Concurrent resistance and high-intensity interval training (RT-HIIT) provided significant beneficial effects in terms of counteracting cancer-related physical and total fatigue [5], reducing anthracycline-induced pain-hypersensitivity, and improving muscle strength in patients undergoing adjuvant treatment for breast cancer.

In concordance with previous findings, aerobic training resulted in sustained cardiorespiratory fitness [4]. Interestingly, cardiorespiratory fitness in the RT-HIIT group was maintained despite only a total of 9 min of HIIT per session, contradicting the results from Courneya et al. [24], where only the higher dose aerobic training (50–60 min, 3 days/week, 60–75% of  $VO_{2peak}$ ) resulted in a maintained



**Table 2** Mean values and standard deviations at the first measurement and at 16 weeks, and between-group differences for all outcome measures for participants receiving taxane- and non-taxane-based treatment

Outcome	Arm	N	Mean SD		Mean SD		p value
			Pre-interve	ntion	Post-intervention		F /
Estimated VO <sub>2peak</sub> Taxanes (L min <sup>-1</sup> )	RT-HIIT	43	2.24	0.53	2.23	0.60	0.845
Estimated VO <sub>2peak</sub> Taxanes (E mm )	AT-HIIT	45	2.06	0.51	1.99	0.51	0.043
	UC	30	2.27	0.52	1.95	0.48	< 0.001
Estimated VO <sub>2peak</sub> Non-taxanes (L min <sup>-1</sup> )	RT-HIIT	30	2.27	0.48	2.18	0.43	0.140
Estimated VO <sub>2peak</sub> (Von-taxanes (E min )	AT-HIIT	25	2.17	0.40	2.19	0.43	0.709
	UC	21	2.06	0.53	1.92	0.57	0.709
Isometric mid-thigh pull Taxanes (kg)	RT-HIIT	38	89.35	29.07	105.67	36.39	< 0.001
isometre mid-ungi pun Taxanes (kg)	AT-HIIT	40	75.68	23.56	83.94	21.83	< 0.001
	UC	30	90.07	20.08	85.47	21.92	0.153
Isometric mid-thigh pull Non-taxanes (kg)	RT-HIIT	27	84.26	30.52	96.27	26.91	0.133
isometric find-ungil pun ivon-taxanes (kg)	AT-HIIT	22	83.19	27.61	91.95	24.18	0.001
	UC	21	88.25	31.77	85.96	31.10	0.396
Hondowin gurgory side Toyones (leg)			28.16	4.98			0.003
Handgrip surgery side Taxanes (kg)	RT-HIIT	43	27.73		29.49	5.33	0.687
	AT-HIIT	45 25	29.40	5.09	27.56	5.60	0.087
Hardada and a state of the Tourist (La)	UC	35		6.26	28.31	5.81	
Handgrip non-surgery side Taxanes (kg)	RT-HIIT	43	27.54	4.69	28.47	5.33	0.012
	AT-HIIT	45	27.32	5.81	27.06	6.10	0.570
W 1 : (1 N )	UC	35	28.43	6.70	27.23	6.52	0.011
Handgrip surgery side Non-taxanes (kg)	RT-HIIT	31	28.73	5.18	29.35	5.26	0.169
	AT-HIIT	27	29.63	4.59	28.94	4.72	0.113
	UC	25	28.42	6.09	26.89	5.76	0.033
Handgrip non-surgery side Non-taxanes (kg)	RT-HIIT	31	27.95	5.32	28.27	5.90	0.540
	AT-HIIT	27	28.78	4.71	28.00	4.28	0.139
	UC	25	28.52	6.34	27.12	6.20	0.064
Body mass Taxanes (kg)	RT-HIIT	43	67.16	9.96	68.31	10.19	0.008
	AT-HIIT	45	67.40	10.97	67.61	10.93	0.514
	UC	35	68.77	10.02	71.46	10.46	< 0.001
Body mass Non-taxanes (kg)	RT-HIIT	31	70.72	12.89	70.64	12.44	0.868
	AT-HIIT	27	68.10	16.05	69.18	18.06	0.073
	UC	25	69.48	12.38	71.62	13.94	0.002
Hemoglobin Taxanes (g/L)	RT-HIIT	43	133.42	7.04	108.68	7.33	< 0.001
	AT-HIIT	45	135.00	7.96	108.85	9.61	< 0.001
	UC	35	131.29	9.82	106.60	7.91	< 0.001
Hemoglobin Non-taxanes (g/L)	RT-HIIT	31	130.42	7.88	112.55	7.36	< 0.001
	AT-HIIT	27	133.59	10.59	114.39	9.68	< 0.001
	UC	25	130.56	9.43	111.05	9.97	< 0.001
PPT trapezius Taxanes (kPa)	RT-HIIT	41	424.67	159.92	441.65	163.41	0.374
	AT-HIIT	45	396.12	135.40	381.91	109.08	0.427
	UC	35	395.39	139.88	401.13	127.37	0.681
PPT gluteus Taxanes (kPa)	RT-HIIT	41	408.29	162.91	434.36	129.89	0.214
	AT-HIIT	45	410.69	195.77	400.39	128.21	0.631
	UC	35	434.66	153.58	415.69	132.43	0.349
PPT trapezius Non-taxanes (kPa)	RT-HIIT	29	411.43	115.63	457.95	115.63	0.053
	AT-HIIT	25	421.78	185.3	398.95	146.09	0.365
	UC	24	410.52	127.52	314.89	102.69	< 0.001
PPT gluteus Non-taxanes (kPa)	RT-HIIT	29	437.86	114.38	450.39	142.29	0.503
	AT-HIIT	25	443.58	201.15	436.37	173.63	0.773
	UC	24	420.83	128.67	309.84	130.19	< 0.001



Table 2 (continued)

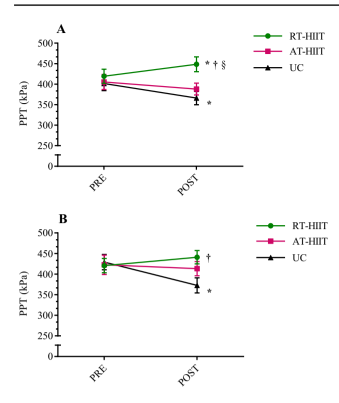
Outcome	Adjusted mean change (95% CI):RT-HIIT versus UC	p value	ES	Adjusted mean change (95% CI):AT-HIIT versus UC	p value	ES	Adjusted mean change (95% CI):RT-HIIT versus AT-HIIT	p value	ES
Estimated VO <sub>2peak</sub> Taxanes (L min <sup>-1</sup> )	0.30 (0.13 to 0.48)	< 0.001	0.59	0.22 (0.04 to 0.39)	0.013	0.49	0.09 (- 0.07 to 0.25)	0.554	0.12
$\begin{array}{c} \text{Estimated VO}_{2\text{peak}}\text{Non-}\\ \text{taxanes}\;(\text{L}\;\text{min}^{-1}) \end{array}$	0.10 (- 0.08 to 0.29)	0.516	0.10	0.19 (0.01 to 0.38)	0.047	0.13	- 0.09 (- 0.26 to 0.09)	0.656	- 0.04
Isometric mid-thigh pull Taxanes (kg)	20.88 (11.92 to 29.85)	< 0.001	0.82	12.08 (2.98 to 21.18)	0.005	0.58	8.80 (0.26 to 17.34)	0.041	0.30
Isometric mid-thigh pull Non-taxanes (kg)	13.52 (4.13 to 22.90)	0.002	0.46	10.05 (0.20 to 19.90)	0.044	0.37	3.47 (- 5.79 to 12.72)	1.000	0.11
Handgrip surgery side Taxanes (kg)	2.30 (0.80 to 3.80)	0.001	0.43	0.76 (- 0.73 to 2.24)	0.661	0.16	1.55 (0.15 to 2.94)	0.025	0.30
Handgrip non-surgery side Taxanes (kg)	2.08 (0.59 to 3.57)	0.003	0.37	0.87 (- 0.61 to 2.34)	0.471	0.15	1.21 (- 0.18 to 2.60)	0.109	0.22
Handgrip surgery side Non-taxanes (kg)	2.20 (0.48 to 3.91)	0.007	0.38	1.01 (- 0.77 to 2.79)	0.509	0.16	1.19 (- 0.50 to 2.87)	0.264	0.27
Handgrip non-surgery side Non-taxanes (kg)	1.63 (- 0.31 to 3.57)	0.128	0.30	0.66 (- 1.34 to 2.66)	1.000	0.11	0.97 (- 0.93 to 2.88)	0.642	0.22
Body mass Taxanes (kg)	- 1.57 (- 3.19 to 0.06)	0.062	- 0.15	- 2.50 (- 4.10 to - 0.89)	0.001	- 0.23	0.93 (- 0.59 to 2.45)	0.420	0.09
Body mass Non-taxa- nes (kg)	- 2.30 (- 4.15 to - 0.44)	0.010	- 0.17	- 0.99 (- 2.91 to 0.93)	0.628	- 0.07	- 1.30 (- 3.13 to 0.52)	0.254	- 0.08
Hemoglobin Taxanes (g/L)	1.07 (- 3.08 to 5.22)	1.000	0.01	0.49 (- 3.66 to 4.64)	1.000	0.17	0.58 (- 3.30 to 4.46)	1.000	0.19
Hemoglobin Non- taxanes (g/L)	1.56 (- 3.85 to 6.96)	1.000	0.19	2.14 (- 3.49 to 7.76)	1.000	0.03	- 0.58 (- 5.93 to 4.77)	1.000	- 0.14
PPT trapezius Taxanes (kPa)	21.7 (- 33.0 to 76.4)	1.000	0.07	- 19.7 (- 73.1 to 33.7)	1.000	- 0.13	41.4 (- 9.9 to 92.7)	0.158	0.19
PPT gluteus Taxanes (kPa)	31.6 (- 23.8 to 87.0)	0.507	0.27	- 3.6 (- 57.8 to 50.7)	1.000	- 0.06	35.2 (- 16.7 to 87.0)	0.308	0.18
PPT trapezius Non- taxanes (kPa)	142.5 (77.6 to 207.4)	< 0.001	1.17	77.4 (10.2 to 144.7)	0.019	0.45	65.1 (0.9 to 129.3)	0.046	0.45
PPT gluteus Non- taxanes (kPa)	129.4 (57.9 to 198.8)	< 0.001	1.02	110.3 (37.3 to 184.3)	0.001	0.61	18.1 (- 51.5 to 87.7)	1.000	0.12

SD standard deviation, ES, effect size, PPT pressure-pain threshold,  $VO_{2peak}$  peak oxygen consumption, RT-HIIT resistance and high-intensity interval training, UC usual care, p < 0.05 is highlighted in bold

cardiorespiratory fitness. This suggests that HIIT provides an effective and time-saving training strategy resulting in the same beneficial preservation of cardiorespiratory fitness as high-volume aerobic training, similar to findings in healthy individuals [11]. Results from the subgroup analyses showed that the declines in  $VO_{2peak}$  (L min<sup>-1</sup>) for the UC group were more severe for those receiving taxanes (-14%) compared to those not receiving taxanes (-7%). A previous exercise trial including periodic high and low-intensity components as well as relaxation exercise training reported on the challenges of preserving cardiorespiratory fitness during taxanebased treatment and found significant declines in cardiorespiratory fitness for both exercise and control groups [25]. In the current trial, both exercise interventions were able to counteract the decline in estimated  $VO_{2peak}$  in patients receiving taxanes, suggesting the importance of HIIT performed throughout the treatment period.

We previously showed that UC reported an increase in CRF measured by the EORTC-OLO-C30 over the intervention, while no increase was found in the RT-HIIT and AT-HIIT groups [5]. Moreover, RT-HIIT was able to counteract the increase in physical and total CRF found in the UC group as measured by the Piper Fatigue Scale [5]. This is in line with previous trials showing that exercise training during chemotherapy was able to counteract CRF [4, 26]. Suggested mechanisms for CRF comprise reduced cardiorespiratory fitness and Hb [3]. Given the general decline in Hb and that aerobic fitness was equally maintained in both exercise groups, the physical CRF component is most likely multifactorial. It is currently unknown whether CRF is mainly induced by central or peripheral factors; however, recent studies point toward CRF being centrally mediated [27]. Studies on patients with advanced stage cancer suffering from CRF showed that these patients were unable to voluntarily recruit as much muscle as healthy controls [27],





**Fig. 2** Pressure-pain thresholds (PPT) for a trapezius muscle, **b** gluteus muscle. *RT-HIIT* resistance and high-intensity interval training, *AT-HIIT* moderate-intensity aerobic and high-intensity interval training, *UC* usual care. \*p < 0.05 at post versus pre measurement; †p < 0.05 compared to UC; \*p < 0.05 between RT-HIIT and AT-HIIT. Data is presented as mean and standard error of the mean. No statistically significant differences were found at baseline between groups

and had an earlier motor task failure [28]. The resistance-training component of RT-HIIT likely induced central neuro-muscular adaptations through increased motor neuron firing frequencies [29]. In line with this, we found a weak inverse association between change in self-reported CRF and change in lower-limb muscle strength (r = -0.28; p < 0.001). This is in concordance with findings from a study on patients with

Table 3 Pearson productmoment correlations between change in cancer-related fatigue and pain (EORTC-QLQ-C30) and change in physical outcome

found between maximal handgrip strength and self-reported
CRF in breast cancer survivors [31]. We did not however
find associations between handgrip strength and cardiores-
piratory fitness measures with changes in self-reported CRF.
A study by Thorsen et al. also failed to show associations
between cardiorespiratory fitness (assessed by the same sub-
maximal cycle test as in our trial) and self-reported CRF
[32]. It may be speculated that stronger associations would
be found between time to fatigue assessments and CRF.
However, based on the current assessments, our findings
indicate that lower-limb muscle strength may be an impor-
tant underlying component of the multifactorial symptom
CRF. Given that CRF can persist for up to ten years after a
cancer diagnosis, and predicts shorter survival [3], imple-
menting combined high-intensity intervals with resistance
exercise regimens, similar to RT-HIIT, during chemotherapy
may be critical for reducing the burden of this symptom.
In agreement with previous resistance-training inter-
vantions [4] we found a moderate affect size (0.66) for

advanced cancer [30]. Weak negative associations have been

ventions [4], we found a moderate effect size (0.66) for changes in lower-limb muscle strength with RT-HIIT compared to UC. Importantly, lower-limb muscle strength was also improved in the AT-HIIT group, not commonly reported with conventional aerobic exercise, supporting a role for HIIT in inducing neuromuscular adaptations. The handgrip strength improvements displayed by the RT-HIIT group offers a prognostic value and is an important correlate of health in survivors of breast cancer [31]. In the current study, strength gains were found with RT-HIIT regardless of chemotherapy regimen, and even slightly greater improvements for those receiving taxane-treatment for lower-limb muscle strength (+ 18.3%) compared to those on taxane-free treatment (+ 14.2%). Our findings are in contrast to those of Courneya [10], reporting that participants in a combined resistance and aerobic training group receiving taxanetreatment did not gain as much muscle strength as those not receiving taxanes. The discrepancy is unclear; however, despite that taxanes induce higher rates of myalgia, arthralgia, and neurosensory effects [9], a recent preclinical study

Outcome	1	2	3	4	5	6	7	8
Self-reported cancer-related fatigue	_							
2. Self-reported pain	0.37*	_						
3. Cardiorespiratory fitness	-0.10	0.00	_					
4. Lower-limb muscle strength	- 0.28*	-0.07	0.25*	-				
5. Handgrip strength (surgery side)	-0.12	-0.07	0.15*	0.23*	_			
6. Handgrip strength (non-surgery side)	-0.13	-0.05	0.22*	0.22*	0.67*	-		
7. Pressure-pain threshold (trapezius muscle)	-0.09	0.03	0.01	0.17*	0.10	0.17*	-	
8. Pressure-pain threshold (gluteus muscle)	- 0.24*	- 0.02	0.03	0.16*	- 0.02	0.11	0.64*	-

 $\it EORTC-QLQ-C30$  European Organization for Research and Cancer Treatment for Cancer Quality of Life Questionnaire, \* $\it p < 0.05$ 



measures

showed no impairments in muscle function in response to taxanes [33].

Women with breast cancer commonly gain about 5 kg body weight during chemotherapy, and few return to their pre-diagnosis weight [34]. Here, no weight gain was observed in either of the intervention groups, except for the RT-HIIT group that received taxanes. The UC group gained body weight regardless of chemotherapy regimen. Since weight gain is associated with comorbidities and recurrence [34], maintaining pre-diagnosis weight is of major importance.

Physical fitness is associated with a high pressure-pain threshold (PPT) [35]. In the current study, the RT-HIIT intervention could completely compensate for the reduced PPT found in the UC group over the intervention. Notably, subgroup analyses showed that the patients receiving taxanes displayed no hyperalgesia over time, despite taxanes being more neurotoxic and associated with higher levels of self-reported pain compared to non-taxane treatments [36]. We speculate that taxanes may cause increased numbness leading to a blunted PPT response. In line with this, loss of vibratory perception has been reported with taxane therapies [37]. Moreover, we found no associations between selfreported pain and PPT. Our findings are in line with a previous study showing no associations between self-reported pain and heat pain threshold in healthy participants [38]. The same study noted that those who reported more pain were found to be more anxious than those who reported less pain, while objectively measured heat pain thresholds did not follow the same pattern. Given that a cancer diagnosis is accompanied by feelings of depression, fear, and anxiety [39], it may be speculated that the participants in our study were influenced by negative affect/anxiety that might have influenced their subjective feeling of pain, which could explain the lack of association between self-reported pain and PPT in the current study as well.

Of note, we found significant weak-to-moderate associations between changes in PPT at the gluteus and trapezius muscles and changes in lower-limb muscle strength as well as between changes in PPT at the trapezius muscle and changes in handgrip strength, indicating that muscle strength/function may be of particular importance in preserving objectively measured pain sensitivity. In contrast, a recent study assessing associations between PPT and muscle strength found no significant correlations [40].

The OptiTrain study is a sufficiently powered, supervised, in-clinic, randomized intervention trialing two types of progressive exercise regimens with validated measures. Our attendance rates are within the range commonly reported in exercise trials [4]. Limitations comprise that the first assessment was performed after one cycle of chemotherapy; however, this also provides some benefit by excluding acute effects of chemotherapy from analyses comparing first and

last measurements. Moreover, a selection bias may have been introduced by the relatively large number of UC-patients declining participation directly after randomization. This aspect should generally be considered when drawing conclusions from exercise intervention studies.

#### **Conclusions**

A 16-week supervised concurrent resistance and high-intensity interval training intervention (RT-HIIT) significantly improved muscle strength, and prevented hyperalgesia. Moreover, the RT-HIIT intervention was as efficient as AT-HIIT in maintaining body mass and cardiorespiratory fitness in women with early breast cancer receiving adjuvant chemotherapy. Findings from the current study also show that participants in the RT-HIIT group displayed similar beneficial effects from the exercise intervention regardless of receiving taxane or taxanefree treatment. This renders us to recommend that women receiving chemotherapy for primary breast cancer should be provided with knowledge of and access to a concurrent resistance and high-intensity interval training regimen.

Acknowledgements The authors would like to thank the OptiTrain participants, the staff at the oncological rehabilitation center (Karolinska University Hospital, Stockholm, Sweden) for use of facilities, biostatisticians Eva Hagel and Daniel Olsson (Karolinska Institutet) for statistical advice, and Daniele Cardinale at the Swedish Sports Confederation (Lidingö, Sweden) for technical support.

**Funding** This work was supported by the Swedish Cancer Society (130452 to Y.W.), the Cancer Society of Stockholm (131242 to Y.W.), the Swedish Cancer and Traffic Accident Foundation (F-C-001225 to Y.W.), and the Swedish Society for Medical Research (SLS 50514 to H.R.).

#### **Compliance with ethical standards**

Conflict of interest The authors declare no conflict of interest.

**Open Access** This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

#### References

- International Agency for Research on Cancer: Cancer fact sheets Breast cancer (2016). https://www.iarc.fr/. Accessed 9 Mar 2017
- Davis MP, Walsh D (2010) Mechanisms of fatigue. J Support Oncol 8(4):164–174
- Cramp F, Daniel J (2008) Exercise for the management of cancer-related fatigue in adults. Cochrane Database Syst Rev 2:CD006145. https://doi.org/10.1002/14651858.cd006145.pub2
- 4. Furmaniak AC, Menig M, Markes MH (2016) Exercise for women receiving adjuvant therapy for breast cancer. Cochrane



- Database Syst Rev 9:5001. https://doi.org/10.1002/14651858.cd005001.pub3
- Mijwel S, Backman M, Bolam KA, Jervaeus A, Sundberg CJ, Margolin S, Browall M, Rundqvist H, Wengström Y (2017) Adding high-intensity interval training to conventional training modalities: optimizing health-related outcomes during chemotherapy for breast cancer: the OptiTrain randomized controlled trial. Breast Cancer Res Treat. https://doi.org/10.1007/s105 49-017-4571-3
- Caro-Moran E, Fernandez-Lao C, Diaz-Rodriguez L, Cantarero-Villanueva I, Madeleine P, Arroyo-Morales M (2016) Pressure pain sensitivity maps of the neck-shoulder region in breast cancer survivors. Pain Med 17(10):1942–1952. https://doi.org/10.1093/ pm/pnw064
- Jones MD, Booth J, Taylor JL, Barry BK (2014) Aerobic training increases pain tolerance in healthy individuals. Med Sci Sports Exerc 46(8):1640–1647. https://doi.org/10.1249/MSS.00000000 00000273
- Naugle KM, Fillingim RB, Riley JL 3rd (2012) A meta-analytic review of the hypoalgesic effects of exercise. J Pain 13(12):1139– 1150. https://doi.org/10.1016/j.jpain.2012.09.006
- Ho MY, Mackey JR (2014) Presentation and management of docetaxel-related adverse effects in patients with breast cancer. Cancer Manag Res 6:253–259. https://doi.org/10.2147/CMAR .S40601
- Courneya KS, McKenzie DC, Mackey JR, Gelmon K, Reid RD, Friedenreich CM, Ladha AB, Proulx C, Vallance JK, Lane K, Yasui Y, Segal RJ (2008) Moderators of the effects of exercise training in breast cancer patients receiving chemotherapy: a randomized controlled trial. Cancer 112(8):1845–1853. https://doi. org/10.1002/cncr.23379
- Karlsen T, Aamot IL, Haykowsky M, Rognmo O (2017) High intensity interval training for maximizing health outcomes. Prog Cardiovasc Dis 60(1):67–77. https://doi.org/10.1016/j.pcad.2017 .03.006
- Schmitt J, Lindner N, Reuss-Borst M, Holmberg HC, Sperlich B (2016) A 3 week multimodal intervention involving high-intensity interval training in female cancer survivors: a randomized controlled trial. Physiol Rep. https://doi.org/10.14814/phy2.12693
- Buchheit M, Laursen PB (2013) High-intensity interval training, solutions to the programming puzzle. Part II: anaerobic energy, neuromuscular load and practical applications. Sports Med 43(10):927–954. https://doi.org/10.1007/s40279-013-0066-5
- Steckling FM, Farinha JB, Santos DL, Bresciani G, Mortari JA, Stefanello ST, Courtes AA, Duarte T, Duarte MM, Moresco RN, Cardoso MS, Soares FA (2016) High Intensity Interval Training Reduces the Levels of Serum Inflammatory Cytokine on Women with Metabolic Syndrome. Exp Clin Endocrinol Diabetes 124(10):597–601. https://doi.org/10.1055/s-0042-111044
- LaVoy ECP, Fagundes CP, Dantzer R (2016) Exercise, inflammation, and fatigue in cancer survivors. Exerc Immunol Rev 22:82–93
- Wengström Y, Bolam KA, Mijwel S, Sundberg CJ, Backman M, Browall M, Norrbom J, Rundqvist H (2017) Optitrain: a randomised controlled exercise trial for women with breast cancer undergoing chemotherapy. BMC Cancer 17(1):100. https://doi. org/10.1186/s12885-017-3079-x
- Borg GA (1982) Psychophysical bases of perceived exertion. Med Sci Sports Exerc 14(5):377–381
- Uwer L, Rotonda C, Guillemin F, Miny J, Kaminsky MC, Mercier M, Tournier-Rangeard L, Leonard I, Montcuquet P, Rauch P, Conroy T (2011) Responsiveness of EORTC QLQ-C30, QLQ-CR38 and FACT-C quality of life questionnaires in patients with colorectal cancer. Health Qual Life Outcomes 9:70. https://doi.org/10.1186/1477-7525-9-70

- Åstrand I (1960) Aerobic work capacity in men and women with special reference to age. Acta Physiol Scand Suppl 49(169):1–92
- 21. Aguilar-Farias N, Brown WJ, Peeters GM (2014) ActiGraph GT3X+ cut-points for identifying sedentary behaviour in older adults in free-living environments. J Sci Med Sport 17(3):293–299. https://doi.org/10.1016/j.jsams.2013.07.002
- Morris SB (2007) Estimating effect sizes from pretest-posttest-control group designs. Organ Res Methods 11(2):364–386. https://doi.org/10.1177/1094428106291059
- Dong Y, Peng C-YJ (2013) Principled missing data methods for researchers. SpringerPlus 2:222. https://doi.org/10.1186/2193 -1801-2-222
- Courneya KS, McKenzie DC, Mackey JR, Gelmon K, Friedenreich CM, Yasui Y, Reid RD, Cook D, Jespersen D, Proulx C, Dolan LB, Forbes CC, Wooding E, Trinh L, Segal RJ (2013) Effects of exercise dose and type during breast cancer chemotherapy: multicenter randomized trial. JNCI J Natl Cancer Inst 105(23):1821–1832. https://doi.org/10.1093/jnci/djt297
- Møller T, Lillelund C, Andersen C, Bloomquist K, Christensen KB, Ejlertsen B, Nørgaard L, Wiedenbein L, Oturai P, Breitenstein U, Adamsen L (2015) The challenge of preserving cardiorespiratory fitness in physically inactive patients with colon or breast cancer during adjuvant chemotherapy: a randomised feasibility study. BMJ Open Sport Exerc Med 1(1):e000021. https://doi.org/10.1136/bmjsem-2015-000021
- 26. van Waart H, Stuiver MM, van Harten WH, Geleijn E, Kieffer JM, Buffart LM, de Maaker-Berkhof M, Boven E, Schrama J, Geenen MM, Meerum Terwogt JM, van Bochove A, Lustig V, van den Heiligenberg SM, Smorenburg CH, Hellendoorn-van Vreeswijk JA, Sonke GS, Aaronson NK (2015) Effect of low-intensity physical activity and moderate- to high-intensity physical exercise during adjuvant chemotherapy on physical fitness, fatigue, and chemotherapy completion rates: results of the PACES Randomized Clinical Trial. J Clin Oncol 33(17):1918–1927. https://doi.org/10.1200/jco.2014.59.1081
- Yavuzsen T, Davis MP, Ranganathan VK, Walsh D, Siemionow V, Kirkova J, Khoshknabi D, Lagman R, LeGrand S, Yue GH (2009) Cancer-related fatigue: central or peripheral? J Pain Symptom Manag 38(4):587–596. https://doi.org/10.1016/j.jpainsymman.2008.12.003
- Kisiel-Sajewicz K, Davis MP, Siemionow V, Seyidova-Khoshknabi D, Wyant A, Walsh D, Hou J, Yue GH (2012) Lack of muscle contractile property changes at the time of perceived physical exhaustion suggests central mechanisms contributing to early motor task failure in patients with cancer-related fatigue. J Pain Symptom Manag 44(3):351–361. https://doi.org/10.1016/j.jpainsymman.2011.08.007
- Aagaard P, Mayer F (2007) Neuronal adaptations to strength training. Deutsche zeitschrift f
  ür sportmedizin 58(2):50–53
- Schvartsman G, Park M, Liu DD, Yennu S, Bruera E, Hui D (2017) Could objective tests be used to measure fatigue in patients with advanced cancer? J Pain Symptom Manag 54(2):237–244. https://doi.org/10.1016/j.jpainsymman.2016.12.343
- Cantarero-Villanueva I, Fernandez-Lao C, Diaz-Rodriguez L, Fernandez-de-Las-Penas C, Ruiz JR, Arroyo-Morales M (2012) The handgrip strength test as a measure of function in breast cancer survivors: relationship to cancer-related symptoms and physical and physiologic parameters. Am J Phys Med Rehabil 91(9):774–782. https://doi.org/10.1097/PHM.0b013e31825f1538
- 32. Thorsen L, Skovlund E, Strømme SB, Hornslien K, Dahl AA, Fosså SD (2005) Effectiveness of physical activity on



- cardiorespiratory fitness and health-related quality of life in young and middle-aged cancer patients shortly after chemotherapy. J Clin Oncol 23(10):2378–2388. https://doi.org/10.1200/jco.2005.04.106
- Chaillou T, McPeek A, Lanner JT (2017) Docetaxel does not impair skeletal muscle force production in a murine model of cancer chemotherapy. Physiol Rep 5(11):e13261. https://doi. org/10.14814/phy2.13261
- Jiralerspong S, Goodwin PJ (2016) Obesity and breast cancer prognosis: evidence, challenges, and opportunities. J Clin Oncol 34(35):4203–4216
- Lemming D, Borsbo B, Sjors A, Lind EB, Arendt-Nielsen L, Graven-Nielsen T, Gerdle B (2015) Single-point but not tonic cuff pressure pain sensitivity is associated with level of physical fitness–a study of non-athletic healthy subjects. PLoS ONE 10(5):e0125432. https://doi.org/10.1371/journal.pone.0125432

- Majithia N, Loprinzi CL, Smith TJ (2016) New practical approaches to chemotherapy-induced neuropathic pain: prevention, assessment, and treatment. Oncology 30(11):1020–1029
- Hilkens PH, Verweij J, Vecht CJ, Stoter G, van den Bent MJ (1997) Clinical characteristics of severe peripheral neuropathy induced by docetaxel (Taxotere). Ann Oncol 8(2):187–190
- Edwards RR, Fillingim RB (2007) Self-reported pain sensitivity:
   Lack of correlation with pain threshold and tolerance. Eur J Pain 11(5):594–598. https://doi.org/10.1016/j.ejpain.2006.09.008
- 39. Baqutayan SMS (2012) The effect of anxiety on breast cancer patients. Indian J Psychol Med 34(2):119–123. https://doi.org/10.4103/0253-7176.101774
- 40. Alfieri FM, Lima ARS, Oliveira NC, Portes LA (2017) The influence of physical fitness on pressure pain threshold of elderly women. J Bodyw Mov Ther 21(3):599–604. https://doi.org/10.1016/j.jbmt.2016.09.008

