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META-ANALYSIS

Physical Activity and Mortality in Cancer Survivors: A Systematic Review and Meta-Analysis

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Abstract

Background: Recommendations for improved survival after cancer through physical activity (PA) exist, although the evidence is still emerging. Our primary objective was to conduct a systematic review and meta-analysis of the association between prediagnosis and postdiagnosis PA and survival (cancer-specific, all-cause, and cardiovascular disease mortality) for all cancers and by tumor site. Secondary objectives were to examine the associations within population subgroups, by PA domain, and to determine the optimal dose of PA related to survival.

Methods: PubMed, EMBASE, and SportsDiscus databases were searched from inception to November 1, 2018. DerSimonian-Laird random-effects models were used to estimate the summary hazard ratios (HRs) and 95% confidence intervals (CI) for primary and secondary analyses and to conduct dose-response analyses.

Results: Evidence from 136 studies showed improved survival outcomes with highest vs lowest levels of prediagnosis or postdiagnosis total or recreational PA for all-cancers combined (cancer specific mortality: HR = 0.82, 95% CI = 0.79 to 0.86, and HR = 0.63, 95% CI = 0.53 to 0.75, respectively) as well as for 11 specific cancer sites. For breast and colorectal cancers, greater reductions were observed for postdiagnosis PA (HR = 0.58-0.63) compared with prediagnosis PA (HR = 0.80-0.86) for cancer-specific and all-cause mortality. Survival benefits through PA were observed in most subgroups (within sex, body mass index, menopausal status, colorectal subtypes, and PA domain) examined. Inverse dose-response relationships between PA and breast cancer-specific and all-cause mortality were observed, with steep reductions in hazards to 10–15 metabolic equivalent hours per week. Conclusion: Higher prediagnosis and postdiagnosis levels of PA were associated with improved survival outcomes for at least 11 cancer types, providing support for global promotion of PA guidelines following cancer.

The role of physical activity (PA) in cancer prevention is well recognized, with recent publications by the World Cancer Research Fund/American Institute for Cancer Research (1) and the 2018 Physical Activity Guidelines for Americans Report highlighting its importance to global health (2). Since the mid-2000s, there has been an exponential increase in studies evaluating the link between PA and survival outcomes that has resulted in some reviews on this topic (3). Although published reviews have explored the relationship between PA and survival (cancer-specific or all-cause mortality) following breast (4–9), colorectal (6,10), or all cancer (11,12), to date there have been no systematic reviews and meta-analyses examining all

available cancer sites (including all-cancer as well as specific cancer sites) with cancer-specific and all-cause mortality outcomes. In addition, cardiovascular disease (CVD) is receiving increasing research attention as a leading cause of mortality for those with cancer. Yet, despite the known benefits through PA on CVD risk and survival, there are no available reviews evaluating cardiovascular mortality following any cancer.

In part, as a consequence of the exponential growth in PA and cancer survival epidemiological research, the momentum behind endorsing and promoting PA in the prevention and management of cancer has also grown (13). Concurrently, however, concerns have been raised about whether there is

sufficient evidence to support the benefits of PA participation for all people with cancer or, alternatively, whether the evidence supports benefit through PA only for specific cancer types or subgroups within cancer types (that is, is dependent on sex, body mass index (BMI), menopausal status, or subtypes within a specific cancer). In addition, the extent to which the evidence can guide recommendations around PA domain (ie, total, recreational [leisure time], occupational, household) and dose of PA and for whom is unclear (14). Hence, there is a need for rigorous review of the rapidly evolving evidence base. As such, the primary objective of this systematic review and meta-analysis was to evaluate the association between prediagnosis and postdiagnosis PA and survival (primary outcomes: cancer-specific mortality, all-cause mortality, and CVD mortality) for all cancer and by specific cancer sites by using data from all available observational epidemiologic studies and randomized, controlled trials. Secondary objectives included assessing these associations by sex, BMI, menopausal status, and colorectal cancer subtype; evaluating the associations between different domains of PA (ie, total, recreational [leisure time], occupational, household) and survival outcomes; and determining the dose-response relationship between PA and cancer survival.

Methods

This systematic review and meta-analysis was performed in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (15). Additionally, the protocol was registered in PROSPERO (registration number: CRD42018103290).

Literature Search Strategy

PubMed, EMBASE, and SportDiscus were searched from inception to July 5, 2018, using the search strategy "(physical activity OR motor activity OR exercise) AND (cancer OR neoplasm* OR carcinoma OR adenocarcinoma OR sarcoma OR tumor) AND (mortality OR recurrence OR progression OR outcome* OR survival) AND (survivors OR survivor OR survivorship OR patients OR patient)." Keywords (including any associated synonyms) along with medical subject headings for PA, cancer, and mortality were included. There were no restrictions by date, language, or geographical region. Reference lists of all included studies and relevant review articles were searched manually to identify additional studies, and e-alert notifications in PubMed captured additional articles through November 1, 2018.

Eligibility Screening

Eligibility was assessed independently and in duplicate using a two-stage process. First, two independent reviewers (CRS and ML, acknowledgments) screened title and abstracts of all captured literature. Studies were considered for full-text review if the title or abstract indicated that the exposure was PA and the outcome was related to survival outcomes following cancer (survival, mortality, recurrence, progression, etc) in human populations. If relevance was uncertain, the study was carried forward for full-text review. Second, two independent reviewers (CRS and either RKP, NM, or RU, acknowledgments) reviewed the remaining studies in their entirety. Inclusion criteria for full-text review were as follows: 1) the original peer-reviewed published research was available; 2) the exposure was PA,

presented with a comparator group (ie, not continuously); 3) one or more mortality outcomes were reported (ie, cancerspecific mortality, all-cause mortality in cancer patients, CVD mortality in cancer patients); 4) the outcomes reported included a point estimate of risk, hazards, or odds ratios; 5) the study design was observational cohort or randomized trial (case reports and reviews were excluded).

Agreement between the two reviewers was quantified at the full-text review stage using percentage agreement and kappa statistics. Disagreements were resolved by consensus-based discussion between reviewers. In the event that there were multiple publications describing the same population with the same domain of PA exposure and mortality outcome, with no new subgroups of interest presented, the article presenting the largest sample size was retained in the review.

Data Extraction

A data collection form, developed specifically for this review, was used to extract and record author, publication year, study name, location, sample size, number of deaths, recruitment years, date of last follow-up, follow-up period, method of PA assessment, and outcome ascertainment source from eligible publications. We additionally extracted the following variables: cancer type, outcome type, timing of PA, domain of PA, high and low activity categories, activity units, hazard estimates and 95% confidence intervals for the highest vs lowest category of PA from the most adjusted model, population subgroups data on sex, BMI (kg/m²), menopausal status, and estimates by colorectal cancer subsite and by domain of PA. We calculated the reciprocal of the reported point estimate if the lowest vs the highest level of PA was presented. When "floating" confidence intervals were reported, we converted them to conventional confidence intervals with a reference category (16). We contacted six authors (regarding eight papers) via e-mail up to two times to request information that was essential for meta-analysis; four authors replied.

Decision rules for data extraction were established to align with our primary aim and ensure consistent extraction of the exposure of interest: physical activity. For example, if multiple estimates were presented for different activity intensities, we extracted, in priority order, the point estimate for all intensities, moderately vigorous, vigorous, moderate, and finally light intensities. If multiple domains of PA were reported, we extracted, in priority order, the point estimate for total, recreational, occupational, and finally, household PA. If multiple estimates were presented for different life-periods prediagnosis, we extracted the estimate closest to diagnosis, rather than lifetime PA, to capture the short-term effects of exercise. Finally, if multiple estimates were provided for different units of activity, we extracted, in hierarchical order, the following: metabolic equivalent duration (MET; one MET is considered to be the resting metabolic rate achieved during quiet sitting [17]), hours per week, energy expenditure (kilocalories or kilojoules), frequency (times per day), and ordinal or rank (ie, scale of 1-10, categories).

Study Quality Assessment

A single reviewer (CRS) used the Newcastle-Ottawa Scale to assess the quality of each included study (18). This scale assesses the quality of included studies with scores ranging from zero (indicating poor-quality studies) to nine (indicating high-quality studies). The scores come from three domains: selection,

comparability, and outcome. The domain of selection was worth a maximum of four points based on sample selection (two points if the sample was representative of the exposed cohort and one point if the sample was composed of a selected group of individuals, ie, nurses, volunteers); ascertainment of exposure (one point if PA was ascertained through interview or actigraphy and zero points if self-administered); and outcome (one point if outcome was not present at start of study). The domain of comparability was worth a maximum of two points, with one point being awarded if models controlled for age, and an additional point awarded if models controlled for additional confounders. Finally, the domain of outcome was worth a maximum of three points based on outcome assessment (one point if outcome was obtained through record linkage), length of follow-up (one point if study had a follow-up time of more than three years), and loss to follow-up (one point if loss to follow-up was described, or if study had complete follow-up).

Statistical Analysis

To account for heterogeneity within the included studies, estimates were combined only if they pertained to the same cancer type, outcome type (cancer-specific, all-cause, or CVDspecific mortality), and timing of PA (prediagnosis or postdiagnosis). To account further for the inherent between-study heterogeneity in the population of patients, we used DerSimonian and Laird random-effects models to derive summary estimates of hazards depicted graphically with forest plots (19). Studies were represented once per meta-analysis except when results were only available for subgroup (ie, by sex). In these instances, each subgroup was treated as an independent study within random-effects models to acknowledge clinical heterogeneity and to reduce within-study confounding. Meta-regression and stratified analyses were performed to ensure that summary estimates did not differ by time-scale (ie, healthy cohorts vs cancer survivor cohorts) (20). Sensitivity analyses were performed, removing each study one by one to examine the impact of combining randomized, controlled trials with observational studies. Subgroup meta-analyses were conducted across strata of cancer type, outcome type, and timing of PA by domain of PA (total, recreational and/or leisure time, transportation, occupational, household), BMI ($<25 \text{ kg/m}^2$, $\ge25 \text{ kg/m}^2$), sex (male, female), menopausal status (premenopausal, postmenopausal; where studies presented results by age, we used a cut point of 55 years whereby younger than 55 years was classified as premenopausal and older than 55 years was classified as postmenopausal; limited to breast cancer), and colorectal cancer subsite (colon, rectum). Where there were sufficient studies presenting estimates based on recreational PA volume in MET hours per week, we performed random-effects dose-response analyses (21). We applied the midpoint of each exposure category or the limit for open-ended exposure categories (eg, 10-20 was assigned a value of 15; <3 was assigned a value of 1.5).

Heterogeneity was assessed using I² statistics, which serve to describe the percentage of variation across studies due to heterogeneity rather than chance; I² values of 25%, 50%, and 75% indicate low, moderate, and high levels of heterogeneity, respectively (22). Publication bias was assessed pertaining to our primary objective with three or more estimates qualitatively through visual inspection of funnel plots and quantitatively using the Begg rank correlation test and Egger regression test for funnel plot asymmetry (23,24). All analyses were conducted using Stata software (version 15.1; StataCorp LP, College Station,

TX); P values less than .05 were considered to be statistically significant and all tests were two-sided.

Results

Literature Search

We identified 15 760 records from our database search, five from PubMed e-alerts, and 31 through other sources such as reference lists, relevant review articles, and literature summary documents maintained by authors (Figure 1). After removing duplicates, 11 996 titles or abstracts remained and 967 were eligible for full-text screening. Full-text screening by two independent reviewers resulted in 97.5% agreement on inclusion or exclusion (kappa = 0.857). A total of 136 studies remained for inclusion in this systematic review and meta-analysis.

Study Characteristics

The study design, sample size, outcomes, and methods for PA assessment for the 136 included studies are shown in Table 1. Of these, nine studies reported on multiple cancer sites, 38 on all-cancer sites combined, 39 on breast cancer, 19 on colorectal cancer, nine on prostate cancer, four each for ovarian and pancreatic cancers, three each on endometrial and hematologic cancers, two for lung cancer, and one each for bladder cancer, cervical, childhood, kidney cancers, malignant glioma, and melanoma. To improve the precision of our estimates, we combined cervical, endometrial, and ovarian cancers as "female reproductive" cancers and leukemia, lymphoma, myeloma, and other hematopoietic cancers as "hematologic" cancers. The included studies were primarily of high quality (scores >7), with 38 studies receiving perfect scores on the Newcastle-Ottawa quality assessment (Table 1). The most common reasons for reductions on the quality assessment scale were the use of selfadministered questionnaires to report PA behaviors (56% of studies used participant-reported or retrospective data collection to ascertain PA levels) and having nonrepresentative population samples (15% of included studies).

Primary Results

Figures 2 and 3 display forest plots of the summary hazard ratios for the highest vs lowest amount of prediagnosis and postdiagnosis PA for all cancers and specific cancer sites on cancer-specific mortality and all-cause mortality, respectively. Evidence from 136 studies contributed to findings showing reduced hazards of mortality for those in the highest vs lowest levels of prediagnosis and/or postdiagnosis total or recreational PA for all cancers combined (cancer-specific mortality: hazard ratio [HR] = 0.82, 95% confidence interval [CI] = 0.79 to 0.86, and $HR\,{=}\,0.63,\,95\%$ CI ${=}\,0.53$ to 0.75, respectively). Statistically significantly reduced hazards were also found for 11 cancer types depending on timing of PA (prediagnosis and postdiagnosis) and mortality outcome (cancer-specific and all-cause mortality). Specifically, higher prediagnosis PA was protective against cancer-specific mortality following breast, colorectal, hematologic, liver, lung, and stomach cancer, and higher postdiagnosis PA was protective against cancer-specific mortality following breast, colorectal, and prostate cancer (Figure 2). For all-cause mortality, higher prediagnosis PA was protective against breast, colorectal, hematologic, and prostate cancer, and higher postdiagnosis PA was protective following breast, childhood,

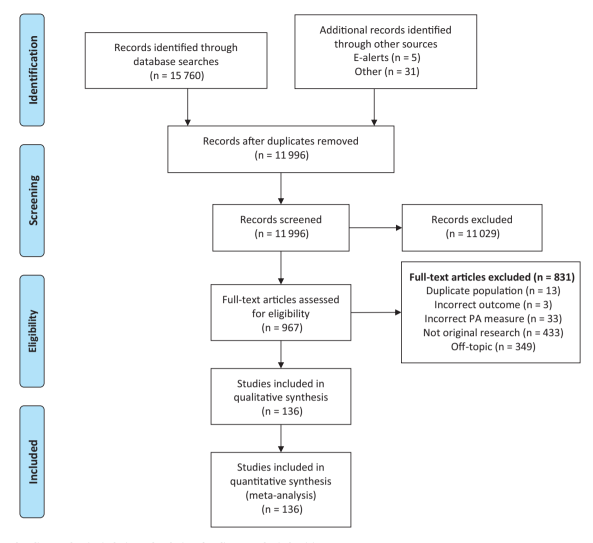


Figure 1. Flow diagram showing inclusion and exclusion of studies. PA = physical activity.

colorectal, gynecologic, glioma, hematologic, kidney, lung, prostate, and stomach cancer (Figure 3). Breast and colorectal cancer sites had the largest number of contributing studies, and results suggest that greater reductions were observed for postdiagnosis PA both for cancer-specific and all-cause mortality (HR = 0.58-0.63) compared with mortality reductions observed with prediagnosis PA (HR = 0.80–0.86). Summary estimates did not differ by time scale (Supplementary Table 1, available online), and thus healthy cohorts and cancer survival cohorts were combined in the results. Further, removal of randomized, controlled trials did not change the results (data not shown).

When considering the association between PA and CVD mortality and given the small number of studies, prediagnosis and postdiagnosis PA were combined to create a single estimate. The summary hazard ratios for all-cancer (n=3), childhood cancer (n = 1), and colorectal cancer (n = 4) were 0.60 (95% CI = 0.50-0.73), 0.89 (95% CI = 0.49-1.61), and 0.60 (95% CI = 0.40-1.61) 0.91), respectively. No cancer sites were found to have statistically significant increased mortality hazards (for any mortality outcome) associated with higher levels of PA (Supplementary Table 2, available online).

After visual examination of funnel plots and P values from the Begg and Egger tests, there was evidence for publication

bias only for postdiagnosis PA and colorectal cancer-specific mortality (P < .05) (results not shown).

Subgroup Analysis Results

Subgroup analyses by sex, BMI, menopausal status (in breast cancer), and colorectal subtype are presented in Table 2. Overall, hazards of cancer-specific and all-cause mortality for those undertaking higher vs lower prediagnosis and/or postdiagnosis PA were reduced both for men and women (all cancers and within colorectal cancer), those with lower BMI (<25 kg/m²; for all cancers, and within breast and colorectal but not within prostate cancer), prediagnosis and postmenopausal women (except for the association for premenopausal women and breast cancer-specific mortality), and colorectal subtypes, with trends toward stronger effect for postdiagnosis PA (HR = 0.37-0.88) vs prediagnosis PA (HR = 0.75-1.53). There was some suggestion (based on differences in effect size observed across colorectal, breast, and hematological cancer groups) that benefit through postdiagnosis PA to all-cause mortality survival was greater for those with BMI less than 25 kg/m² (HR = 0.49-0.57;

Table 1. Characteristics of the included studies in the systematic review and meta-analysis on physical activity and cancer mortality, by cancer site*

Name of study	Cancer type	No. deaths	No. with cancer	No. in analytic sample	Outcome type	Physical activity assessment	Subgroups used	Quality score (of 9)
	All, lung, hematopoietic, stomach, pancreas	832	I	6702	Cancer-specific	Prediagnosis	1	7
	All, lung, colorectal, hematopoietic, stomach, pancreas, colorectal	1499	I	11663	Cancer-specific	Prediagnosis	CRC subsite	7
vedish Oesophagea and Cardia Cancer Studv	l Es	I	280	280	All-cause	Prediagnosis	I	6
	All, colorectal, liver, lung, breast	2185	I	416175	Cancer-specific	Prediagnosis	Age, sex	∞
National Institutes of Health-AARP Diet and Health Study	All, lymphocytic leukemia, liver, oral cavity and pharynx, non-Hodgkin lymphoma, esophagus, myeloma, lung, myeloid, monocytic leukemia, stomach, ovarian, prostate, bladder, breast, brain, endometrial, pan-	15 001	1	293511	Cancer-specific	Prediagnosis	I	∞
BioBank Japan Project	Es	816	1939	1939	All-cause	Postdiagnosis	I	6
orean Metabolic Syndrome Mortality Study	All, esophagus, head and neck, liver, lung, colorectal, pancreas, kidney, stomach, prostate, breast, cervix	7539	I	303428	Cancer-specific	Prediagnosis	No overall, by sex	∞
National Institutes of Health–AARP Diet and Health Study	Hematologic, non-Hodgkin lymphoma, myeloma, leukemia	2606	5182	5182	All-cause, can- cer-specific	Prediagnosis, postdiagnosis	Age, sex, BMI	∞
	All, breast, prostate, colorectal, uterine	3528	13 997	13997	All-cause, can- cer-specific, CVD-specific	Postdiagnosis	1	ത
v. =	Canada Health Survey All Mortality Follow-up Study	229	1	12917	Cancer-specific	Prediagnosis	I	6
erobics Center Longitudinal Study	All	223	I	32421	Cancer-specific	Prediagnosis	No overall, by sex	7
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Author, year, country, reference	Name of study		Cancer type	No. deaths	No. with cancer	No. in analytic sample	Outcome type	Physical activity assessment	Subgroups used	Quality score (of 9)
Rosengren, 1997,	Multifactor Primary	All		723	I	7142	Cancer-specific	Prediagnosis	Men only	∞
Kilander, 2001,	-	All		216	1	2285	Cancer-specific	Prediagnosis	Men only	∞
Hu, 2005, Finland	I	All		2039	I	47 212	Cancer-specific	Prediagnosis	No overall, by sex	∞
(38) Schnohr, 2006,	Copenhagen City	All		632	1	4894	Cancer-specific	Prediagnosis	I	∞
Matthews, 2007,	Heart Study Shanghai Women's	All		537	I	67 143	Cancer-specific	Prediagnosis	Women only	0
Crima (40) Orsini, 2008, Sweden (41)	Health Study Cohort of Swedish Men	All		901	I	37 633	Cancer-specific	Prediagnosis	No overall, by BMI,	∞
van Dam, 2008, United States	Nurses' Health Study	All		4527	I	77 782	Cancer-specific	Prediagnosis	Women only	7
(42) Hamer, 2009,	Scottish Health	All		78	293	293	All-cause	Postdiagnosis	Type of PA	0
Scotland (43) Autenrieth, 2011,	Surveys MONICA/KORA	All		326	I	4672	Cancer-specific	Prediagnosis	Type of PA	0
Germany (44) Borch, 2011,	Augsburg Survey (S2) Norwegian Women	All		1584	I	66136	Cancer-specific	Prediagnosis	Women only	∞
Norway (45)	and Cancer Cohort Study									
Laukkanen, 2011, Finland (46)	Eastern Finnish Follow-up Study	All		181	I	2560	Cancer-specific	Prediagnosis	Men only	6
McCullough, 2011, United States (47)	Cancer Prevention Study-II Nutrition	All		5874	I	111966	Cancer-specific	Prediagnosis	No overall, by sex	∞
Lin, 2012, Taiwan	Taichung Diabetes	All		122	1	2686	Cancer-specific	Prediagnosis	I	∞
(48) Mok, 2012, Korea	Study Severance Cohort	All		1060	3555	29 636	Cancer-specific	Prediagnosis	No overall, by sex	∞
(49) Parekh, 2012, United States	Study Third National Health and Nutrition	All		098	I	15535	Cancer-specific	Prediagnosis	BMI	0
(50) Inoue-Choi, 2013,	Examination Survey Iowa Women's Health	All		461	2017	2017	All-cause, can-	Postdiagnosis	Women only	∞
United States (51) Vergnaud, 2013.	Study European Prospective	T V		9388	I	378864	cer-specific, CVD-specific Cancer-specific	Prediagnosis	Sex	∞
Europe (52)	Investigation into Cancer and Nutrition						•	0		

						No.		Dhvisical		Origlity
Author, year, coun-				No.	No. with	analytic	Outcome	activity		score
try, reference	Name of study		Cancer type	deaths	cancer	sample	type	assessment	Subgroups used	(6 Jo)
Wang, 2013, China (53)	Shanghai Men's Health Studv	All		1053	I	61477	Cancer-specific	Prediagnosis	Men only	6
Yu, 2013, China (54)		All		452	I	2867	Cancer-specific	Prediagnosis	No overall, by sex	∞
Gunnell, 2014, Anstralia (55)	Busselton Health	All		164	528	2320	Cancer-specific	Prediagnosis	I	∞
Hardee, 2014,	Aerobics Center	All		121	2863	2863	All-cause	Postdiagnosis	I	∞
United States (50) Hastert, 2014, Thited States (57)	Longitudinal Study Vitamins and Lifestyle Study	All		1595	I	57841	Cancer-specific	Prediagnosis	I	∞
Lee, 2014, United States (58)	Judy Harvard Alumni Health Study	All		777	1021	1021	All-cause, can- cer-specific,	Postdiagnosis	Men only	7
Wanner, 2014, Switzerland (59)	MONICA/National Research Program	All		1351	I	17 663	Cancer-specific	Prediagnosis	Sex	∞
Brown, 2015, United States	Third National Health and Nutrition	All		319	416	416	All-cause	Prediagnosis	I	∞
(60) Kabat, 2015, United States	Examination Survey National Institutes of Health–AARP Diet	All		16 193	73 784	476396	Cancer-specific	Prediagnosis	No overall, by sex	∞
(61) Kraschnewski, 2016,	and Health Study National Health	All		I	I	30162	Cancer-specific	Prediagnosis	I	7
United States (62) Lee, 2016, Korea	Interview Survey Kangbuk Samsung	All		970	I	336560	Cancer-specific	Prediagnosis	Age, sex, BMI	∞
(62) Robsahm, 2016, Norwaw (64)	Oslo Ischemia Study	All		433	758	1997	Cancer-specific	Prediagnosis	Men only, type of PA	∞
Gunnell, 2017, Australia (65)	Western Australia Health and Wellbeing	All		135	1589	4734	All-cause, can- cer-specific	Postdiagnosis	I	6
Kamada, 2017, United States (66)	Surveniance System Women's Health Study	All		748	I	28879	Cancer-specific	Prediagnosis	Women only	∞
O'Donovan, 2017, United Kingdom (67)	Health Survey for England and Scottish Health	All		2526	I	63591	Cancer-specific	Prediagnosis	I	6
	ourvey								(5)	(continued)

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Author, year, country, reference	Name of study	Cancer type	No. deaths	No. with cancer	No. in analytic sample	Outcome type	Physical activity assessment	Subgroups used	Quality score (of 9)
Vainshelboim, 2017,	Veterans Exercise	All	447	1013	4034	Cancer-specific	Prediagnosis	Men only	7
United states (98) Dohrn, 2018, Sweden (69)	sesung stuay Sweden Attitude Behaviour and	All	27	I	851	Cancer-specific	Prediagnosis	I	6
Liu, 2018, China (70)	Change Study Shanghai Men's Health Study and	All	3512	I	120727	Cancer-specific	Prediagnosis	Sex	Ø
Patel, 2018, United States (71)	Shanghai Women's Health Study Cancer Prevention Study-II Nutrition	All	13 186	I	139 255	Cancer-specific	Prediagnosis	I	∞
Studies assessing only bladder cancer Liss, 2017, United National Hea	bladder cancer National Health Information Survey	Bladder	83	I	222163	Cancer-specific	Prediagnosis	I	ത
Studies assessing only ofeast cancer Rohan, 1995,	01east cancei 	Breast	112	412	412	Cancer-specific	Prediagnosis	Menopausal status	6
Borugian, 2004, Canada (74)	I	Breast	112	603	603	Cancer-specific	Prediagnosis	Menopausal status	∞
Enger, 2004,	I	Breast	251	717	717	Cancer-specific	Prediagnosis	I	6
Officed States (73) Holmes, 2005, Thirted States (76)	Nurses' Health Study	Breast	463	2987	2987	All-cause, can-	Postdiagnosis	BMI, menopausal	7
Abrahamson, 2006,	ı	Breast	290	1264	1264	All-cause	Prediagnosis	BMI	∞
Dal Maso, 2008, Italy (78)	I	Breast	503	1453	1453	All-cause, can-	Prediagnosis	Type of PA	6
Holick, 2008, United States (79)	Collaborative Women's Longevity	Breast	412	4482	4482	All-cause, can- cer-specific	Postdiagnosis	BMI, age	∞
Irwin, 2008, United States (80)	Health, Eating, Activity, and Lifestyle Study	Breast	164	933	933	All-cause, can- cer-specific	Prediagnosis, postdiagnosis	BMI, menopausal status	ത
Friedenreich, 2009, Canada (81)		Breast	341	1225	1225	All-cause, can- cer-specific	Prediagnosis	Type of PA	6
Sternfeld, 2009, United States (82)	Life After Cancer Epidemiology Study	Breast	187	1970	1970	All-cause, can- cer-specific	Postdiagnosis	BMI	∞
West-Wright, 2009, United States (83)	California Teachers Study	Breast	460	3539	3539	All-cause, can- cer-specific	Prediagnosis	BMI	7
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Author, year, coun-			No.	No. with	analytic	Outcome	activity		score
try, reference	Name of study	Cancer type	deaths	cancer	sample	type	assessment	Subgroups used	(6 Jo)
Emaus, 2010,	Norwegian Counties	Breast	429	1364	1364	All-cause, can-	Prediagnosis	BMI, menopausal	∞
Norway (84)	Study					cer-specific		status	
Hellmann, 2010,	Copenhagen City	Breast	323	528	528	All-cause, can-	Prediagnosis	Menopausal status	∞
Denmark (85)	Heart Study					cer-specific			
Keegan, 2010,	Breast Cancer Family	Breast	725	4153	4153	All-cause	Prediagnosis	BMI	œ
United States,	Registry								
Canada, Australia									
(86)			7	0	0	-			Ó
Bertram, 2011,	Women's Healthy	Breast	163	2361	2361	All-cause	Postdiagnosis	l	∞
United States	Eating and Living								
(87)	Study								
Chen, 2011, China	Shanghai Breast	Breast	436	4826	4826	All-cause	Postdiagnosis	BMI, menopausal	6
(88)	Cancer Survival							status	
	Study								
Irwin, 2011, United	Women's Health	Breast	350	4643	4643	All-cause, can-	Prediagnosis,	BMI	∞
States (89)	Initiative					cer-specific	postdiagnosis		
Beasley, 2012,	After Breast Cancer	Breast	1468	13 302	13302	All-cause, can-	Postdiagnosis	BMI, menopausal	7
United States,	Pooling Project:					cer-specific		status	
China (9)	LACE-NHS-SBCSS-					•			
	WHEL								
Cleveland, 2012,	Long Island Breast	Breast	196	1508	1508	All-canse, can-	Prediagnosis	BMI, menopausal	6
United States	Cancer Study					cer-specific)	status	
(06)	Project					•			
Schmidt, 2013,	MARIE Study	Breast	367	3393	3393	All-cause, can-	Prediagnosis	BMI	0
Germany (91)	•					cer-specific)		
Tao, 2013, United	Western New York	Breast	170	1170	1170	All-cause, can-	Prediagnosis	1	6
States (92)	Exposure and Breast					cer-specific)		
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Williams, 2013,	National Kunners and	breast	111	I	/9124	Cancer-specinc	Frediagnosis	I	`
Oillea States	Waikels nealul								
(93)	survey	ę.	0	4	,			Š	Ó
Bradsnaw, 2014,	Long Island Breast	breast	4.20	1423	1423	All-cause, can-	Postdiagnosis	BMI	ת
United States (94)	Cancer Study					cer-specific			
Courneya, 2014,	Supervised Trial of	Breast	24	242	242	All-cause	Postdiagnosis	I	6
Canada (95)	Aerobic vs								
	Resistance Training								
de Glas, 2014,	Tamoxifen	Breast	80	521	521	All-cause, can-	Prediagnosis	Age	9
Netherlands (96)	Exemestane					cer-specific			
	Adjuvant								
	Multicenter								
	Lifestyle Study								
								5)	(continued)

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Autnor, year, country, reference	Name of study	ŭ	Cancer type	no. deaths	No. With cancer	analyuc sample	Outcome type	activity assessment	Subgroups used	score (of 9)
Keegan, 2014,	Neighborhoods and	Breast		915	4345	4345	All-cause, can-	Prediagnosis	I	6
United States (97)	Breast Cancer Study						cer-specific			
Williams, 2014, United States (98)	National Runners' and Walkers' Health	Breast		46	986	986	Cancer-specific	Postdiagnosis	I	_
	Survey									
Bao, 2015, China	Shanghai Breast	Breast		128	518	518	All-cause	Postdiagnosis	Triple negative only	6
(66)	Cancer Survival Study									
Borch, 2015,	Norwegian Women	Breast		197	1327	1327	All-cause, can-	Prediagnosis,	BMI, menopausal	∞
Norway (100)	and Cancer Cohort						cer-specific	postdiagnosis	status	
1,1, 2015 IInited	Study California Breast	Broset		1347	4608	4608	All-carres	Drediomocie	RMI menonenisel	σ
States (101)	Cancer Survivorship	Dieast		(FCT	500	900	cer-specific	i rediagilosis	status)
	Consortium									
Pinkston, 2015,	New Mexico Women's	Breast		240	540	1283	All-cause, can-	Prediagnosis	No overall, by race	6
United States (102)	Health Study						cer-specinc		and type or PA	
Ammitzholl 2016	Diet Cancer and	Breact		144	959	959	All-cause	Postdiagnosis	Type of PA	7
Denmark (103)	Health Study			1)			20019		
Jones, 2016, United	Life After Cancer	Breast		405	6211	6211	Cancer-specific	Postdiagnosis	I	∞
States (104)	Epidemiology and									
	Pathways studies									
McCullough, 2017,	Long Island Breast	Breast		486	1254	1254	All-cause, can-	Prediagnosis	I	7
United States (105)	Cancer Study						cer-specific			
Cifu, 2018, United	National Institutes of	Breast		1162	7088	7088	Cancer-specific	Prediagnosis	I	∞
States (106)	Health-AARP Diet									
JO200 2010	and Health Study	Droot		90	722	722	Δ11 σ211σ2	Doct-diognosis	DMI mononing	7
nayes, 2018, Australia (107)	Exercise for nearth Trials	breast		97	727	23/	All-Cause	Fostalagnosis	bivii, menopausai status	
Maliniak, 2018,	Cancer Prevention	Breast		1771	5254	5254	All-cause, can-	Prediagnosis,	No overall, by age	∞
United States	Study-II Nutrition						cer-specific	postdiagnosis		
(108)	Cohort									
Palesh, 2018,	I	Breast		93	103	103	All-cause	Postdiagnosis	I	∞
United States										
(109)										
Parada, 2019,	Carolina Breast Cancer	Breast		717	1808	1808	All-cause	Prediagnosis	I	6
United States	Study									
(110)	برميديم أممنيهمم									
Kim 2016	cervicai caiicei —	Cervical		30	860	860	All-cause	Postdiagnosis	I	00
Korea (111)				3				000000000000000000000000000000000000000)
Studies assessing only childhood cancer	childhood cancer									
Scott, 2018, United	Childhood Cancer	Childhood cancer	l cancer	1063	15 450	15450	All-cause, CVD-	Postdiagnosis	I	∞
States Canada	Survivor Study						specific			
(112)										(Forest way

(2000)									
Author, year, country, reference	Name of study	Cancer type	No. deaths	No. with cancer	No. in analytic sample	Outcome type	Physical activity assessment	Subgroups used	Quality score (of 9)
Studies assessing only colorectal cancer Meyerhardt, 2006, Nurses' Health United States	r colorectal cancer Nurses' Health Study	Colorectal	132	573	573	All-cause, can- cer-specific	Prediagnosis; postdiagnosis	Women only, age, BMI, colon vs	7
(113) Huxley, 2007, China, Hong Kong, Japan, Korea, Singapore, Taiwan, Thailand, Australia, New	Asia Pacific Cohort Studies Collaboration	Colorectal	751	I	539 201	Cancer-specific	Prediagnosis	rectum	φ
Meyerhardt, 2009, United States	Health Professionals Follow-Up Study	Colorectal	258	899	899	All-cause, can- cer-specific	Postdiagnosis	Men only	7
Baade, 2011,	I	Colorectal	462	1825	1825	All-cause, CVD-	Postdiagnosis	Age, sex, BMI, colon	∞
Morrison, 2011, United Kingdom	Whitehall Study	Colorectal	450	I	17 949	Specific Cancer-specific	Prediagnosis	No overall, by colon vs rectum	7
Kuiper, 2012, United States	Women's Health Initiative	Colorectal	265	1339	1339	All-cause, can- cer-specific	Prediagnosis, postdiagnosis	Women only	∞
Boyle, 2013, Australia (119)	Western Australia Bowel Health Studv	Colorectal	224	879	879	All-cause, can- cer-specific	Prediagnosis	Sex, colon vs rectum	∞
Campbell, 2013, United States (120)	Cancer Prevention Study-II Nutrition Cohort	Colorectal	846	2293	2293	All-cause, can- cer-specific, CVD-specific	Prediagnosis, postdiagnosis	Age, sex, BMI, colon vs rectum	∞
Pelser, 2014, United States (121)	National Institutes of Health-AARP Diet and Health Study	Colorectal	1727	5727	5727	All-cause, can- cer-specific, CVD-specific	Prediagnosis,	No overall, by colon vs rectum	∞
Arem, 2015, United States (122)	National Institutes of Health-AARP Diet and Health Study	Colorectal	1541	3797	3797	All-cause, can- cer-specific, CVD-specific	Prediagnosis, postdiagnosis	I	∞
Hardikar, 2015, United States (123)	Colon Cancer Family Registry-Seattle	Colorectal	408	1309	1309	All-cause, can- cer-specific	Prediagnosis	Colon vs rectum	∞
Romaguera, 2015, Europe (124)	European Prospective Investigation into Cancer and Nutrition	Colorectal	1113	3292	3292	All-cause, can- cer-specific	Prediagnosis	I	∞
Mok, 2016, Korea (125)	Korean Metabolic Syndrome Mortality Study	Colorectal	469	ı	226089	Cancer-specific	Prediagnosis	No overall, by sex, colon vs rectum, age, BMI	∞
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Author, year, country, reference	Name of study	Cancer type	No. deaths	No. with cancer	No. in analytic sample	Outcome type	Physical activity assessment	Pesn sdnogguog	Quality score (of 9)
Thong, 2016, Netherlands (126)	Patient-Reported Outcomes Following Initial Treatment and Long-Term Evaluation of Survivorship	Colorectal	249	1552	1552	All-cause	Postdiagnosis	I	σ
Ratjen, 2017, Germany (127)	- Carago	Colorectal	200	1376	1376	All-cause	Postdiagnosis	Type of PA, sex, age, BMI, colon vs	∞
Walter, 2017, Germany (128)	DACHS Study	Colorectal	898	3121	3121	All-cause, can- cer-specific	Prediagnosis	Age, sex, colon vs rectum, BMI	0
Jayasekara, 2018, Australia (129)	Melbourne Collaborative Cohort Study	Colorectal	339	724	724	All-cause, can- cer-specific	Prediagnosis	Colon vs rectum	Q
Phipps, 2018, United States	North Central Cancer Treatment Group	Colorectal	205	1992	1992	All-cause	Prediagnosis	Colon only	7
van Blangan, 2018, United States	Cancer and Leukemia Group B 89803	Colorectal	299	992	992	All-cause	Postdiagnosis	Colon only	∞
(121) Studies assessing only endometrial cancer Arem, 2013, United National Institutes States (132) Health-AARP Di	endometrial cancer National Institutes of Health-AARP Diet and Health Studv	Endometrial	312	1400	1400	All-cause	Prediagnosis	I	∞
Arem, 2013, United States (133)	Women's Health Initiative	Endometrial	163	983	983	All-cause, can- cer-specific	Prediagnosis	I	∞
Arem, 2016, United States (134)	National Institutes of Health–AARP Diet and Health Study	Endometrial	91	280	580	All-cause	Postdiagnosis	I	∞
Studies assessing only malignant glioma Ruden, 2011, United States (135)	malignant glioma —	Malignant glioma	149	243	243	All-cause	Postdiagnosis	I	7
Studies assessing only hematologic cancers Wiskemann, 2015, — Germany (136)	nematologic cancers —	Leukemia	44	103	103	All-cause	Postdiagnosis	I	9
Boyle, 2017, Canada (137)	I	Lymphoma	169	413	413	All-cause, can- cer-specific	Prediagnosis	I	9
Pophali, 2018, United States (138)	Lymphoma SPORE Molecular Epidemiology Resource	Lymphoma	863	3060	3060	All-cause, can-	Prediagnosis	I	∞

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Name of study Cancer type deaths cancer sample type assessment National Institutes of National National Institutes of National Institutes of National Institutes of All Institutes of National Institutes of N	Author vear coun-			Ŋ	No with	No. in	Outcome	Physical activity		Quality
Note Kidney 175 667 667 All-cause, can Postdiagnosis	try, reference	Name of study	Cancer type	deaths	cancer	sample	type	assessment	Subgroups used	(6 Jo)
Study Lung 77 118 118 All-cause Postdiagnosis Lung 512 1466 1466 All-cause Postdiagnosis Lung 512 1466 1466 All-cause Postdiagnosis Lung 512 1466 1466 All-cause Prediagnosis Lung 6arch 67 67 Carce-specific Prediagnosis na Ovarian 346 635 635 Cancer-specific Prediagnosis norer Ovarian 346 600 600 All-cause, can-crapecific Prediagnosis carc-specific 368 264 264 All-cause, can-crapecific Prediagnosis carc-specific 368 264 264 All-cause, can-crapecific Prediagnosis carc-specific 30 264 264 All-cause, can-crapecific Prediagnosis carc-specific 402 - 32687 Cancer-specific Prediagnosis aborative Pancreatic 4	Studies assessing only Schmid, 2018,	kidney cancer National Institutes of	Kidney	175	299	299	All-cause, can-	Postdiagnosis	I	∞
gry and card and another and another an	United States (139)	Health-AARP Diet and Health Study					cer-specific			
gy and Lung Lung 512 1466 All-cause All-cause Postdiagnosis Pung earch All-cause All-cause Prediagnosis Pung earch All-cause All-cause Prediagnosis Pung earch All-cause Prediagnosis Pung man 396 635 Cancer-specific Prediagnosis Puncer All-cause Prediagnosis Prediagnosis Puncer All-cause Prediagnosis Pancreatic 80 264 264 All-cause Prediagnosis Pancreatic 212 - 32687 Cancer-specific Prediagnosis Aby Pancreatic 402 - 100932 Cancer-specific Prediagnosis Bancreatic 1710 - 1290000 Cancer-specific Prediagnosis Bancreatic 1710 - 1290000 Cancer-specific Prediagnosis Bancreatic 1710 - 1290000 Cancer-specific Prediagnosis Bancreatic	Studies assessing only Jones 2012 United	lung cancer —	Ling	77	118	118	All-cause	Postdiagnosis	I	7
gar and Lung Lung and earth and a sarch and a s	States (140)		0			9				
sgy and Lung 1 Lung All-cause, can-can-car-specific Prediagnosis nment Melanoma 341 2465 2465 All-cause, can-cer-specific Prediagnosis nment Ovarian 396 635 635 Cancer-specific Prediagnosis ncer Ovarian 346 600 600 All-cause, can-car-specific Prediagnosis sgy Study cer-specific Prediagnosis Prediagnosis sgy Study cer-specific Prediagnosis spy Study cer-specific Prediagnosis spy Study cer-specific Prediagnosis spy Study cer-specific Prediagnosis spy Study cer-specific Prediagnosis shy	Sloan, 2016,	Mayo Clinic	Lung	512	1466	1466	All-cause	Postdiagnosis	I	7
nument Melanoma 341 2465 2465 All-cause, can-cer-specific Prediagnosis nument Ovarian 396 635 635 Cancer-specific Prediagnosis a Ovarian 346 600 600 All-cause, can-cer-specific Prediagnosis alth Ovarian 346 600 600 All-cause, can-cer-specific Prediagnosis ican Ovarian 80 264 264 All-cause positic Prediagnosis cer er 1100 32 Cancer-specific Prediagnosis daborative Pancreatic 402 - 100 32 Cancer-specific Prediagnosis dy - 1290 000 Cancer-specific Prediagnosis ady - 1290 000 Cancer-specific Prediagnosis	United States	Epidemiology and								
nment Melanoma 341 2465 2465 All-cause, can- prediagnosis a Ovarian 396 635 635 Cancer-specific Prediagnosis alth Ovarian 346 600 600 All-cause, can- lith Ovarian 80 264 264 All-cause, can- prediagnosis car-specific Prediagnosis car-specific Prediagnosis car-specific Prediagnosis alth Ovarian 346 600 600 All-cause, can- postdiagnosis car-specific Prediagnosis aborative Pancreatic 402 — 32.687 Cancer-specific Prediagnosis aborative Pancreatic 402 — 100932 Cancer-specific Prediagnosis aborative Pancreatic 702 — 1290000 Cancer-specific Prediagnosis aborative Pancreatic 702 — 1290000 Cancer-specific Prediagnosis aborative Pancreatic 702 — 1290000 Cancer-specific Prediagnosis	(111)	Cancer Research								
nment Melanoma 341 2465 All-cause, can-cer-specific Prediagnosis ncer Ovarian 396 635 635 Cancer-specific Prediagnosis ncer Ovarian 346 600 600 All-cause Prediagnosis sith Ovarian 346 600 600 All-cause, can-predific Prediagnosis ican Ovarian 346 600 600 All-cause Prediagnosis gy Study cer 264 264 All-cause Prediagnosis er ni Health Pancreatic 222 All-cause Prediagnosis aborative Pancreatic 402 - 100 932 Cancer-specific Prediagnosis ady Pancreatic 1710 - 1290 000 Cancer-specific Prediagnosis dy - 1290 000 Cancer-specific Prediagnosis dy - 1290 000 Cancer-specific Prediagnosis dy - 129		Program								
nument Melanoma 341 2465 All-cause, can-repecific Prediagnosis ovarian 396 635 635 Cancer-specific Prediagnosis ncer Ovarian 346 600 600 All-cause Prediagnosis sith Ovarian 346 600 600 All-cause Prediagnosis sigs Study ser 264 264 All-cause Prediagnosis ser ni Health Pancreatic 212 - 32 687 Cancer-specific Prediagnosis aborative Pancreatic 402 - 100932 Cancer-specific Prediagnosis dy Pancreatic 1710 - 1290 000 Cancer-specific Prediagnosis dy - 1290 000 Cancer-specific Prediagnosis dy - 1290 000 Cancer-specific Prediagnosis	Studies assessing only	' melanoma								
nna Ovarian Ovarian	Schwitzer, 2017,	Genes, Environment	Melanoma	341	2465	2465	All-cause, can-	Prediagnosis	I	6
ncer 396 635 635 Cancer-specific Prediagnosis ncer Ovarian 346 600 600 All-cause, can-repecific Prediagnosis alth Ovarian 346 600 600 All-cause, can-repecific Prediagnosis egy Study 264 264 All-cause pecific Prediagnosis cer 10093 Cancer-specific Prediagnosis aborative Pancreatic 222 23.687 Cancer-specific Prediagnosis dy 1710 1290 000 Cancer-specific Prediagnosis ady Pancreatic 1710 1290 000 Cancer-specific Prediagnosis	Australia,	and Melanoma					cer-specific			
ncer Ovarian 396 635 635 635 Cancer-specific Prediagnosis alth Ovarian 346 600 600 All-cause Prediagnosis ican Ovarian 80 264 264 All-cause Prediagnosis cer-specific Prediagnosis day - 100932 Cancer-specific Prediagnosis day Pancreatic - 1290000 Cancer-specific Prediagnosis day Pancreatic - 1710 - 1290000 Cancer-specific Prediagnosis	Canada, Italy,	Study								
alth Ovarian 396 635 635 Cancer-specific Prediagnosis ncer Ovarian 238 638 1321 All-cause Prediagnosis alth Ovarian 346 600 600 All-cause, can-cer-specific Prediagnosis ican Ovarian 80 264 264 All-cause Prediagnosis gy Study 212 - 32 687 Cancer-specific Prediagnosis aborative Pancreatic 402 - 100 932 Cancer-specific Prediagnosis aly Pancreatic 1710 - 1290 000 Cancer-specific Prediagnosis udy Pancreatic 52 - 30 826 Cancer-specific Prediagnosis	United States									
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North Carolina Ovarian 396 635 638 Cancer-specific Prediagnosis Ovarian Cancer Study Women's Health Ovarian Ovarian 346 600 600 All-cause, can- Initiative Initiative A African American Ovarian Ovarian 80 264 264 All-cause Prediagnosis, Epidemiology Study Inpancreatic cancer College Alumni Health Pancreatic Cancer College Alumni Health Pancreatic A02 - 100932 Cancer-specific Prediagnosis Study Inpancreatic Cancer College Alumni Health Pancreatic Cancer College Alumni Health Pancreatic Cancer College Alumni Health Pancreatic Cancer Study Inpancreatic Cancer College Alumni Health Pancreatic Cancer Study Inpancreatic Cancer College Alumni Health Pancreatic Cancer Study Inpancreatic Cancer Cancer College Alumni Health Pancreatic Cancer Specific Prediagnosis Cancer-specific Prediagnosis Inpancreatic Cancer Cancer Cancer Specific Prediagnosis Cancer Specific Prediagnosis Inpancreatic Cancer Cancer Cancer Specific Prediagnosis Inpancreatic Cancer C	Studies assessing only	ovarian cancer					!			
North Carolina Ovarian Ovarian 238 638 1321 All-cause Prediagnosis Study Women's Health Ovarian American African American Ovarian Cancer Epidemiology Study Japanese Collaborative Study Japanese Collaborative Study Million Women Study Million Women Study Takayama Study Takayama Study Ovarian Ovarian Ovarian 346 600 600 All-cause, can- Sudy Japanese Collaborative Pancreatic Cohort Study Million Women Study Takayama Study T	Yang, 2008,	I	Ovarian	396	635	635	Cancer-specific	Prediagnosis	I	∞
North Carolina Ovarian 238 638 1321 All-cause Prediagnosis Ovarian Cancer Study Women's Health Ovarian 346 600 600 All-cause, can-prediagnosis cancer Cancer Cancer College Alumni Health Pancreatic Cancer Specific Prediagnosis Cohort Study Alumni Million Women Study Pancreatic Cancer Specific Prediagnosis Cancer Specific Prediagnosis Takayama Study Pancreatic Cancer Specific Prediagnosis Cancer Specific Prediagnosis	Sweden (143)									
Study Women's Health Ovarian Momen's Health Ovarian Momen's Health Ovarian Momen's Health Ovarian African American Cancer Cancer Cancer Cancer Callege Alumni Health Pancreatic Colort Study Million Women Study Mallon Women Study Pancreatic Takayama Study Pancreatic Ovarian 346 600 600 All-cause, can- cer-specific Active Active Active All-cause All-cause Prediagnosis Cancer-specific Prediagnosis Prediagnosis Cancer-specific Prediagnosis Prediagnosis Cancer-specific Prediagnosis Prediagnosis	Moorman, 2011,	North Carolina	Ovarian	238	638	1321	All-cause	Prediagnosis	Ι	6
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Women's Health Ovarian 346 600 600 All-cause, can-repedition Prediagnosis d African American Cancer Care-specific Prediagnosis Cancer Cancer Prediagnosis Epidemiology Study Pancreatic 212 — 32 687 Cancer-specific Prediagnosis College Alumni Health Pancreatic 402 — 100 932 Cancer-specific Prediagnosis Study Japanese Collaborative Pancreatic 402 — 1290 000 Cancer-specific Prediagnosis Million Women Study Pancreatic Takayama Study Pancreatic Prediagnosis Prediagnosis Takayama Study Pancreatic 52 — 30 826 Cancer-specific Prediagnosis	(144)	Study								
n Ovarian 80 264 264 All-cause Prediagnosis, postdiagnosis, postdiagnosis Study Health Pancreatic 712 — 32.687 Cancer-specific Prediagnosis Orative Pancreatic 402 — 100.932 Cancer-specific Prediagnosis Study Pancreatic 1710 — 1290.000 Cancer-specific Prediagnosis I Pancreatic 52 — 30.826 Cancer-specific Prediagnosis	Zhou, 2014, United	Women's Health	Ovarian	346	009	009	All-cause, can-	Prediagnosis	I	∞
Study Health Pancreatic Pancreatic Study Health Pancreatic Prediagnosis Prediagnosis Prediagnosis Prediagnosis Prediagnosis	States (145)	Initiative	-	Ċ		Č	cer-specific			Ċ
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Health Pancreatic 212 — 32.687 Cancer-specific Prediagnosis orative Pancreatic 402 — 100.932 Cancer-specific Prediagnosis Study Pancreatic 1710 — 1290.000 Cancer-specific Prediagnosis 7 Pancreatic 52 — 30.826 Cancer-specific Prediagnosis	States (146)	Cancer						postdiagnosis		
Health Pancreatic 212 — 32 687 Cancer-specific Prediagnosis orative Pancreatic 402 — 100 932 Cancer-specific Prediagnosis Study Pancreatic 1710 — 1290 000 Cancer-specific Prediagnosis I Pancreatic 52 — 30 826 Cancer-specific Prediagnosis		Epidemiology Study								
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Japanese Collaborative Pancreauc 402 — 100 932 Cancer-specinc Prediagnosis Cohort Study Pancreatic 1710 — 1290 000 Cancer-specific Prediagnosis Takayama Study Pancreatic 52 — 30 826 Cancer-specific Prediagnosis	States (147)	study	:			0			;	ć
Conort Study Million Women Study Pancreatic 1710 — 1290 000 Cancer-specific Prediagnosis Takayama Study Pancreatic 52 — 30826 Cancer-specific Prediagnosis	Lin, 2007, Japan	Japanese Collaborative	Fancreatic	402		100932	Cancer-specific	Prediagnosis	No overall, by sex	×
Million Women Study Pancreauc 1/10 — 1290 000 Cancer-specinc Prediagnosis m Takayama Study Pancreatic 52 — 30 826 Cancer-specific Prediagnosis	(148)	Conort study		7		0				Ċ
m Takayama Study Pancreatic 52 — 30826 Cancer-specific Prediagnosis	Stevens, 2009,	Million Women Study	Fancreatic	1/10		1.290.000	Cancer-specific	Prediagnosis	Women only	×
Takayama Study Pancreatic 52 — 30826 Cancer-specific Prediagnosis	United Kingdom (149)									
Japan (150)	Nakamura, 2011,	Takayama Study	Pancreatic	52	I	30826	Cancer-specific	Prediagnosis	No overall, by sex	∞
	Japan (150)									Ī

(continued)

Author, year, country, reference	Name of study	Gancer type	No. deaths	No. with cancer	No. in analytic sample	Outcome type	Physical activity assessment	Subgroups used	Quality score (of 9)
Studies assessing only prostate cancer Nilsen, 2006, HUNT Study (r prostate cancer HUNT Study (Norway)	Prostate	354	957	29110	Cancer-specific	Prediagnosis	I	∞
Norway (151) Crespo, 2008 (152)	Puerto Rico Heart Health Program	Prostate	167	I	9780	Cancer-specific	Prediagnosis	BMI, age	6
Puerto Rico Orsini, 2009,	Cohort of Swedish	Prostate	190	I	45887	Cancer-specific	Prediagnosis	Type of PA	∞
Batty, 2011, United	Mhitehall Study	Prostate	578	I	17934	Cancer-specific	Prediagnosis	Type of PA	7
Kenfield, 2011, United States	Health Professionals Follow-Up Study	Prostate	548	2705	2705	All-cause, can- cer-specific	Postdiagnosis	I	7
(155) Bonn, 2015, Sweden (156)	Progression in Cancer	Prostate	561	4623	4623	All-cause, can-	Postdiagnosis	Type of PA	∞
Friedenreich, 2016,		Prostate	458	830	830	All-cause, can-	Prediagnosis,	Type of PA	6
Tai, 2016, Taiwan (158)	I	Prostate	48	809	809	Cancer-specific	Prediagnosis	I	7
Wang, 2017, United States (159)	Cancer Prevention Study-II Nutrition Cohort	Prostate	454	7328	7328	All-cause, can- cer-specific	Prediagnosis, postdiagnosis	ſ	∞

 $^*BMI = body\ mass\ index;\ CRC = colorectal\ cancer,\ CVD = cardiovas cular\ disease,\ PA = physical\ activity.$

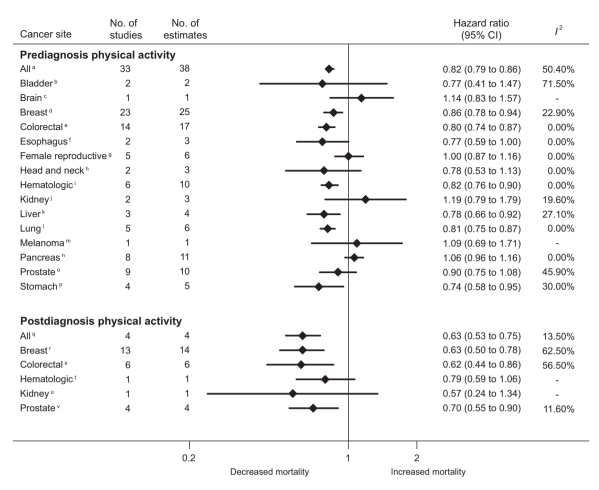


Figure 2. Summary hazard ratios for the highest vs lowest levels of prediagnosis and postdiagnosis physical activity and cancer-specific mortality by cancer site (each estimate denotes a separate meta-analysis performed; if only one estimate is present, then no meta-analyses were conducted and the individual point estimate is reported). aRefs. (25,26,28,29,31,34-39,41,42,44-46,48-50,52,54,55,57,59,62-64,66-71). Refs. (29,72). aRefs. (29,72). dRefs. (28,31,73-75,78,80,81,83-85,89,91-93,96,97,100-100). 102,105,106,108). eRefs. (26,28,31,113,114,117-120,122-124,128,129). eRefs. (29,31). eRefs. (29,31,132,143,145). eRefs. (29,31). eRefs. (29,31 (28,29,31). ¹Refs. (25,26,28,29,31). ^mRefs. (142). ^mRefs. (25,26,29,31,147-150). ^oRefs. (29,31,151-154,157-159). ^pRefs. (25,26,29,31). ^qRefs. (33,51,55,58). ^rRefs. (9,76,79,80,82,88,89,93,94,96,100,104,108). *Refs. (113,115,116,118,120,122). 'Refs. (32). "Refs. (139). 'Refs. (155–157,159). CI = confidence interval.

all P < .05) compared with those with BMI greater than 25 kg/m² (HR = 0.64-0.71; P < .05-0.112).

PA Domain Results

Additional subgroup analyses by domain of PA (total, recreational, transportation, occupational, and household) are presented in Table 3. For prediagnosis PA, the domains of recreational and total PA estimates were consistently associated with reduced hazards of mortality for all-cancer, breast, and colorectal cancer-specific mortality (P < .05). Results remained inconsistent for the less-studied domains of transportation, occupational, and household PA (HR = 0.64-1.65).

Dose-Response Analyses

We restricted the analysis of dose-response to breast cancer studies because few studies examined these associations for other cancer sites. There was a linear association between prediagnosis PA dose and all-cause mortality (P for nonlinearity = .53) (Figure 4C). Evidence of nonlinear associations was found (P for nonlinearity <.05) between prediagnosis and postdiagnosis

PA and breast cancer-specific mortality (Figure 4, A and B, respectively) and postdiagnosis PA and all-cause mortality (Figure 4D). As seen in Figure 4B, the dose-response curve for postdiagnosis PA and all-cause mortality shows the largest reductions in mortality. Compared with no recreational PA, 5, 10, 20, 30, and 65 MET hours per week reduced all-cause mortality by 22%, 43%, 59%, 69%, and 108%, respectively. The steep reductions in mortality seen in Figure 4, A, B, and D, become less pronounced when PA dose is 10-15 MET hours per week or greater. The upper bounds of Figure 4C are less precise because of few contributing studies at higher levels of PA.

Discussion

In this first ever analysis, to our knowledge, of the association between PA and cancer survival that included all cancer sites, we found evidence from 136 studies conducted to date for improved survival outcomes for all cancer and 11 cancer sites associated with prediagnosis or postcancer diagnosis PA. Although the most consistent and strong evidence for a role of PA in cancer survival was found for breast and colorectal cancer, there is also clear evidence for improved prostate cancer-specific survival with

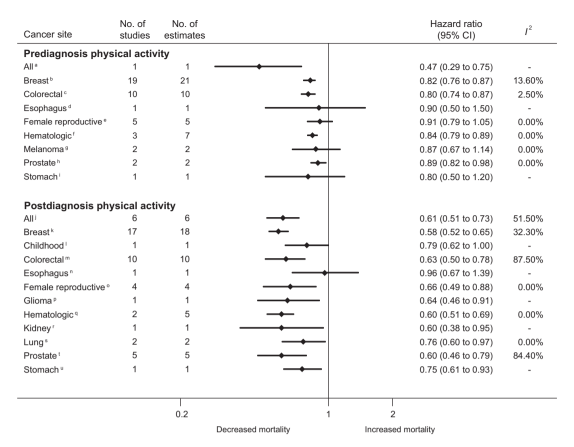


Figure 3. Summary hazard ratios for the highest vs lowest levels of prediagnosis and postdiagnosis physical activity and all-cause mortality in cancer survivors by cancer site (each estimate denotes a separate meta-analysis performed; if only one estimate is present, then no meta-analyses were conducted and the individual point estimate is reported). ^aRefs. (60). ^bRefs. (77,78,80,81,83-86,89,91,92,96,97,100-102,105,108,110). ^cRefs. (113,118-120,122-124,128-130). ^dRefs. (27). ^eRefs. (132,144-146). ^fRefs. (32,137,138). ^gRefs. (27,142). ^hRefs. (157,159). ⁱRefs. (27). ^jRefs. (33,43,51,55,56,58). ^kRefs. (9,33,76,79,80,82,87–89,94–96,100,103,107–109). ⁱRefs. (112). ^mRefs. (33,113,115,116,118,120,122,126,127,131). Refs. (30). Refs. (33,111,134,146). PRefs. (135). Refs. (136,139). Refs. (139). Refs. (140,141). Refs. (33,155-157,159). Refs. (30). CI = confidence interval.

postdiagnosis PA. In addition, there is emerging evidence for a beneficial effect of prediagnosis PA on cancer-specific survival for liver, lung, hematologic, esophageal, and stomach cancers. Compared with prediagnosis PA, postdiagnosis PA was associated with greater reductions both in cancer-specific and allcause mortality, with greater than 30% reductions in hazards for all-cause mortality observed in studies of all cancer, breast, colorectal, female reproductive, glioma, kidney, lung, prostate, and stomach cancers (HR = 0.58-0.76).

This study extends the results found in previous metaanalyses of PA and cancer survival (5,6,8-12), with our results for breast and colorectal cancer similar in magnitude to those previously reported (4-6,10) (prediagnosis and postdiagnosis PA HR = \sim 0.80 and 0.60, respectively, for cancer-specific and all-cause mortality). Findings reported here also indicate that PA contributes to survival benefits for prostate, lung, liver, hematologic, stomach, esophageal, and female reproductive cancers. Conversely, there was no evidence of harm from higher PA levels, even for cancers associated with poor prognosis (eg, lung cancer) or melanoma, which is the only cancer site for which higher levels of PA have been associated with higher risk of development.

Using data from studies involving women with breast cancer, we found a nonlinear relationship between increasing postdiagnosis PA levels and breast cancer-specific and all-cause

mortality hazards, up to about 10-15 MET hours per week. This level is consistent with approximately 150 weekly minutes of moderate-intensity PA or 75 weekly minutes of vigorousintensity PA and fits with the amount of PA recommended by the World Health Organization for healthy adults (160). This amount of PA is also typically endorsed and recommended by international cancer and clinical groups for those with cancer (13). Our findings also suggest that the clinical relevance of any potential survival benefit accrued through PA levels beyond 15 MET hours per week becomes less clear.

Questions remain regarding what represents the optimal dose, domain, and timing of activity for people with cancer and what these associations are for specific cancer sites or population subgroups. Findings from this meta-analysis show that there is clear evidence that postdiagnosis PA is an important independent prognostic factor distinct from prediagnosis activity levels. In addition, there is some preliminary evidence from three RCTs that exercise during treatment is also an important predictor of mortality outcomes (95,107,136). PA is also beneficial, irrespective of menopausal status, BMI, and sex, although being overweight or obese may attenuate the survival benefit. These findings highlight the need to combine weight (particularly fat mass) loss and PA interventions postcancer for those with BMIs greater than 25 kg/m². Currently, there are insufficient data to support specific recommendations related to

Table 2. Subgroup meta-analyses of the association between physical activity and cancer mortality, separately by sex, BMI, menopausal status, and colorectal subsite*

]	Prediagnosis physical	Postdiagnosis physical activity					
Subgroup	No. of studies/ No. of estimates	HR (95% CI)	P	I ²	No. of studies/ No. of estimates	HR (95% CI)	P	I^2
Cancer-specific mortality								
Sex								
All cancers (male)	18/18	0.80 (0.74 to 0.87)	<.001	75.50%	1/1	0.62 (0.44 to 0.87)	.006	-
All cancers (female)	16/16	0.86 (0.79 to 0.93)	<.001	61.70%	1/1	0.72 (0.47 to 1.10)	.130	-
Colorectal (male)	3/3	0.85 (0.53 to 1.34)	.478	76.50%	2/2	0.70 (0.38 to 1.28)	.247	66.60%
Colorectal (female)	5/5	0.67 (0.54 to 0.84)	.001	0.00%	3/3	0.50 (0.27 to 0.90)	.020	58.10%
BMI								
All cancers (<25 kg/m²)	3/3	0.77 (0.62 to 0.96)	.018	0.00%	_	_	_	-
All cancers (≥25 kg/m²)	2/2	0.91 (0.66 to 1.25)	.568	0.00%	_	_	_	_
Breast (<25 kg/m²)	4/4	0.92 (0.58 to 1.23)	.56	42.60%	7/7	0.59 (0.44 to 0.78)	<.001	49.70%
Breast (≥25 kg/m²)	4/4	0.76 (0.48 to 1.22)	.258	73.40%	7/8	0.61 (0.50 to 0.75)	<.001	50.20%
Colorectal (<25 kg/m²)	2/3	0.75 (0.59 to 0.96)	.021	19.20%	2/2	0.37 (0.07 to 1.94)	.239	71.80%
Colorectal (≥25 kg/m²)	2/3	0.79 (0.61 to 1.02)	.070	0.00%	2/2	0.78 (0.34 to 1.66)	.485	66.80%
Prostate (<25 kg/m²)	1/1	1.07 (0.55 to 2.11)	.844	_	_		_	_
Prostate (≥25 kg/m²)	1/1	1.53 (0.81 to 2.91)	.192	_	_	_	_	_
Menopausal status		,						
Breast (premenopausal)	5/5	1.11 (0.90 to 1.37)	.310	0.00%	5/5	0.65 (0.47 to 0.89)	.008	45.50%
Breast (postmenopausal)	7/7	0.93 (0.79 to 1.09)	.347	0.00%	7/7	0.68 (0.55 to 0.84)	<.001	48.60%
Colorectal subsite	•	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			·	,		
Colon	8/9	0.94 (0.80 to 1.11)	.448	34.80%	2/2	0.76 (0.58 to 0.99)	.044	0.00%
Rectum	8/9	0.79 (0.67 to 0.94)	.007	0.00%	2/2	0.60 (0.19 to 1.88)	.378	71.00%
All-cause mortality in cancer s	-•-	(-, -	()		,
Sex	a1111010							
All cancers (male)	_	_	_	_	1/1	0.52 (0.42 to 0.65)	<.001	_
All cancers (female)	_	_	_	_	1/1	0.62 (0.47 to 0.83)	.001	_
Colorectal (male)	3/3	0.73 (0.62 to 0.87)	<.001	0.00%	3/3	0.67 (0.56 to 0.80)	<.001	0.00%
Colorectal (female)	5/5	0.73 (0.59 to 0.91)	.006	18.00%	4/4	0.45 (0.30 to 0.68)	<.001	49.40%
BMI	3, 3	0.75 (0.55 to 0.51)	.000	10.0070	1/ 1	0.15 (0.50 to 0.00)	<.001	15.1070
Breast (<25 kg/m ²)	7/7	0.74 (0.60 to 0.91)	.005	56.70%	7/7	0.49 (0.35 to 0.68)	<.001	64.20%
Breast (\geq 25 kg/m ²)	7/8	0.81 (0.71 to 0.93)	.002	0.00%	7/11	0.70 (0.60 to 0.82)	<.001	24.30%
Colorectal (<25 kg/m ²)	1/1	0.78 (0.58 to 1.05)	.101	-	2/2	0.57 (0.45 to 0.73)	<.001	0.00%
Colorectal (≥25 kg/m²)	2/2	0.78 (0.58 to 1.03) 0.73 (0.58 to 0.92)	.009	0.00%	2/2	0.71 (0.47 to 1.08)	.112	38.60%
Hematologic (<25 kg/m²)	1/1	0.80 (0.67 to 0.96)	.015	-	1/1	0.54 (0.36 to 0.79)	.002	-
Hematologic (≥25 kg/m²)	1/1	0.83 (0.74 to 0.93)	.001	_	1/1	0.64 (0.50 to 0.82)	<.001	_
Menopausal status	1/ 1	0.03 (0.74 (0 0.33)	.001	-	1/ 1	0.04 (0.30 t0 0.62)	<.001	_
Breast (premenopausal)	4/4	0.86 (0.61 to 1.22)	.394	30.70%	4/4	0.77 (0.58 to 1.02)	.065	28.60%
Breast (premenopausal)	4/4 6/6	0.86 (0.81 to 1.22) 0.81 (0.70 to 0.94)	.006	30.70%	4/4 5/5	0.69 (0.63 to 0.77)	<.001	0.00%
\L /	0/0	0.61 (0.70 to 0.94)	.000	31.30%	5/5	0.03 (0.03 t0 0.77)	<.001	0.00%
Colorectal subsite	7/7	0.04 (0.71 += 0.00)	027	FC C00/	2/2	0 FC (0 40 to 0 75)	- 001	42.30%
Colon	7/7	0.84 (0.71 to 0.99)	.037	56.60%	3/3	0.56 (0.42 to 0.75)	<.001	
Rectum	6/6	0.84 (0.70 to 1.00)	.056	23.00%	2/2	0.88 (0.67 to 1.14)	.321	0.00%

^{*}BMI = body mass index; CI = confidence interval; HR = hazard ratio.

domain and dose of activity. For example, from a survival perspective, these epidemiologic findings support a PA dose of at least 10 METs, but not whether that dose is accumulated through recreational, transportation, occupational, or household activity, or mixed mode (aerobic vs resistance vs combined exercise) or specific intensity (moderate vs vigorous vs mixed). Nonetheless, findings are sufficiently compelling to support additional epidemiologic research, particularly on understudied cancer sites, subgroups within cancer sites, and more comprehensive measurement of PA (including during and posttreatment and domain, type, intensity, duration, and frequency). Further, these findings support the need for adequately powered, randomized, controlled exercise interventions that seek to

evaluate the impact of modifying recreational PA on cancer outcomes (161-164).

The magnitude of the effect of PA on cancer-specific and allcause mortality outcomes ranged from 0.46 to 1.19 for prediagnosis PA and cancer-specific survival, whereas for postdiagnosis activity the range was narrower and stronger (0.57-0.79 for cancer-specific survival). The range of effect sizes observed was similar for prediagnosis and postdiagnosis activity when considering all-cause mortality outcomes. For prediagnosis activity, estimates ranged from 0.47 to 0.92, and for postdiagnosis, the range was 0.37-0.96. Of interest, however, was that for cancer sites for which there were greater than 10 contributing point estimates (which occurred for all cancers combined, breast,

Table 3. Subgroup meta-analyses of the association between physical activity and cancer mortality, separately by domain of physical activity

		No. of studies/				No. of studies/			
Site	PA type	No. of estimates	HR (95% CI)	P	I^2	No. of estimates	HR (95% CI)	P	I^2
Cancer-speci	fic mortality								
All	Total	12/16	0.83 (0.75 to 0.92)	<.001	48.10%	2/2	0.66 (0.50 to 0.86)	.002	0.00%
	Recreational	24/27	0.82 (0.77 to 0.86)	<.001	68.20%	2/2	0.50 (0.24 to 1.02)	.057	67.80%
	Transportation	2/2	0.94 (0.82 to 1.07)	.362	0.00%	_	_	_	_
	Occupational	2/2	1.18 (0.70 to 1.98)	.530	61.00%	_	_	_	_
	Household	1/1	0.90 (0.54 to 1.49)	.684	_	_	_	_	_
Breast	Total	5/6	0.79 (0.63 to 0.99)	.043	0.00%	3/3	0.75 (0.47 to 1.21)	.236	0.00%
	Recreational	19/21	0.84 (0.75 to 0.94)	.002	35.40%	10/11	0.61 (0.47 to 0.78)	<.001	70.40%
	Transportation	_	· —	_	_	_		_	_
	Occupational	2/2	1.03 (0.80 to 1.33)	.802	0.00%	_	_	_	_
	Household	1/1	1.25 (0.81 to 1.94)	.317	_	_	_	_	_
Colorectal	Total	2/2	0.84 (0.73 to 0.96)	.010	0.00%	1/1	0.88 (0.68 to 1.15)	.340	_
	Recreational	10/12	0.78 (0.70 to 0.87)	<.001	0.00%	5/7	0.48 (0.34 to 0.67)	<.001	10.50%
	Transportation	1/2	1.00 (0.63 to 1.58)	.989	0.00%	_		_	_
	Occupational	_	· — ,	_	_	_	_	_	_
	Household	_	_	_	_		_	_	_
Prostate	Total	3/3	0.94 (0.70 to 1.27)	.697	7.20%	2/2	0.55 (0.36 to 0.87)	.010	0.00%
	Recreational	7/7	0.85 (0.70 to 1.04)	.108	44.70%	3/3	0.71 (0.56 to 0.91)	.007	14.30%
	Transportation	1/1	1.65 (0.87 to 3.14)	.127	_	1/1	0.64 (0.43 to 0.95)	.025	_
	Occupational	2/2	0.89 (0.59 to 1.35)	.580	0.00%	1/1	0.90 (0.53 to 1.54)	.700	_
	Household	1/1	0.78 (0.49 to 1.24)	.294	_	2/2	1.02 (0.76 to 1.36)	.911	0.00%
All-cause mo	rtality in cancer su	ırvivors	,				,		
All	Total	_	_	_	_	3/3	0.55 (0.47 to 0.65)	<.001	0.00%
	Recreational	1/1	0.47 (0.29 to 0.75)	.002	_	5/5	0.63 (0.50 to 0.79)	<.001	50.80%
	Transportation	_		_	_	_		_	_
	Occupational	_	_	_	_	_	_	_	_
	Household	_	_	_	_	1/1	1.04 (0.60 to 1.80)	.889	_
Breast	Total	5/6	0.84 (0.67 to 1.05)	.126	32.80%	6/6	0.60 (0.47 to 0.75)	<.001	0.00%
	Recreational	16/18	0.81 (0.76 to 0.87)	<.001	16.70%	11/12	0.58 (0.51 to 0.66)	<.001	47.10%
	Transportation	_	` —	_	_	_	` _	_	_
	Occupational	2/2	1.09 (0.88 to 1.35)	.421	0.00%	_		_	_
	Household	1/1	1.46 (1.02 to 2.09)	.039	_	1/1	0.93 (0.55 to 1.55)	.784	_
Colorectal	Total	2/2	0.92 (0.80 to 1.06)	.237	0.00%	3/3	0.77 (0.57 to 1.03)	.080	84.60%
	Recreational	8/8	0.76 (0.70 to 0.84)	<.001	0.00%	7/9	0.58 (0.49 to 0.69)	<.001	11.60%
	Transportation	_	_ ′	_	_	_		_	_
	Occupational	_	_	_	_	_	_	_	_
	Household	_	_	_	_	1/1	0.83 (0.55 to 1.23)	.364	_
Prostate	Total	1/1	1.02 (0.77 to 1.35)	.89		2/2	0.47 (0.31 to 0.71)	<.001	68.90%
	Recreational	2/2	0.87 (0.80 to 0.96)	.004	0.00%	4/4	0.69 (0.56 to 0.85)	<.001	71.80%
	Transportation	_	—	_	_	1/1	0.64 (0.43 to 0.94)	.025	_
	Occupational	1/1	1.35 (1.00 to 1.81)	.047	_	1/1	0.64 (0.47 to 0.91)	.011	_
	Household	1/1	0.91 (0.70 to 1.18)	.474	_	2/2	0.82 (0.70 to 0.97)	.023	0.00%

 $^{^*}CI = confidence interval; HR = hazard ratio; PA = physical activity.$

colorectal, and prostate cancers), there was greater consistency of the evidence. This range of effect sizes for cancer-specific survival was reduced to 0.80–0.90 for prediagnosis PA and 0.62–0.70 for postdiagnosis PA, and for all-cause survival, the range was 0.80–0.82 for prediagnosis PA and 0.58–0.63 for postdiagnosis PA. Hence, as the evidence base is accumulating, despite differences in study populations, study designs, and PA assessment methods, there is remarkable consistency of the effects of prediagnosis and postdiagnosis PA across various cancer sites.

Despite the exponential increase in the number of studies conducted on this topic since the mid-2000s, there is still a paucity of evidence for most cancer sites with only breast, colorectal, and prostate cancers approaching the number of studies required per site for meta-analyses by site and within

population subgroups. To understand whether current differences observed in effect size are cancer specific or due to imprecision, more research beyond these top three cancer sites is needed. Additional limitations of this meta-analysis include the heterogeneous PA assessment methods. We mitigated, as much as possible, the impact of different PA assessment methods by selecting, wherever possible, point estimates expressed in units of MET hours per week. In addition, differences in adjustment for confounding and examination of effect modification also make comparisons across studies more challenging and can adversely influence the precision of summary estimates reported. We examined this issue with our quality assessment of the 136 included studies, which determined that these studies, overall, had high quality of conduct, adding credibility to the findings reported here.

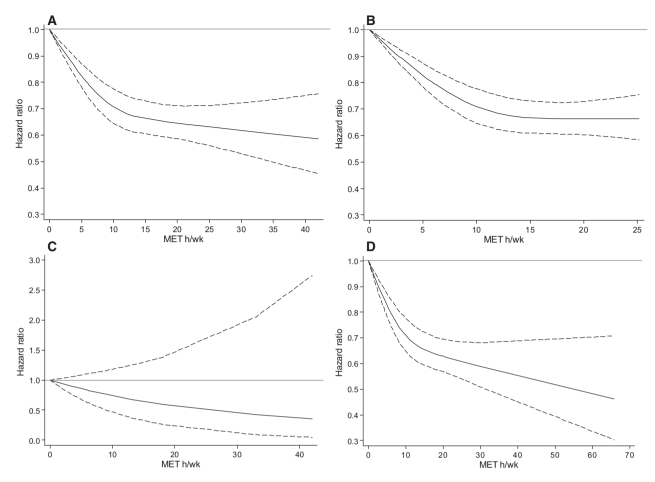


Figure 4. Random-effects dose-response curves for recreational physical activity in breast cancer survivors. A) Prediagnosis physical activity and breast cancer-specific mortality (n = 7 sets of data from six studies); B) postdiagnosis physical activity and breast cancer-specific mortality (n = 7 sets of data from six studies); C) prediagnosis physical activity and all-cause mortality (n = 5 sets of data from four studies); D) postdiagnosis physical activity and all-cause mortality (n = 8 sets of data from seven studies). MET = metabolic equivalent.

We were unable to examine the associations between PA and cancer recurrence, progressions, or other cancer outcomes because of the heterogeneous definitions used across the source studies. Likewise, an interest in precision exercise oncology is to examine how cancer population subgroups, defined by clinical or pathologic characteristics, respond to PA (165). To date, few studies have examined these clinicopathologic subgroups to identify which populations might benefit more from PA. With additional research on this topic and the prerequisite that future studies follow standardized definitions of outcomes (eg, STEEP guidelines) and comprehensively report patient and tumor characteristics, analyses by specific outcomes will also be possible and highly informative (166). Finally, future studies are needed that use the highest quality of PA assessment with objective and self-report measures and the reporting in MET hours per week to permit additional evaluations of the dose-response effects in other cancer sites.

In summary, we found strong evidence that PA before or after cancer diagnosis was associated with statistically significant decreased hazards of cancer-specific and all-cause mortality in at least 11 different cancer sites. In addition, we found that hazard of CVD mortality among cancer survivors was also reduced with PA. As such, these findings confirm the importance of promoting PA after cancer and suggest that in doing so, there is huge potential for patient and public health gain through PA.

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CMF, SCH, and CRS designed and conceptualized the study; CRS conducted the literature search and eligibility review, abstracted the study details and results, contacted authors for additional details, conducted the analysis, prepared the tables and figures, and drafted the study methods and results; CMF wrote the final paper with input from SCH and WYC. WYC also provided input on subgroup analyses. All authors reviewed and approved the final draft. The corresponding author had full access

to all the data in the study and had final responsibility for the decision to submit for publication.

The authors declare no competing interests.

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