UCSF

UC San Francisco Previously Published Works

Title

Post-diagnostic health behaviour scores and risk of prostate cancer progression and mortality

Permalink

https://escholarship.org/uc/item/82w3610k

Journal

British Journal of Cancer, 129(2)

ISSN

0007-0920

Authors

Langlais, Crystal S Graff, Rebecca E Van Blarigan, Erin L et al.

Publication Date

2023-08-10

DOI

10.1038/s41416-023-02283-1

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at https://creativecommons.org/licenses/by/4.0/

Peer reviewed

ARTICLE OPEN



Epidemiology

Post-diagnostic health behaviour scores and risk of prostate cancer progression and mortality

Crystal S. Langlais 1.4[™], Rebecca E. Graff 1.2, Erin L. Van Blarigan 1.2,3, John M. Neuhaus 1, Janet E. Cowan 3, Jeanette M. Broering 3, Peter Carroll 2.3, Stacey A. Kenfield 1.2,3,5 and June M. Chan 1.2,3,5

© The Author(s) 2023

BACKGROUND: Individual behaviours are associated with prostate cancer (PC) progression. Behavioural scores, comprised of multiple risk factors, allow assessment of the combined impact of multiple behaviours.

METHODS: We examined the association between six a priori scores and risk of PC progression and mortality among 2156 men with PC in the Cancer of the Prostate Strategic Urologic Research Endeavor (CaPSURE) cohort: two scores developed based on the PC survivorship literature ('2021 Score [+ Diet]'); a score developed based on pre-diagnostic PC literature ('2015 Score'); and three scores based on US recommendations for cancer prevention ('WCRF/AICR Score') and survival ('ACS Score [+ Alcohol]'). Hazard ratios (HRs) and 95% confidence intervals (Cls) were estimated for progression and PC mortality via parametric survival models (interval censoring) and Cox models, respectively.

RESULTS: Over a median (IQR) of 6.4 (1.3, 13.7) years, we observed 192 progression and 73 PC mortality events. Higher (i.e., healthier) 2021 Score + Diet and WCRF/AICR Scores were inversely associated with risk of PC progression (2021 + Diet: $HR_{continuous} = 0.76$, 95% CI: 0.63–0.90. WCRF/AICR: $HR_{continuous} = 0.83$, 95% CI: 0.67–1.02) and mortality (2021 + Diet: $HR_{continuous} = 0.65$, 95% CI: 0.45–0.93. WCRF/AICR: $HR_{continuous} = 0.71$; 95% CI: 0.57–0.89). The ACS Score + Alcohol was only associated with progression ($HR_{continuous} = 0.89$, 95% CI: 0.81–0.98) while the 2021 Score was only associated with PC mortality ($HR_{continuous} = 0.62$, 95% CI: 0.45–0.85). The 2015 was not associated with PC progression or mortality.

CONCLUSION: Findings strengthen the evidence that behavioural modifications following a prostate cancer diagnosis may improve clinical outcomes.

British Journal of Cancer (2023) 129:346-355; https://doi.org/10.1038/s41416-023-02283-1

BACKGROUND

Prostate cancer is the most commonly diagnosed cancer among men in the United States (US), with 248,530 new cases expected to have occurred in 2021 [1, 2]. Currently, there are over 3.6 million prostate cancer survivors in the US [3]. Though the 5-year survival rate for prostate cancer approaches 100%, there remains uncertainty regarding which cancers will eventually progress, and prostate cancer remains the second leading cause of cancer death among US men [2, 4, 5]. To inform interventions and mitigate risk of progression and prostate cancer-specific mortality (PCSM) for the large population of men living with the disease, there is a need to better understand how behavioural factors *after* diagnosis influence disease progression.

Several studies have linked modifiable risk factors with prostate cancer progression and PCSM [6]. However, prior reports have predominantly focused on individual exposures, which do not fully reflect the complex relationships among multiple diet and other behavioural factors [7, 8]. For example, physical activity may offset some of the negative effects of unhealthy dietary choices

[9]. Therefore, scores that reflect multiple behavioural factors may be more strongly associated with outcomes among men with prostate cancer than individual health habits.

Our team previously conducted an extensive review summarising the literature on post-diagnostic behaviours and prostate cancer progression and PCSM [6]. Using that report, we developed a prostate cancer-specific behavioural scores ("2021 Score [+ Diet]"). Here, we examine the association of these scores in relation to risk of progression and PCSM among men with non-metastatic prostate cancer in the Cancer of the Prostate Strategic Urologic Research Endeavor (CaPSURE) cohort. Further, for completeness and comparability with other studies, we evaluated associations of four other scores developed to inform the risk of cancer onset or progression to understand if adherence to general cancer prevention or survivorship guidelines may improve outcomes following a prostate cancer diagnosis. One of these scores was developed by members of our team to predict the risk of developing incident lethal prostate cancer based on pre-diagnostic behaviours ("2015 Score") [10]. It is distinct from the 2021 Scores focused on post-

¹Department of Epidemiology and Biostatistics, University of California, San Francisco, CA, USA. ²Helen Diller Family Comprehensive Cancer Center, University of California, San Francisco, CA, USA. ³Department of Urology, University of California, San Francisco, CA, USA. ⁴Present address: IQVIA, Durham, NC, USA. ⁵These authors contributed equally: Stacey A. Kenfield, June M. Chan. [™]email: crystal.langlais@ucsf.edu

Received: 24 August 2022 Revised: 11 April 2023 Accepted: 14 April 2023

Published online: 22 May 2023

diagnostic behaviours as the known behavioural risk factors for prostate cancer risk and progression differ. The other three are operationalized versions of the American Cancer Society (ACS) cancer survivorship recommendations [11, 12] and the World Cancer Research Fund (WCRF)/American Institute for Cancer Research (AICR) cancer prevention recommendations [13, 14]. We hypothesised that men with healthier lifestyles (i.e., higher scores) would have lower risk of disease progression and mortality.

METHODS

Study sample

CaPSURE is a longitudinal observational cohort of 15,310 men with biopsyproven prostate cancer. Men diagnosed between 1999 and 2018 at any of 43 participating urology practices across the US were eligible. Participating urologists provided data on clinical and pathological features, treatments, and clinical follow-up. Additional details on CaPSURE are reported elsewhere [15]. The study was conducted in accordance with the Belmont Report and U.S. Common Rule under local Institutional Review Board approval, with all participants providing written informed consent.

The CaPSURE Diet and Lifestyle (CDL) sub-study—consisting of a comprehensive lifestyle questionnaire and full-length food frequency questionnaire (FFQ)—was administered at three time points between 2004 and 2016; a total of 2891 men participated in at least one administration. For the subset of men who completed more than one questionnaire (n = 443), only the first administration (closest to diagnosis date) was used. We excluded men with last clinical follow-up or documented progression prior to completing their first CDL questionnaire (n = 551). Consistent with the recommended approach to address implausible energy intakes [16], we excluded men with extreme (<800 kcal/day or >4200 kcal/day) or unknown caloric intake (n = 153) and/or missing ≥ 70 FFQ items (n = 20). Finally, we excluded men without a discernable clinical T-stage (n = 100) or with a clinical T-stage >T3a (n = 8) and those with death from unknown cause (n = 3). These exclusions left us with a sample size of 2056 men for our primary analyses of prostate cancer progression. Following the exclusion of men with documented progression prior to completing the first questionnaire, the subsequent exclusions resulted in the loss of 23 events, 2 of which were PCSM. For PCSM analyses, men who were excluded due to documented progression prior to completion of the CDL questionnaire were included—death could not occur prior to completing the questionnaire—resulting in a sample size of 2447 men.

Diet and lifestyle questionnaire

Dietary intake was self-reported on a validated [17–19] semiquantitative FFQ, wherein men reported how frequently they consumed a standard unit or portion size of approximately 140 different items. The nine frequency options ranged from never or less than once per month to six or more times per day. FFQ data were sent to the Nutrition Department at the Harvard T.H. Chan School of Public Health, which calculated total intake of nutrients, including total caloric intake and grams of whole grains, fibre, and alcohol. Nutrient intake was calculated by multiplying the nutrient value in the specified portion size of each item on the FFQ by its frequency of intake and then summing across all items. Nutrient values were obtained from the US Department of Agriculture databases [20] supplemented with other sources.

The survey asked men if they had smoked 20 packs of cigarettes or more in their lifetime. If they responded "yes", they were asked to report additional details regarding their smoking history. Men who responded "no" were considered never smokers.

Men completed a validated physical activity questionnaire which asked them to report their average weekly time spent doing nine types of aerobic and resistance training activities over the prior year [21]. Ten frequency options could be selected, ranging from 0 minutes to 11 or more hours per week. Participants were also asked about their regular walking pace and ability/frequency of climbing stairs.

Other information collected on the survey included height and weight [used to calculate body mass index (BMI; kg/m²)]; education level; a brief medical history, including family history of prostate cancer; and a detailed history of the use of vitamins and supplements.

Behavioural scores

Six a priori scores were evaluated, as described below. All scores were oriented such that increasing values reflected healthier behaviours. Please see Supplemental Tables S1 and S2 for additional details.

2021 score. The 2021 (post-diagnostic) Score was based on an extensive literature review conducted in 2021, summarising behaviours following a prostate cancer diagnosis associated with risk of recurrence, progression, and/or PCSM [6]. To determine the factors for inclusion in the score, we searched PubMed using the terms "prostate cancer" and "progression or mortality" in combination with terms describing individual lifestyle factors. Factors considered for the score included those that (1) exhibited a statistically significant association with metastases or PCSM in at least one study and (2) were corroborated by at least one additional study with an association in the same direction, whether or not statistically significant. In total, we identified seven such factors—smoking status [22-31]; BMI [32–48]; physical activity [49–53]; and intake of saturated fat [54–56], whole milk [57, 58], wine [59, 60], and processed meat [61, 62]. The three nondietary factors demonstrated the strongest evidence in the literature review. We examined two versions of the 2021 Score, one without ("2021 Score") and one with ("2021 Score + Diet") the dietary components. The points per behaviour component ranged from 0 to 1 (see Supplement Table S1), with the points for the four dietary components (whole milk, alcohol, red and processed meat, saturated fat) averaged using the arithmetic mean to create a single dietary sub-score ranging from 0 to 1. This approach was consistent with the operationalization of the ACS recommendations into the ACS Score [13]. The point values were based on where the risk associated with prostate cancer outcomes appeared to change in the literature. The points for each component were summed to create the total 2021 Score (range: 0–3) and 2021 Score + Diet (range: 0–4) for each participant.

2015 score. Our team previously developed the 2015 (pre-diagnostic) Score to identify the risk of developing lethal prostate cancer among healthy men, based on the evidence available circa 2014 [10]. The six components—smoking status, BMI, physical activity, fatty fish intake, tomato intake, and processed red meat intake—were each scored as 0 or 1 based on cut-points associated with risk as reported in the literature at the time of score creation. The sub-scores were then summed to create the total 2015 Score (range: 0–6). Components of the 2015 Score were identified based on the existing literature available in 2015.

ACS score. To create a primary and an alternative ACS Score, we expanded on the operationalization of the ACS Nutrition and Physical Activity Guidelines for Cancer Survivors developed by McCullough et al. [13]. Each of the three components—BMI, physical activity, and dietary—were scored from 0 to 2 and then summed to create the primary ACS Score (range: 0–6). The dietary component included total servings and variety of fruits and vegetables, red and processed meat intake, and whole grain intake. We expanded to include strength training when assigning physical activity points, consistent with the guidelines. The "ACS Score + Alcohol" additionally included alcohol intake, scored from 0 to 2 (with the highest score for moderate alcohol intake: >0 to 2 servings/day), reflecting the inclusion of alcohol in the ACS recommendations for cancer prevention but not cancer survival (alternative score range: 0–8).

WCRF/AICR score. The WCRF/AICR Cancer Prevention Recommendations were operationalized based on published scoring guidelines [11, 12] and included BMI; physical activity; and intake of alcohol, sugar-sweetened beverages, fruits/vegetables, fibre, red and processed meat, and adapted ultra-processed foods (range: 0–7).

Outcome

The primary outcome was time to prostate cancer progression, defined as biochemical recurrence, secondary treatment, bone metastases, or PCSM, as applied previously [9, 53, 63]. Given the small number of PCSM events (n=73) in this cohort, PCSM was evaluated as a secondary outcome.

Biochemical recurrence was defined as two consecutive prostate-specific antigen (PSA) readings ≥ 0.2 ng/mL following radical prostatectomy or a rise of 2.0 ng/mL above post-radiation nadir on two consecutive PSA readings; the date of recurrence was recorded as the date of the second elevated PSA. Secondary treatment was defined as any treatment started at least 6 months following primary treatment. Bone metastases included prostate cancer progression to bone, advancement to TNM stage M1b, a positive bone scan, and radiation to treat bone metastases. Cause of death was determined by the registry data coordinating centre and through confirmation by either the vital statistics official death certificate from the state in which the death occurred or by the National Centre for Health Statistics National Death Index [64]. Deaths were attributed to prostate cancer if the death certificate

included ICD-9 code 185 [(metastatic) malignant neoplasm of prostate] as the primary or secondary cause of death.

Time to progression was measured from completion date of the CDL guestionnaire to the date of progression (first event of biochemical recurrence, secondary treatment, bone metastases, or PCSM). For men with documented non-PCSM progression (i.e., recurrence, secondary treatment, or bone metastasis failure events), the censoring interval (i.e., window in which the event occurred) was bound by the last normal clinical visit (left limit) and the clinical visit documenting evidence of progression (right limit). For men who died from prostate cancer, the left and right limit were both date of death. Men without documented progression or PCSM were censored at their last date of follow-up or death (other cause); thus, the left limit of their censoring interval was defined by the last clinical follow-up date or date of death (non-PCSM), respectively, and the right limit was undefined (i.e., censored). Clinical follow-up was last consistently assessed across all CaPSURE sites on January 31, 2019; 26 men had a last known clinical follow-up date beyond this date and were administratively censored on that date.

Statistical analysis

Parametric survival models with a Weibull distribution were used to accommodate interval censoring associated with uncertainty in actual date of prostate cancer progression [65]. Because the date of death is known for PCSM (i.e., interval censoring was not an issue), we utilised Cox proportional hazards models rather than parametric survival methods when assessing the PCSM outcome. Proportional hazards assumptions were assessed visually by plotting the scaled Schoenfeld residuals against follow-up time.

We fit survival models using both continuous scores (per 1-unit change) and tertiles of scores. All models were clustered by CaPSURE clinical site with robust standard errors used to calculate confidence intervals (CI). Simple models were adjusted for time between diagnosis and participants' first CDL questionnaire (continuous) and age at diagnosis. A directed acyclic graph (DAG) was developed to reflect our understanding of the complex relationship of interest; variables identified in the DAG were included as covariates in the fully adjusted models, as appropriate [66, 67]. Fully adjusted models were additionally adjusted for clinical T-stage (T1, T2, T3a), Gleason score (<7, 7, >7), diagnostic PSA level (≤6 ng/mL, >6 to 10 ng/mL, >10 ng/mL), primary treatment (radical prostatectomy, radiation, hormonal therapy, watchful waiting/active surveillance, other), family history of prostate cancer in a brother or father (yes, no), self-identified and physician-reported race (white, non-white), selenium supplement use (non-user; <140 μg/day; ≥140 μg/day; user with unknown daily dosage), total caloric intake (continuous, kcal/d), and the following variables if not part of the score of interest: whole milk intake (≤4 servings/week, >4 servings/week), wine intake (3–14 servings/week, <3 or ≥14 servings/ week), alcohol intake (non-drinker, >0-2 servings/day, >2 servings/day), red and processed meat intake (quartiles), tomato intake (continuous, servings/day), dark fish intake (continuous, servings/day), and smoking (never, quit ≥10 years prior, quit <10 years prior, current). We further considered adjustment for comorbidities (diabetes, stroke, prior myocardial infarction, or other heart disease; yes/no) but the magnitudes of the estimates changed very little with adjustment, so these variables were not included in the final models.

We assessed potential interaction between each of the scores and age at diagnosis (<65 years, ≥65 years) and, separately, stage at diagnosis (T1, T2–T3a) by adding interaction terms with the scores in the models and using Wald tests. Given statistically non-significant Wald tests and small magnitudes of estimated interaction regression coefficients, interaction terms were not included in the final models. We examined goodness-of-fit of the survival models using Cox–Snell residual plots. Across all scores, goodness-of-fit was best in the fully adjusted models, with decreasing fit in the tails. Fully adjusted models for progression were also run using exponential distributions, which produced Cox–Snell residual plots that demonstrated poorer fit than Weibull models and thus were not reported.

Sensitivity analyses

First, we were concerned about confounding due to PSA surveillance after diagnosis (i.e., men with healthier behaviours may be more likely to be monitored via PSA tests, potentially creating a positive correlation between healthy lifestyle habits and risk of progression). To address this, Poisson regression was utilised to compare the number of PSA visits to tertile of each of the six scores, with the lowest tertile (i.e., the least healthy group) as the reference. Total follow-up time was used as an offset in these models.

Second, whereas our primary analyses used time of the CDL questionnaire completion as time zero—which necessitates excluding men who experienced an event prior to the survey—sensitivity analyses re-assigned time zero as time of diagnosis. These analyses assumed that the responses on the CDL questionnaire were consistent with what would have been measured at the date of diagnosis. Men excluded from our primary analyses due to documented progression prior to CDL questionnaire were included in these sensitivity analyses, resulting in an analytic sample of 2447 men. For this approach, we first assessed whether there was an interaction between year of diagnosis and each of the behavioural scores by adding an interaction term with the scores in the models and using Wald tests; no evidence of interaction was found.

Third, we were interested in understanding how competing events (i.e., deaths due to causes other than prostate cancer) impacted our primary results. Methods to address competing events in the presence of interval censoring are not readily available or accessible. Thus, we ran Cox proportional hazards models for progression and compared these results to Fine-Gray analyses accounting for other deaths as a competing risk. Proportional hazards assumptions were assessed visually by plotting the scaled Schoenfeld residuals against follow-up time.

Lastly, missingness in the covariates resulted in a loss of events in our fully adjusted models. Specifically, men with missing data for any of the score components were excluded from the primary analysis for that score: n = 60 for 2021 Score, n = 60 for the 2021 Score with Diet, n = 83 for 2015 Score, n = 40 for ACS Score, n = 70 for the ACS Score with Alcohol, and n = 43 for WCRF/AICR Score. To understand the impact of this missingness on our primary results, we performed sensitivity analyses utilising multiple imputation to handle missing data [68], which assumes that data are missing at random. We assessed the plausibility of this assumption by summarising participant characteristics by missingness status for each of the six scores. We performed multiple imputation via chained equations using the *chained* command in Stata to first generate 25 imputed datasets. We then fit survival models across all 25 imputed datasets and pooled the results using Rubin's Rules [69]. Our imputed models included fully observed variables (CaPSURE clinical site, age at diagnosis, BMI, days of follow-up, total energy intake, tomato intake, days from CDL return to the left interval of follow-up time, race, diagnostic T-stage, and family history of prostate cancer) and variables with incomplete values (diagnostic PSA and Gleason score; total alcohol, whole milk, dark fish, total wine, and red and processed meat intake; each of the scores; smoking status; and primary treatment).

All statistical analyses were performed using Stata version 17 (StataCorp, College Station, TX) using a two-sided alpha level of 0.05 to assess statistical significance.

RESULTS

In our main analyses, the 2056 men who met inclusion criteria were followed for a median of 6.4 years (IQR: 1.3, 12.7) after completing the CDL questionnaire, for a total of 13,102 person-years. During the follow-up period, 192 had documented progression, including 168 (88%) with biochemical recurrence, 7 (4%) with bone metastases, and 17 (9%) deaths related to prostate cancer as the first recorded event (there were 73 PCSM events in total). There were 384 all-cause deaths during the observation period. Baseline characteristics by tertile of each of the four primary scores are shown in Table 1. Most participants identified as white race with a diagnostic T-stage of 1 and Gleason grade <7 and underwent radical prostatectomy as their primary treatment. Characteristics were balanced across tertiles of the scores.

Progression

Those with higher 2021 Scores had a non-statistically significant lower risk of progression (HR_{cont}: 0.84, 95% CI: 0.65–1.08); however, in models assessing score tertiles, there was no clear association with progression (HR_{2 vs} $_1$: 0.90, 95% CI: 0.62–1.32; HR_{3 vs} $_1$: 0.79, 95% CI: 0.50–1.24; $P_{\rm trend}=0.30$). Including dietary factors in the 2021 Score (2021 Score + Diet) strengthened the associations: HR_{cont}: 0.76, 95% CI: 0.63–0.90; HR_{2 vs} $_1$: 0.82, 95% CI: 0.62–1.08 and HR_{3 vs} $_1$: 0.67; 95% CI: 0.44–1.02 ($P_{\rm trend}=0.06$) (Table 2 and Fig. 1).

Neither the 2015 Score (HR_{cont}: 0.89, 95% CI: 0.80–1.00; HR_{2 vs 1}: 0.90, 95% CI: 0.60–1.35; HR_{3 vs 1}: 0.57, 95% CI: 0.30–1.09; $P_{\text{trend}} =$

Table 1. Patient and clinical characteristics of men diagnosed with non-metastatic prostate cancer by tertile of health behaviour scores.

	2021 Score ^a	e e		2015 Score ^a	e e		ACS Score ^a			WCRF/AICR Score ^a	Scorea	
N ^b :	1996			1973			2016			2013		
Tertile: Point range:	1st 0-2	2nd 2.5	3rd 3	1st 0-3	2nd 4	3rd 5-6	1st 0-2	2nd 2.5–3	3rd 3.5-6	1st 0.75-3.25	2nd 3.5-4	3rd 4.25-7
u	1033	640	323	1259	475	239	710	610	969	805	260	648
Age ^c (years), mean (SD)	64.5 (7.9)	64.0 (7.9)	64.7 (8.3)	64.6 (7.9)	64.2 (7.9)	63.3 (8.1)	64.0 (7.8)	64.8 (8.0)	64.3 (8.1)	64.4 (7.8)	64.4 (7.9)	64.3 (8.2)
White, <i>n</i> (%)	979 (95)	(26) 609	310 (96)	1202 (95)	455 (96)	220 (92)	672 (95)	285 (96)	(92)	764 (95)	531 (95)	620 (96)
T-Stage ^c , <i>n</i> (%)												
11	598 (58)	367 (57)	184 (57)	(52) (69	284 (60)	156 (65)	407 (57)	336 (55)	416 (60)	483 (60)	324 (58)	350 (54)
12	426 (41)	268 (42)	134 (41)	550 (44)	186 (39)	81 (34)	297 (42)	270 (44)	271 (39)	315 (39)	230 (41)	292 (45)
T3a	9 (1)	5 (1)	5 (2)	12 (1)	5 (1)	2 (1)	(1)	4 (1)	9 (1)	7 (1)	(1)	6 (1)
Gleason ^c , <i>n</i> (%)												
<7	(99) 089	435 (69)	223 (69)	816 (65)	332 (70)	174 (73)	472 (67)	404 (67)	470 (68)	518 (65)	388 (70)	440 (68)
7	278 (27)	164 (26)	75 (23)	343 (28)	115 (24)	51 (21)	178 (25)	168 (28)	182 (26)	219 (28)	138 (25)	167 (26)
>7	(2)	35 (6)	23 (7)	87 (7)	25 (5)	13 (5)	55 (8)	31 (5)	40 (6)	59 (7)	30 (5)	37 (6)
PSA ^c (ng/ml), median (IQR)	5.6 (4.4, 8.0)	5.5 (4.3, 7.6)	5.3 (4.3, 7.5)	5.6 (4.4, 8.0)	5.5 (4.5, 7.6)	5.2 (4.1, 7.6)	5.6 (4.4, 8.0)	5.6 (4.4, 7.7)	5.4 (4.3, 7.8)	5.7 (4.4, 8.0)	5.5 (4.3, 7.7)	5.4 (4.3, 7.7)
Primary treatment, n (%)												
Radical prostatectomy	604 (60)	402 (65)	210 (67)	769 (62)	282 (61)	153 (67)	407 (59)	376 (63)	448 (66)	473 (60)	340 (62)	413 (66)
AS/WW	26 (6)	39 (6)	23 (7)	64 (5)	33 (7)	19 (8)	32 (5)	37 (6)	49 (7)	40 (5)	36 (7)	42 (7)
RT/brachytherapy	245 (24)	129 (21)	58 (19)	281 (23)	101 (22)	43 (19)	175 (25)	129 (22)	131 (19)	189 (24)	123 (22)	124 (20)
Hormone therapy	61 (6)	26 (4)	11 (4)	61 (5)	25 (5)	9 (4)	40 (6)	40 (7)	21 (3)	46 (6)	30 (5)	25 (4)
Other	47 (5)	21 (3)	11 (4)	57 (5)	19 (4)	4 (2)	37 (5)	14 (2)	27 (4)	35 (4)	22 (4)	22 (4)
Family history of PC,	186 (18)	141 (22)	79 (24)	246 (20)	105 (22)	51 (21)	143 (20)	115 (19)	151 (22)	171 (21)	121 (22)	116 (18)

²2021 Score included smoking, and physical activity. 2015 Score included smoking, BMI, physical activity, fatty fish intake, tomato intake, and processed red meat intake. ACS Score included BMI, physical activity, alcohol intake, red and processed meat intake, and whole grain intake. WCRF/AICR Score included BMI, physical activity, alcohol intake, sugar-sweetened beverage intake, percentage ACS American Cancer Society, AICR American Institute for Cancer Research, AS/WW active surveillance/watchful waiting, aUPFs adapted ultraprocessedfoods, BMI body mass index, IQR interquartile range, PC prostate cancer, RT radiation therapy, SD standard deviation, WCRF World Cancer Research Fund.

of calories from aUPFs, fiber intake, whole fruit and vegetable intake, and red and processed meat intake. b Missingness on score components resulted in missing scores: n = 60 for 2021 Score, n = 83 for 2015 Score, n = 40 for ACS Score, and n = 43 for AICR Score.

At prostate cancer diagnosis.

Table 2. Post-diagnostic health behaviour scores and the risk of prostate cancer progression among men with non-metastatic prostate cancer, estimated via parametric (Weibull) survival models.

2021 Score—HR	(95% CI)												
	Events	N	Continuous			1st Tertile	2nd Tertile			3rd Te	ertile		$P_{\rm trend}$
						(0-2 pts)	(2.5 pt	ts)		(3 pts))		
Simple ^a	188	1996	0.79	(0.66,	0.95)	Ref	0.77	(0.57,	1.04)	0.69	(0.46,	1.04)	0. 07
Fully adjusted ^b	146	1615	0.84	(0.65,	1.08)	Ref	0.90	(0.62,	1.32)	0.79	(0.50,	1.24)	0.30
2021 Score + Die	et—HR (95	% CI)											
	Events	N	Contin	nuous		1st Tertile	2nd Te	ertile		3rd Te	rtile		P_{trend}
						(0.25-2.25 pts)	(2.5-3	pts)		(3.25-	4 pts)		
Simple ^a	188	1996	0.78	(0.68,	0.90)	Ref	0.83	(0.67,	1.04)	0.67	(0.46,	0.97)	0.03
Fully adjusted ^b	151	1673	0.76	(0.63,	0.90)	Ref	0.82	(0.62,	1.08)	0.67	(0.44,	1.02)	0.06
2015 Score—HR	(95% CI)												
	Events	N	Contin	nuous		1st Tertile	2nd Te	ertile		3rd Te	rtile		P_{trend}
						(0-3 pts)	(4 pts)			(5–6 p	ots)		
Simple ^a	183	1973	0.85	(0.78,	0.94)	Ref	0.82	(0.59,	1.14)	0.56	(0.33,	0.94)	0.03
Fully adjusted ^b	141	1611	0.89	(0.80,	1.00)	Ref	0.90	(0.60,	1.35)	0.57	(0.30,	1.09)	0.09
ACS Score—HR ((95% CI)												
	Events	N	Contin	nuous		1st Tertile	2nd Te	ertile		3rd Te	ertile		$P_{\rm trend}$
			Continuous 6 0.89 (0.80, 0.98)			(0-2 pts)	(2.5-3	pts)		(3.5–6	pts)		
Simple ^a	188	2016	0.89	(0.80,	0.98)	Ref	0.93	(0.68,	1.28)	0.71	(0.53,	0.94)	0.02
Fully adjusted ^b	146	1614	0.93	(0.82,	1.05)	Ref	1.19	(0.82,	1.71)	0.83	(0.58,	1.18)	0.30
ACS Score + Alco	ohol—HR (95% CI)											
	Events N		Continuous		1st Tertile	2 nd Te	rtile		3rd Te	ertile		P_{trend}	
			Continuous			(0-3.5 pts)	(4–5 pts)			(5.5-8 pts)			
Simple ^a	182	1986	0.88	(0.80,	0.96)	Ref	0.77	(0.56,	1.05)	0.49	(0.29,	0.83)	0.008
Fully adjusted ^b	146	1614	0.89	(0.81,	0.98)	Ref	0.97	(0.71,	1.32)	0.48	(0.28,	0.82)	0.007
WCRF/AICR Score	e—HR (959	% CI)											
	Events N		Continuous		1st Tertile	2nd Te	ertile		3rd Te	ertile		P_{trend}	
						(0.75-3.25 pts)	(3.5–4	pts)		(4.25–	7 pts)		
Simple ^a	188	2013	0.85	(0.72,	1.01)	Ref	0.79	(0.52,	1.20)	0.63	(0.43,	0.91)	0.01
Fully adjusted ^b	146	1618	0.83	(0.67,	1.02)	Ref	0.89	(0.51,	1.55)	0.60	(0.36,	1.01)	0.05

ACS American Cancer Society, Adj adjusted, AICR American Institute for Cancer Research, CI confidence interval, HR hazard ratio, pts points, WCRF World Cancer Research Fund, wk week.

0.091) nor the ACS Score were associated with risk of prostate cancer progression (HR_{cont}: 0.93; 95% Cl: 0.82–1.05; HR_{2 vs 1}: 1.19, 95% Cl: 0.82–1.71; HR_{3 vs 1}: 0.83; 95% Cl: 0.58–1.18; $P_{\rm trend}=0.30$). The ACS Score + Alcohol, however, demonstrated evidence of an inverse association with risk of progression (HR_{cont}: 0.89, 95% Cl: 0.81–0.98; HR_{2 vs 1}: 0.97, 95% Cl: 0.71–1.32; HR_{3 vs 1}: 0.48, 95% Cl: 0.28–0.82; $P_{\rm trend}=0.007$). The WCRF/AICR Score was also inversely associated with risk of prostate cancer progression (HR_{cont}: 0.83, 95% Cl: 0.67–1.02; HR_{2 vs 1}: 0.89, 95% Cl; 0.51–1.55; HR_{3 vs 1}: 0.60, 95% Cl: 0.36–1.01; $P_{\rm trend}=0.05$) (Table 2 and Fig. 1).

Prostate cancer-specific mortality

The 2021 Score was statistically significantly associated with a lower risk of PCSM (HR_{cont}: 0.62, 95% CI: 0.45–0.85). However, in models assessing score tertiles, there was no clear association with

PCSM (HR $_{2 \text{ vs}}$ 1: 0.50, 95% CI: 0.23–1.06; HR $_{3 \text{ vs}}$ 1: 0.71, 95% CI: 0.38-1.33; $P_{\text{trend}} =$ 0.28). When dietary factors were included (2021 Score + Diet), associations with PCSM were statistically significant in continuous (HR: 0.65; 95% CI: 0.45, 0.93) and tertile models, showing a 59% reduced risk of PCSM among those with the highest versus the lowest tertile of score (HR $_{3 \text{ vs}}$ 1: 0.41, 95% CI: 0.20, 0.85; $P_{\text{trend}} =$ 0.02) (Table 3 and Fig. 1).

There was no association with PCSM for the 2015 (HR_{cont} : 0.81, 95% CI: 0.63–1.04), ACS (HR_{cont} : 0.82, 95% CI: 0.65–1.04) or ACS + Alcohol (HR_{cont} : 0.92, 95% CI: 0.74–1.15) Scores (Table 3 and Fig. 1). The WCRF/AICR Score was inversely associated with risk of PCSM (HR_{cont} : 0.71, 95% CI: 0.57–0.89), amounting to a 48% (HR: 0.52, 95% CI: 0.33–0.81; $P_{trend}=0.004$) lower risk among those with the highest versus lowest tertile of score (Table 3 and Fig. 1).

^aSimple models were adjusted for time between diagnosis and date of first CDL questionnaire (continuous), age at diagnosis (continuous) and CaPSURE clinical site.

^bFully adjusted models were additionally adjusted for clinical T-stage (T1, T2, T3), Gleason score (<7, 7, >7), diagnostic PSA value (≤6, >6–10, >10–20), primary treatment (radical prostatectomy, active surveillance/watchful waiting, radiotherapy/brachytherapy, hormone therapy, other), family history of prostate cancer in brother of father (yes, no), race (white, non-white), total caloric intake (continuous), plus the following variables (if not part of the score): whole milk intake (≤4 servings/wk, >4 servings/wk), wine intake (3–14 servings/wk, <3 or >14 servings/wk), total alcohol intake (non-drinker, >0–2 serving/day, >2 servings/day), red and processed meat intake (quartiles), tomato intake (continuous), dark meat fish intake (continuous), selenium supplement use (non-user, <140 μg/day, ≥140 μg/day, user with unknown daily dosage), smoking (never, quit ≥10 years prior, quit <10 years prior, current).

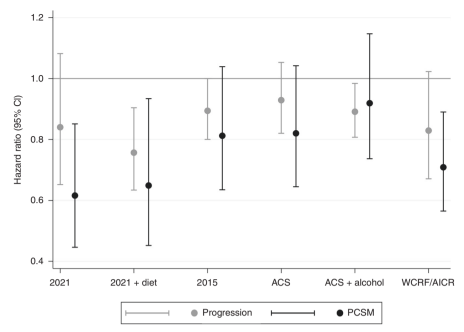


Fig. 1 Post-diagnostic health behaviour scores and the risk of prostate cancer progression and mortality among men with non-metastatic prostate cancer. Visual summary of the HR and 95% CI for the risk of progression (light grey) and PCSM (dark grey) per 1-unit increase (i.e, healthier score) in each of the 6 behavioural scores examined.

Sensitivity analyses

Across all scores, there was no evidence that men with higher behavioural scores presented more frequently for PSA monitoring following a diagnosis (data not shown). In models that imposed date of diagnosis as time zero, the trends were similar across all scores (Supplemental Table S3). The results from Cox proportional hazards models for progression were similar to those from the parametric (Weibull) survival and there was no evidence that competing events impacted the results (Supplemental Table S4). Multiple imputation resulted in 2056 complete records and retention of all 192 events in multivariable models. Across all scores, with the exception of age, characteristics were similar between men with and without missingness, providing some evidence that data were missing at random (Supplemental Table S5). The results following imputation were similar to those obtained from the complete-case analysis. With the larger sample sizes, however, the confidence intervals tightened, resulting in statistically significant estimates across all scores (Supplemental Table S6).

DISCUSSION

In this prospective study, we examined associations of behavioural risk scores with prostate cancer progression and PCSM among men diagnosed with non-metastatic prostate cancer. For each 1-unit increase (i.e., healthier) in the 2021 Score + Diet and the ACS Score + Alcohol, men had a statistically significant 24% and 11% lower risk of progression, respectively. The WCRF/AICR Score was also associated with a (statistically non-significant) reduced risk of progression, demonstrating a 17% lower risk of progression per point increase. Men in the highest tertile of the 2021 Score + Diet and WCRF/AICR Score had a 59% and 48%, respectively, lower risk of dying from prostate cancer compared to those in the lowest tertile.

The difference in associations observed between the two outcomes may reflect different mechanisms driving recurrence versus PCSM. Indeed, 94% of men with biochemical recurrence in this cohort did not die from prostate cancer during study follow-up. Another explanation is confounding: healthier men may

present more often for PSA monitoring and thus be more likely to have biochemical recurrence detected which may spuriously attenuate associations. We attempted to evaluate whether such confounding bias impacted our results and did not observe different screening behaviours based on score levels. Nevertheless, we cannot rule out confounding.

Importantly, components varied across behavioural scores and were used differently within scores. For example, the ACS Score + Alcohol assigned the highest (i.e., healthiest) points for moderate alcohol intake, whereas WCRF/AICR Score preserved highest points for no alcohol intake. The 2021 Score + Diet only included moderate consumption of wine in its highest point. Aligned with ACS recommendations, the decision to consume alcohol should be made on an individual basis with a patient's provider [70].

The 2015 Score was developed based on the literature describing the risk of developing lethal prostate cancer among disease-free men; [10] our team previously reported that this score was associated with a 68% lower risk (5–6 points vs. 0–1 points) of developing lethal prostate cancer among disease-free men [10]. However, our results, in combination with existing evidence, suggest that behavioural factors associated with *developing* prostate cancer may differ from those associated with *progression* and mortality following a diagnosis [71, 72].

While not all statistically significantly, all scores were *inversely* associated with both outcomes, supporting our hypothesis that higher scores/healthier lifestyle patterns would be protective for progression and PCSM. Considering similarities and differences in score composition may provide insights for further scientific exploration. For example, while both the 2021 Score + Diet and WCRF/AICR Score had statistically significant inverse associations with PCSM, the 2021 Score + Diet was associated with a slightly larger reduction in risk. This suggests that while general "healthy" diet recommendations for total cancer prevention (e.g., WCRF/AIRC) are good, additional specific guidance (e.g., limited intake of whole milk) for preventing *prostate cancer* death may be warranted. Also, the 2021 Score + Diet was the only score to consider saturated fat intake in addition to specific high-fat foods. This information may guide hypothesis

Table 3. Post-diagnostic health behaviour scores and the risk of prostate cancer mortality among men with non-metastatic prostate cancer, estimated via Cox proportional hazards models^a.

2021 Score—HR	(95% CI)												
	Events	Ν	Contin	iuous		1st Tertile	2nd Te	ertile		3rd Te	rtile		P_{trend}
			7 0.62 (0.45, 0.8 CI) Continuous 9 0.76 (0.55, 1.0 7 0.65 (0.45, 0.9 Continuous 6 0.86 (0.71, 1.0 4 0.81 (0.63, 1.0 Continuous 6 0.84 (0.69, 1.0 5 0.82 (0.65, 1.0 6 CI) Continuous 9 0.87 (0.74, 1.0 Continuous 9 0.87 (0.74, 1.0 Continuous			(0-2 pts)	(2.5 pt	ts)		(3 pts))		
Simple ^b	69	2369	0.73	(0.52,	1.03)	Ref	0.64	(0.39,	1.04)	0.73	(0.40,	1.35)	0.32
Fully adjusted ^c	51	1877	0.62	(0.45,	0.85)	Ref	0.50	(0.23,	1.06)	0.71	(0.38,	1.33)	0.28
2021 Score with	Diet—HR	(95% CI)											
	Events	N	Contin	iuous		1st Tertile	2nd Te	ertile		3rd Te	rtile		P_{trend}
						(0.25-2.25 pts)	(2.5-3	pts)		(3.25-	4 pts)		
Simple ^b	69	2369	0.76	(0.55,	1.04)	Ref	0.64	(0.37,	1.09)	0.59	(0.33,	1.06)	0.078
Fully adjusted ^c	52	1947	0.65	(0.45,	0.93)	Ref	0.55	(0.30,	1.02)	0.41	(0.20,	0.85)	0.02
2015 Score—HR	(95% CI)												
	Events	N	Contin	iuous		1st Tertile	2nd Te	ertile		3rd Te	rtile		P_{trend}
						(0-3 pts)	(4 pts)			(5–6 p	ts)		
Simple ^b	65	2336	0.86	(0.71,	1.04)	Ref	0.96	(0.64,	1.43)	0.72	(0.33,	1.57)	0.41
Fully adjusted ^c	49	1874	0.81	(0.63,	1.04)	Ref	0.94	(0.65,	1.37)	0.61	(0.18,	2.03)	0.42
ACS Score—HR	(95% CI)												
	Events	N	Contin	iuous		1st Tertile	2nd Te	ertile		3rd Te	rtile		$P_{\rm trend}$
			Continuous Continuous			(0-2 pts)	(2.5-3	pts)		(3.5–6	pts)		
Simple ^b	69	2396	0.84	(0.69,	1.02)	Ref	0.74	(0.42,	1.32)	0.64	(0.35,	1.15)	0.13
Fully adjusted ^c	50	1875	0.82	(0.65,	1.04)	Ref	0.65	(0.36,	1.17)	0.58	(0.29,	1.15)	0.12
ACS Score with	alcohol—H	R (95% C	:1)										
	Events	N	Contin	iuous		1st Tertile	2nd Tertile			3rd Te	rtile		$P_{\rm trend}$
			Continuous			(0-3.5 pts)	(4–5 pts)			(5.5–8			
Simple ^b	68	2359	0.87	(0.74,	1.03)	Ref	0.74	(0.41,	1.33)	0.73	(0.37,	1.42)	0.35
Fully adjusted ^c	50	1875	0.92	(0.74,	1.15)	Ref	1.07	(0.49,	2.37)	0.82	(0.35,	1.93)	0.64
WCRF/AICR Scor	e—HR (95	% CI)											
	Events	Ν	Contin	uous		1st Tertile	2nd Te	ertile		3rd Te	P_{trend}		
						(0.75-3.25 pts)	(3.5–4	pts)		(4.25–	7 pts)		
Simple ^b	69	2395	0.80	(0.69,	0.94)	Ref	0.66	(0.37,	1.17)	0.57	(0.38,	0.84)	0.004
Fully adjusted ^c	50	1883	0.71	(0.57,	0.89)	Ref	0.51	(0.25,	1.03)	0.52	(0.33,	0.81)	0.004

ACS American Cancer Society, Adj adjusted, AICR American Institute for Cancer Research, CI confidence interval, HR hazard ratio, pts points, WCRF World Cancer Research Fund, wk week.

generation regarding biological mechanisms linking health behaviours to PC outcomes. Confirmation of the performance of the 2021 Score + Diet in independent study populations is warranted, and could lead to more tailored recommendations for patients with prostate cancer.

There are several limitations of our analyses to consider. Men in our study predominately identified as white race (95%), were well educated (77% with at least some college), and were insured (97%), meaning these results are not generalisable to all men with prostate cancer. Social determinants of health and their impacts on health and disease status cannot be addressed in this cohort. While we observed some statistically significant inverse associations for PCSM, this was a secondary outcome given the limited number of events. Though we made efforts to address potential

biases in this study (e.g., multiple imputation to address missingness, modelling PSA surveillance behaviour as a function of behavioural scores to address confounding issues), these approaches are not without their own assumptions, and thus we cannot rule out bias entirely. Finally, the post-diagnostic literature that drove the creation of the 2021 Score (with and without diet) came from a limited number of study populations, which included CaPSURE [6]. This fact underscores the importance of confirming these findings in other populations.

In conclusion, among men diagnosed with non-metastatic prostate cancer, a behavioural score developed based on the current post-diagnostic literature (2021 Score Including Diet) was associated with a 24% lower risk of progression and 35% lower risk of PCSM per one-unit increase in the score. Men diagnosed

^aA total of 2447 men met inclusion for PCSM analyses, as men who were excluded from progression analysis due to having a documented progression event prior to questionnaire were included in this analysis.

bSimple models were adjusted for time between diagnosis and date of first CDL questionnaire (continuous) and age at diagnosis (continuous).

^cFully adjusted models were additionally adjusted for clinical T-stage (T1, T2, T3), Gleason score (<7, 7, >7), diagnostic PSA value (≤6, >6–10, >10–20), primary treatment (radical prostatectomy, active surveillance/watchful waiting, radiotherapy/brachytherapy, hormone therapy, other), family history of prostate cancer in brother of father (yes, no), race (white, non-white), total caloric intake (continuous), plus the following variables (if not part of the score): whole milk intake (≤4 servings/wk, >4 servings/wk), wine intake (3–14 servings/wk), <3 or >14 servings/wk), total alcohol intake (non-drinker, >0–2 serving/day, >2 servings/day), red and processed meat intake (quartiles), tomato intake (continuous), dark meat fish intake (continuous), selenium supplement use (non-user, <140 µg/day, ≥140 µg/day, user with unknown daily dosage), smoking (never, quit ≥10 years prior, quit <10 years prior, current).

with non-metastatic prostate cancer may improve survivorship by adhering to post-diagnostic prostate cancer-specific dietary recommendations—avoiding/limiting the consumption of whole milk, red and processed meats, and saturated fat, while allowing moderate consumption of wine—in addition to the general recommendations to avoid smoking, maintain a healthy body size, and engage in regular physical activity.

DATA AVAILABILITY

Data and code can be made available on request.

REFERENCES

- Siegel RL, Miller KD, Fuchs HE, Jemal A. Cancer statistics, 2021. CA Cancer J Clin. 2021;71:7–33.
- American Cancer Society, Cancer Facts and Figures 2021. https:// www.cancer.org/content/dam/cancer-org/research/cancer-facts-and-statistics/ annual-cancer-facts-and-figures/2021/cancer-facts-and-figures-2021.pdf. Accessed 09/07/2021.
- American Cancer Society, Cancer Facts and Figures 2019-2021. 2019. https:// www.cancer.org/content/dam/cancer-org/research/cancer-facts-and-statistics/ cancer-treatment-and-survivorship-facts-and-figures/cancer-treatment-andsurvivorship-facts-and-figures-2019-2021.pdf. Accessed 01/28/2022.
- Negoita S, Feuer EJ, Mariotto A, Cronin KA, Petkov VI, Hussey SK, et al. Annual report to the nation on the status of cancer, part II: recent changes in prostate cancer trends and disease characteristics. Cancer. 2018;124:2801–14.
- Torre LA, Siegel RL, Ward EM, Jemal A. Global cancer incidence and mortality rates and trends-an update. Cancer Epidemiol Biomark Prev. 2016;25:16–27.
- Langlais CS, Graff RE, Van Blarigan EL, Palmer NR, Washington SL 3rd, Chan JM, et al. Post-diagnostic dietary and lifestyle factors and prostate cancer recurrence, progression, and mortality. Curr Oncol Rep. 2021;23:37.
- 7. Hu FB, Willett WC. Current and future landscape of nutritional epidemiologic research. J Am Med Assoc. 2018;320:2073–4.
- 8. Hu FB. Dietary pattern analysis: a new direction in nutritional epidemiology. Curr Opin Lipido. 2002;13:3–9.
- Tat D, Kenfield SA, Cowan JE, Broering JM, Carroll PR, Van Blarigan EL, et al. Milk and other dairy foods in relation to prostate cancer recurrence: data from the cancer of the prostate strategic urologic research endeavor (CaPSURE). Prostate. 2018;78:32–9.
- Kenfield SA, Batista JL, Jahn JL, Downer MK, Van Blarigan EL, Sesso HD, et al. Development and application of a lifestyle score for prevention of lethal prostate cancer. J Natl Cancer Inst. 2016;108:div329.
- Shams-White MM, Brockton NT, Mitrou P, Romaguera D, Brown S, Bender A, et al.
 Operationalizing the 2018 World Cancer Research Fund/American Institute for Cancer Research (WCRF/AICR) cancer prevention recommendations: a standar-dized Scoring System. Nutrients. 2019;11:1572.
- Shams-White MM, Romaguera D, Mitrou P, Reedy J, Bender A, Brockton NT. Further guidance in implementing the standardized 2018 World Cancer Research Fund/American Institute for Cancer Research (WCRF/AICR) Score. Cancer Epidemiol Biomark Prev. 2020;29:889–94.
- McCullough ML, Patel AV, Kushi LH, Patel R, Willett WC, Doyle C, et al. Following cancer prevention guidelines reduces risk of cancer, cardiovascular disease, and all-cause mortality. Cancer Epidemiol Biomark Prev. 2011;20:1089–97.
- Van Blarigan EL, Fuchs CS, Niedzwiecki D, Zhang S, Saltz LB, Mayer RJ, et al. Association of survival with adherence to the american cancer society nutrition and physical activity guidelines for cancer survivors after colon cancer diagnosis: the CALGB 89803/Alliance trial. JAMA Oncol. 2018;4:783–90.
- Lubeck DP, Litwin MS, Henning JM, Stier DM, Mazonson P, Fisk R, et al. The CaPSURE database: a methodology for clinical practice and research in prostate cancer. CaPSURE Research Panel. Cancer of the Prostate Strategic Urologic Research Endeavor. Urology. 1996;48:773–7.
- Willett W. Nutritional epidemiology. Vol. 40. third edn. New York, USA: Oxford University Press; 2013.
- Feskanich D, Rimm EB, Giovannucci EL, Colditz GA, Stampfer MJ, Litin LB, et al. Reproducibility and validity of food intake measurements from a semiquantitative food frequency questionnaire. J Am Diet Assoc. 1993;93:790–6.
- Hu FB, Rimm E, Smith-Warner SA, Feskanich D, Stampfer MJ, Ascherio A, et al. Reproducibility and validity of dietary patterns assessed with a food-frequency questionnaire. Am J Clin Nutr. 1999;69:243–9.
- Rimm EB, Giovannucci EL, Stampfer MJ, Colditz GA, Litin LB, Willett WC. Reproducibility and validity of an expanded self-administered semiquantitative food frequency questionnaire among male health professionals. Am J Epidemiol. 1992;135:1114–26.

- U.S. Department of Agriculture, Agricultural Research Service. FoodData Central, 2019. https://fdc.nal.usda.gov/. Accessed January 12, 2022.
- Chasan-Taber S, Rimm EB, Stampfer MJ, Spiegelman D, Colditz GA, Giovannucci E, et al. Reproducibility and validity of a self-administered physical activity questionnaire for male health professionals. Epidemiology. 1996;7:81–6.
- Gong Z, Agalliu I, Lin DW, Stanford JL, Kristal AR. Cigarette smoking and prostate cancer-specific mortality following diagnosis in middle-aged men. Cancer Causes Control. 2008;19:25–31.
- 23. Oefelein MG, Resnick MI. Association of tobacco use with hormone refractory disease and survival of patients with prostate cancer. J Urol. 2004;1711:2281–4.
- Pantarotto J, Malone S, Dahrouge S, Gallant V, Eapen L. Smoking is associated with worse outcomes in patients with prostate cancer treated by radical radiotherapy. BJU Int. 2007;99:564–9.
- Moreira DM, Aronson WJ, Terris MK, Kane CJ, Amling CL, Cooperberg MR, et al. Cigarette smoking is associated with an increased risk of biochemical disease recurrence, metastasis, castration-resistant prostate cancer, and mortality after radical prostatectomy: results from the SEARCH database. Cancer. 2014;120:197–204.
- Pickles T, Liu M, Berthelet E, Kim-Sing C, Kwan W, Tyldesley S. The effect of smoking on outcome following external radiation for localized prostate cancer. J Urol. 2004;171:1543–6.
- Joshu CE, Mondul AM, Meinhold CL, Humphreys EB, Han M, Walsh PC, et al. Cigarette smoking and prostate cancer recurrence after prostatectomy. J Natl Cancer Inst. 2011;103:835–8.
- Oh JJ, Hong SK, Jeong CW, Byun SS, Lee SE. Significance of smoking status regarding outcomes after radical prostatectomy. Int Urol Nephrol 2012;44:119–24.
- 29. Ngo TC, Lee JJ, Brooks JD, Nolley R, Ferrari M, Presti JC Jr. Smoking and adverse outcomes at radical prostatectomy. Urol Oncol. 2013;31:749–54.
- Rieken M, Shariat SF, Kluth LA, Fajkovic H, Rink M, Karakiewicz PI, et al. Association of cigarette smoking and smoking cessation with biochemical recurrence of prostate cancer in patients treated with radical prostatectomy. Eur Urol. 2015;68:949–56.
- Steinberger E, Kollmeier M, McBride S, Novak C, Pei X, Zelefsky MJ. Cigarette smoking during external beam radiation therapy for prostate cancer is associated with an increased risk of prostate cancer-specific mortality and treatment-related toxicity. BJU Int. 2015;116:596–603.
- Negrón R, Vásquez A, Nieves M, Guerrios L, Irizarry-Ramírez M. Body mass index affects the diagnosis and progression of prostate cancer in Hispanics. Ethn Dis. 2010;20:S1-168–172.
- 33. Keto CJ, Aronson WJ, Terris MK, Presti JC, Kane CJ, Amling CL, et al. Obesity is associated with castration-resistant disease and metastasis in men treated with androgen deprivation therapy after radical prostatectomy: results from the SEARCH database. BJU Int. 2012;110:492–8.
- Bhindi B, Kulkarni GS, Finelli A, Alibhai SM, Hamilton RJ, Toi A, et al. Obesity is associated with risk of progression for low-risk prostate cancers managed expectantly. Eur Urol. 2014;66:841–8.
- Efstathiou JA, Chen MH, Renshaw AA, Loffredo MJ, D'Amico AV. Influence of body
 mass index on prostate-specific antigen failure after androgen suppression and
 radiation therapy for localized prostate cancer. Cancer. 2007;109:1493–8.
- Spangler E, Zeigler-Johnson CM, Coomes M, Malkowicz SB, Wein A, Rebbeck TR.
 Association of obesity with tumor characteristics and treatment failure of prostate cancer in African-American and European American men. J Urol. 2007;178:1939–44.
- 37. Stroup SP, Cullen J, Auge BK, L'Esperance JO, Kang SK. Effect of obesity on prostate-specific antigen recurrence after radiation therapy for localized prostate cancer as measured by the 2006 Radiation Therapy Oncology Group-American Society for Therapeutic Radiation and Oncology (RTOG-ASTRO) Phoenix consensus definition. Cancer. 2007;110:1003–9.
- Hisasue S, Yanase M, Shindo T, Iwaki H, Fukuta F, Nishida S, et al. Influence of body mass index and total testosterone level on biochemical recurrence following radical prostatectomy. Jpn J Clin Oncol. 2008;38:129–33.
- Magheli A, Rais-Bahrami S, Trock BJ, Humphreys EB, Partin AW, Han M, et al. Impact of body mass index on biochemical recurrence rates after radical prostatectomy: an analysis utilizing propensity score matching. Urology. 2008;72:1246–51.
- Freedland SJ, Isaacs WB, Mangold LA, Yiu SK, Grubb KA, Partin AW, et al. Stronger association between obesity and biochemical progression after radical prostatectomy among men treated in the last 10 years. Clin Cancer Res. 2005;11:2883–8.
- Freedland SJ, Grubb KA, Yiu SK, Humphreys EB, Nielsen ME, Mangold LA, et al. Obesity and risk of biochemical progression following radical prostatectomy at a tertiary care referral center. J Urol. 2005;174:919–22.
- 42. Freedland SJ, Sun L, Kane CJ, Presti JC Jr, Terris MK, Amling CL, et al. Obesity and oncological outcome after radical prostatectomy: impact of prostate-specific antigen-based prostate cancer screening: results from the Shared Equal Access

- Regional Cancer Hospital and Duke Prostate Center databases. BJU Int. 2008:102:969–74.
- Strom SS, Wang X, Pettaway CA, Logothetis CJ, Yamamura Y, Do KA, et al. Obesity, weight gain, and risk of biochemical failure among prostate cancer patients following prostatectomy. Clin Cancer Res. 2005;11:6889–94.
- Strom SS, Kamat AM, Gruschkus SK, Gu Y, Wen S, Cheung MR, et al. Influence of obesity on biochemical and clinical failure after external-beam radiotherapy for localized prostate cancer. Cancer. 2006;107:631–9.
- Kane CJ, Im R, Amling CL, Presti JC Jr, Aronson WJ, Terris MK, et al. Outcomes after radical prostatectomy among men who are candidates for active surveillance: results from the SEARCH database. Urology. 2010;76:695–700.
- Ho T, Gerber L, Aronson WJ, Terris MK, Presti JC, Kane CJ, et al. Obesity, prostatespecific antigen nadir, and biochemical recurrence after radical prostatectomy: biology or technique? Results from the SEARCH database. Eur Urol. 2012;62:910–6.
- Wang LS, Murphy CT, Ruth K, Zaorsky NG, Smaldone MC, Sobczak ML, et al. Impact of obesity on outcomes after definitive dose-escalated intensity-modulated radiotherapy for localized prostate cancer. Cancer. 2015;121:3010–7.
- 48. Langlais CS, Cowan JE, Neuhaus J, Kenfield SA, Van Blarigan EL, Broering JM, et al. Obesity at diagnosis and prostate cancer prognosis and recurrence risk following primary treatment by radical prostatectomy. Cancer Epidemiol Biomark Prev. 2019;28:1917–25.
- Bonn SE, Sjölander A, Lagerros YT, Wiklund F, Stattin P, Holmberg E, et al. Physical activity and survival among men diagnosed with prostate cancer. Cancer Epidemiol Biomark Prev. 2015;24:57–64.
- Wang Y, Jacobs EJ, Gapstur SM, Maliniak ML, Gansler T, McCullough ML, et al. Recreational physical activity in relation to prostate cancer-specific mortality among men with nonmetastatic prostate cancer. Eur Urol. 2017;72:931–9.
- 51. Kenfield SA, Stampfer MJ, Giovannucci E, Chan JM. Physical activity and survival after prostate cancer diagnosis in the health professionals follow-up study. J Clin Oncol. 2011;29:726–32.
- Friedenreich CM, Wang Q, Neilson HK, Kopciuk KA, McGregor SE, Courneya KS. Physical activity and survival after prostate cancer. Eur Urol. 2016;70:576–85.
- Richman EL, Kenfield SA, Stampfer MJ, Paciorek A, Carroll PR, Chan JM. Physical activity after diagnosis and risk of prostate cancer progression: data from the cancer of the prostate strategic urologic research endeavor. Cancer Res. 2011;71:3889–95.
- 54. Meyer F, Bairati I, Shadmani R, Fradet Y, Moore L. Dietary fat and prostate cancer survival. Cancer Causes Control. 1999;10:245–51.
- Epstein MM, Kasperzyk JL, Mucci LA, Giovannucci E, Price A, Wolk A, et al. Dietary fatty acid intake and prostate cancer survival in Örebro County, Sweden. Am J Epidemiol. 2012;176:240–52.
- Van Blarigan EL, Kenfield SA, Yang M, Sesso HD, Ma J, Stampfer MJ, et al. Fat intake after prostate cancer diagnosis and mortality in the Physicians' Health Study. Cancer Causes Control. 2015;26:1117–26.
- Pettersson A, Kasperzyk JL, Kenfield SA, Richman EL, Chan JM, Willett WC, et al. Milk and dairy consumption among men with prostate cancer and risk of metastases and prostate cancer death. Cancer Epidemiol Biomark Prev. 2012;21:428–36.
- 58. Tat D, Kenfield SA, Cowan JE, Broering JM, Carroll PR, Van Blarigan EL, et al. Milk and other dairy foods in relation to prostate cancer recurrence: data from the cancer of the prostate strategic urologic research endeavor (CaPSURE™). Prostate. 2018;78:32–9.
- Farris MS, Courneya KS, Kopciuk KA, McGregor SE, Friedenreich CM. Postdiagnosis alcohol intake and prostate cancer survival: a population-based cohort study. Int J Cancer. 2018;143:253–62.
- Downer MK, Kenfield SA, Stampfer MJ, Wilson KM, Dickerman BA, Giovannucci EL, et al. Alcohol intake and risk of lethal prostate cancer in the health professionals follow-up study. J Clin Oncol. 2019;37:1499–511.
- 61. Richman EL, Kenfield SA, Stampfer MJ, Giovannucci EL, Chan JM. Egg, red meat, and poultry intake and risk of lethal prostate cancer in the prostate-specific antigen-era: incidence and survival. Cancer Prev Res. 2011;4:2110–21.
- Yang M, Kenfield SA, Van Blarigan EL, Batista JL, Sesso HD, Ma J, et al. Dietary patterns after prostate cancer diagnosis in relation to disease-specific and total mortality. Cancer Prev Res. 2015:8:545–51.
- Langlais CS, Chan JM, Kenfield SA, Cowan JE, Graff RE, Broering JM, et al. Postdiagnostic coffee and tea consumption and risk of prostate cancer progression by smoking history. Cancer Causes Control. 2021;32:635–44.
- National Death Index. National Center for Health Statistics, Centers for Disease Control and Prevention, Department of Health and Human Services; 2021. https://www.cdc.gov/nchs/ndi/resources.htm. Accessed January 13, 2022.
- 65. Klein JP, Moeschberger ML. Survival analysis: techniques for censored and truncated data. Berlin/Heidelberg, Germany: Springer; 2003.
- Greenland S, Pearl J, Robins JM. Causal diagrams for epidemiologic research. Epidemiology. 1999;10:37–48.
- Howards PP, Schisterman EF, Poole C, Kaufman JS, Weinberg CR. "Toward a clearer definition of confounding" revisited with directed acyclic graphs. Am J Epidemiol. 2012;176:506–11.

- Harel O, Mitchell EM, Perkins NJ, Cole SR, Tchetgen Tchetgen EJ, Sun B, et al. Multiple imputation for incomplete data in epidemiologic studies. Am J Epidemiol. 2018;187:576–84.
- 69. Rubin DB. Inference and missing data. Biometrika. 1976;63:581-92.
- Rock CL, Doyle C, Demark-Wahnefried W, Meyerhardt J, Courneya KS, Schwartz AL, et al. Nutrition and physical activity guidelines for cancer survivors. CA Cancer J Clin. 2012;62:243–74.
- 71. Giovannucci E, Liu Y, Platz EA, Stampfer MJ, Willett WC. Risk factors for prostate cancer incidence and progression in the health professionals follow-up study. Int J Cancer. 2007;121:1571–8.
- Kensler KH, Rebbeck TR. Cancer progress and priorities: prostate cancer. Cancer Epidemiol Biomark Prev. 2020;29:267–77.

ACKNOWLEDGEMENTS

The authors wish to thank M. Maria Glymour for her contributions to the analysis plan and the CaPSURE participants for making this research possible.

AUTHOR CONTRIBUTIONS

CSL acquired funding to support the work, conceptualised and performed analyses, interpreted the results, and drafted the first and subsequent drafts of the manuscript and tables. REG conceptualised analyses, interpreted the results, and commented on the manuscript. ELVB conceptualised analyses, interpreted the results, and commented on the manuscript. JMN conceptualised analyses, interpreted the results, and commented on the manuscript. JEC curated data and commented on the manuscript. PC acquired CaPSURE funding and commented on the manuscript. SAK conceptualised analyses, supervised research conduct, interpreted results, and commented on the manuscript. JMC conceptualised analyses, supervised research conduct, interpreted results and commented on the manuscript.

FUNDING

Research reported in this publication was supported by the University of California, San Francisco Prostate Cancer Program, the National Cancer Institute of the National Institutes of Health: F31CA247093 (CSL) and K07CA197077 (EVB), and the UCSF Goldberg-Benioff Program in Translational Cancer Biology. CaPSURE was funded by the United States Department of Defense Prostate Cancer Research Program (W81XWH-13-2-0074 and W81XWH-04-1-0850). JMC is the Steven & Christine Burd-Safeway Distinguished Professor. SAK is the Helen Diller Family Chair in Population Science for Urologic Cancer. REG is supported by a Young Investigator Award from the Prostate Cancer Foundation.

COMPETING INTERESTS

The authors declare no competing interests.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

All participants provided written informed consent and the study was conducted in accordance with the Belmont Report and the U.S. Common Rule. The study was approved by the Institutional Review Board of the University of California, San Francisco, and those of participating sites, as required.

CONSENT FOR PUBLICATION

Not applicable.

ADDITIONAL INFORMATION

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41416-023-02283-1.

Correspondence and requests for materials should be addressed to Crystal S. Langlais.

Reprints and permission information is available at http://www.nature.com/

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as 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. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's

Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2023