INTRODUCTION

Despite recent advances in the treatment strategies used for the clinical management of metastatic prostate cancer (PCa), disease survival remains lower than 30%; notably, PCa is the second leading cause of cancer mortality in American males. Notably, men of African descent in the United States and Caribbean have the highest PCa mortality rates compared to men with European ancestry. Although current therapeutics are quite potent and effective, disease resistance, progression to metastasis, therapy-associated toxicities and efficacy-related issues in diverse populations develop over time. Thus, non-toxic and efficacious therapeutic strategies are needed to address these major obstacles for the clinical treatment and management of PCa. In this regard, preclinical and population-based efficacy studies have shown the potential of natural non-toxic nutraceuticals as potent anti-PCa agents. Accordingly, the implementation of nutraceutical intervention and genetic testing in diverse populations might aid in the development and design of precision medicine strategies to reduce the burden of chemotherapy-associated toxicities, suppress disease resistance, and treat both localized and advanced PCa. Consequently, additional large-scale and inclusive clinical studies are required to fully assess efficacy and therapeutic limitations of these agents in PCa. This review discusses the most current clinical research on selected nutraceutical agents and their efficacy in the context of clinico-pathological outcomes and disease susceptibility in diverse PCa clinical and epidemiological studies.

npj Precision Oncology (2018) 2:15; doi:10.1038/s41698-018-0058-x

Prostate cancer (PCa) is the most frequently diagnosed malignancy and second leading cause of cancer mortality in American males. Notably, men of African descent in the United States and Caribbean have the highest PCa mortality rates compared to men with European ancestry. Although current therapeutics are quite potent and effective, disease resistance, progression to metastasis, therapy-associated toxicities and efficacy-related issues in diverse populations develop over time. Thus, non-toxic and efficacious therapeutic strategies are needed to address these major obstacles for the clinical treatment and management of PCa. In this regard, preclinical and population-based efficacy studies have shown the potential of natural non-toxic nutraceuticals as potent anti-PCa agents. Accordingly, the implementation of nutraceutical intervention and genetic testing in diverse populations might aid in the development and design of precision medicine strategies to reduce the burden of chemotherapy-associated toxicities, suppress disease resistance, and treat both localized and advanced PCa. Consequently, additional large-scale and inclusive clinical studies are required to fully assess efficacy and therapeutic limitations of these agents in PCa. This review discusses the most current clinical research on selected nutraceutical agents and their efficacy in the context of clinico-pathological outcomes and disease susceptibility in diverse PCa clinical and epidemiological studies.

Review Article

Nutraceuticals in prostate cancer therapeutic strategies and their neo-adjuvant use in diverse populations

Dominique Reed1, Komal Raina1,2 and Rajesh Agarwal1,2

Prostate cancer (PCa) is the most frequently diagnosed malignancy and second leading cause of cancer mortality in American males. Notably, men of African descent in the United States and Caribbean have the highest PCa mortality rates compared to men with European ancestry. Although current therapeutics are quite potent and effective, disease resistance, progression to metastasis, therapy-associated toxicities and efficacy-related issues in diverse populations develop over time. Thus, non-toxic and efficacious therapeutic strategies are needed to address these major obstacles for the clinical treatment and management of PCa. In this regard, preclinical and population-based efficacy studies have shown the potential of natural non-toxic nutraceuticals as potent anti-PCa agents. Accordingly, the implementation of nutraceutical intervention and genetic testing in diverse populations might aid in the development and design of precision medicine strategies to reduce the burden of chemotherapy-associated toxicities, suppress disease resistance, and treat both localized and advanced PCa. Consequently, additional large-scale and inclusive clinical studies are required to fully assess efficacy and therapeutic limitations of these agents in PCa. This review discusses the most current clinical research on selected nutraceutical agents and their efficacy in the context of clinico-pathological outcomes and disease susceptibility in diverse PCa clinical and epidemiological studies.

npj Precision Oncology (2018) 2:15; doi:10.1038/s41698-018-0058-x

INTRODUCTION

Despite recent advances in the treatment strategies used for the clinical management of metastatic prostate cancer (PCa), disease survival remains lower than 30%; notably, PCa is the second leading cause of cancer mortality in American men.1 Primary interventions for PCa include surgery, adjuvant chemotherapy, hormonal therapy (i.e., androgen deprivation), and/or radiation for advanced disease. Although these therapeutic options are quite potent and effective, disease resistance, progression to metastasis stage, therapy-associated toxicities and efficacy-related issues in diverse populations have developed over time. Thus, efficacious and non-toxic therapeutic strategies are needed to address these major obstacles for the clinical treatment and management of PCa. Past and ongoing chemoprevention/intervention research has recognized the use of nutraceutical agents as a feasible and non-toxic option, which could protect against tumorigenesis as well as enhance the therapeutic response of pre-existing anti-cancer treatments.2

Precision medicine using a nutraceutical approach is one of the key paradigms for the conceptualization and development of non-toxic therapeutic strategies that act synergistically with existing clinical anti-cancer agents. Precision medicine strategies navigate the intricate interplay between genetic, racial, and socio-economic factors, and involve interventions that compensate for individual variations in pharmacogenomic responses to cancer therapies and major PCa risk factors (i.e., age, family history, race, and genetic susceptibility) shown in Fig. 1. Genetic dispositions are critical for identifying individuals at higher risk for disease development or advanced disease and predicting therapeutic response to certain anti-cancer agents. Genetic variations have also shown protective and negative effects on PCa susceptibility. Thus, the efficacy of nutraceutical interventions is heavily influenced by ethnicity and genetic variations of patients. Overall, the relationship between nutraceutical agents, genetic variations, and other chemotherapeutic agents in different populations are not well understood in chemoprevention. Precision medicine strategies could provide a platform to address efficacy issues in PCa chemoprevention/intervention utilizing both chemotherapeutic and nutraceutical agents; however, additional clinical population studies are needed to confirm their efficacy against genetic, dietary, and environmental factors associated with PCa.

Precision medicine strategies utilize genomic analysis of PCa patients to identify and elucidate the pharmacogenomic landscape of patient susceptibility for the implementation of adjunct agents that can enhance the anti-tumor activity of pre-existing chemopreventive agents against advanced and/or resistant disease. This approach can aid in designing more specific treatment regimens for high-risk PCa patients. This review will evaluate the most current clinical research (summarized in Tables 1 and 2) on selected nutraceutical agents (silibinin, grapeseed extract, lycopene, soy isoflavonoids, green/black tea, vitamin E, vitamin D, zinc, and selenium) that may serve as adjunct agents in PCa. The efficacy of these nutraceutical agents will be discussed in the context of their effect on clinico-pathological outcomes (i.e., Gleason score, tumor grade, survival, biochemical recurrence), and disease susceptibility in diverse populations based on clinical and epidemiological data. We will also identify research gaps associated with diverse populations and selected nutraceuticals in PCa.

1Department of Pharmaceutical Sciences, Skaggs School of Pharmacy and Pharmaceutical Sciences, Aurora, CO, USA and 2University of Colorado Cancer Center, University of Colorado Anschutz Medical Campus, Aurora, CO, USA
Correspondence: Rajesh Agarwal (Rajesh.Agarwal@UCDenver.edu)
Received: 5 February 2018 Revised: 18 June 2018 Accepted: 21 June 2018
Published online: 25 July 2018

Published in partnership with The Hormel Institute, University of Minnesota
Silibinin
Over the past two decades, silibinin (flavanolignan from milk thistle “Silybum marianum” seeds) has shown strong antitumorogenic effects against various types of tumors including PCa. The significant anti-cancer efficacy of silibinin observed in preclinical animal models of PCa led to its transition into a phase II clinical trial to evaluate its bioavailability in patients diagnosed with localized PCa disease. Prior to surgery, PCa patients either received 13 g of silybin-phytosome for 14–31 days (n = 6) or served as untreated control subjects (n = 6). High dose of oral silybin-phytosome achieved high plasma concentrations in patients; however, very small amount of silibinin was observed in prostatic tissue due to its short half-life. Although serum prostate-specific antigen (PSA) levels of patients did not achieve a partial or complete response to silibinin treatment, disease stability was maintained in several patients. Thus, larger clinical studies are still needed to be performed with a more bioavailable form of silibinin to validate its biological efficacy as an effective nutraceutical agent for the clinical management of localized or advanced forms of PCa.

Grape seed extract (GSE)
GSE is a complex mixture of polyphenols containing procyanidins and their gallate derivatives together known as the proanthocyanidins. It has shown anti-cancer efficacy against PCa growth and progression in several preclinical models; however, clinical studies have not fully evaluated GSE efficacy in PCa patients. Interestingly, in a 2011 prospective study “VITamins And Lifestyle (VITAL)” (n = 1602; PCa cases; n = 35,239 total participants), analyzing the biological outcomes of intake of several dietary supplements (for approximately 5–10 years), GSE consumption stood out as the one associated with reduced risk for PCa (41% reduction in the risk of mortality among PCa patients relative to non-users). Importantly, we along with a team of medical oncologists have recently initiated a phase II study of GSE product in asymptomatic or minimally symptomatic non-metastatic PCa patients with rising PSA (NCT03087903), wherein GSE efficacy will be examined in a cohort of PCa survivors who have undergone treatment but show signs of rising PSA after local therapies. Given that current clinical studies have not yet identified molecular signatures modulated by GSE in PCa patients and this agent is a widely consumed as a supplement and food additive, more studies are needed to identify synergistic and/or additive interactions of GSE and its constituents with clinically used anti-PCa agents.

Lycopene and other carotenoids
Lycopene is a powerful carotenoid antioxidant with anti-tumor activity and present in red fruits and vegetables (i.e., tomatoes (tomato-based products), grapefruit, watermelons, and papayas). Other carotenoids-related compounds such as α-carotene, β-carotene, β-cryptoxanthin, xanthophyll carotenoids, lutein, and retinol also have shown anti-tumor activity. Interestingly, α-carotene, β-carotene, and β-cryptoxanthin can be converted into

Fig. 1 Nutraceutical efficacy in precision medicine. The schematic above depicts a workflow of experimental designs and assessment parameters required to establish the efficacy of nutraceutical agents. Therapeutic potential of agents must undergo vigorous pharmacodynamics and pharmacokinetics (blue) evaluation of their antioxidant and anti-cancer properties (e.g., anti-proliferation, anti-growth, anti-motility, anti-invasion) in cell models (gray). Targets that regulate altered tumor phenotypes are assessed via cell-focus assays (qRT-PCR, western blot, and immunofluorescence) and high-throughput platforms. Network analyses of targets are performed using omic profiling databases and libraries (brown) to determine gene ontology and identify therapeutic targets in cancer-associated canonical/non-canonical pathways (purple). Therapeutic targets and nutraceutical agents are evaluated in preclinical models and undergo the previous workflow. Next, nutraceutical agents are assessed in epidemiological studies and clinical trials (gray) which can evaluate hereditary, genetic, and environmental factors that range in degree from Phase I (n = 5–10 patients, Pharmacodynamics/Pharmacokinetics parameters), Phase II (2–3 treatment groups including standard treatment + new agent, different doses, safety and toxicity assessments, and Pharmacogenomics in humans or animals), Phase III (comparison between new agent and standard treatments, Pharmacogenomics (blue) (gray), Phase IV (Pharmacogenetic testing and side effects in different populations) to marketing and therapeutic application. After clinical trials have assessed the efficacy of nutraceuticals, these agents can be implemented in current precision medicine clinical therapeutic strategies for patients. The images of the mice and the group of people shown here are created by the authors.
Table 1. Epidemiologic and clinical intervention studies with phytochemical nutraceuticals in prostate cancer

| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|-----------------|------------|--------------|---------------------|---------|-----------|
| Silybin          | Phase II   | Daily oral administration of silybin-phytosome (13 g) in three doses daily prior to surgery | Univ. of Colorado Denver 12 PCa cases Ethnicity: N/a (Years: 2006–2010) | High-dose oral silybin-phytosome achieves high blood concentrations (mean 19.7 µM) transiently, but low levels of silybin are seen in prostate tissue (496.6 pmol/g) | Flaig et al. 4 |
|                 | Non-randomized | PCa patients aged ≥ 18 years with localized PCa scheduled for prostatectomy Time: For 14–31 days | | | |
|                 | Case-control cohort | | | | |
|                 |           | | | | |
| Grape seed extract | Case-control cohort | Daily use of grape seed extract for 5–10 years Men aged 50–76 years, (Surveillance, Epidemiology, and End Results (SEER) program cancer registry) filled base line questionnaire (and then followed for incident PCa) Time: For 7 years and median follow-up time of 6.1 years | VITamins and Lifestyle (VITAL) study cohort Residents in the 13-county area of Western Washington State (n = 1602 PCa cases; 35,239 total participants) Ethnicity: EA [1501 cases, 30,918 ctrls] AA [26 cases, 412 ctrls] Other [57 cases, 1876 ctrls] | No change in serum PSA, IGF-I, and IGFBP-3 levels | |
|                 |           | | | | |
| Grape seed extract | Phase II | Daily oral administration of grape seed extract (Leucoselect Phytotherax, 300 mg) Asymptomatic non-metastatic PCa patients with rising PSA Adults ≥ 18 years of age Histologically confirmed PCa. Evidence of rising PSA, baseline PSA must be ≥ 0.2 ng/mL at the time of screening Time: For 1 year | Univ. of Colorado Denver/Cancer Center Recruitment in progress | | NCT03087903 |
|                 | Open-label, single-arm study | | | | |
|                 |           | | | | |
| Lycopene        | Phase I–II | Daily administration of dietary selenium (55 µg), lycopene (35 mg), and green tea polyphenols (600 mg) Patients diagnosed with multi-focal high-grade prostatic intraepithelial neoplasia (HGPN) and/or atypical small acinar proliferation (ASAP) Time: For 1 month (Phase I), 6 months (Phase II); mean follow-up of 37 months | Univ. of Turin, Italy 10 cases (Phase I), 60 cases (Phase II) Ethnicity: Italian (Years: 2009–2014) | No significant variations in PSA (assessed by International Prostate Symptom Score questionnaire) No significant change in mean PSA and DRE assessments after 6 months Higher PCA diagnoses were in intervention group compared to placebo (p = 0.053) | Gontero et al. 43 |
|                 | Randomized, double-blind, placebo-controlled trial | | | | |
|                 | Case cohort | | | | |
|                 |           | | | | |
| Lycopene and β- cryptoxanthin | Multi-disciplinary | Daily carotenoid intake (dietary and/or supplementation) assessed for 1 year Prior to diagnosis of prostate adenocarcinoma in PCa patients (ages 40–79 years) | Data from North Carolina-Louisiana PCa project Ethnicity: AA [n = 1023] EA [n = 1079] (Years: 2004–2009) | Total lycopene dietary and supplemental intake was inversely related to PCa aggressiveness in EAs (OR = 0.56, 95% CI: 0.34–0.90, highest vs. lowest tertile, p-trend = 0.03) Dietary β-cryptoxanthin intake was inversely related to PCa aggressiveness in AAs (OR = 0.56, 95% CI: 0.36–0.87; p-trend = 0.01) | Antwi et al. 38 |
|                 | Cross-sectional study | | | | |
|                 | Case cohort | | | | |
|                 |           | | | | |
| β-cryptoxanthin, cis-lutein/zeaxanthin, and all-trans-lycopene | Prospective | Weekly intervention of high intake of plant-based foods (whole grains, fruits, vegetables, and legumes (soybean products) and exercise) and low intake of meat and dairy products Patients diagnosed with biochemically recurrent PCa Time: For 6 months (3 months of active intervention followed by monthly boosters) | Midlands Region of South Carolina 39 cases Ethnicity: EA [n = 28 (72%)] AA [n = 11 (28%)] | Plasma levels of β–cryptoxanthin (p = 0.01) and all-trans-lycopene (p = 0.004) were inversely related to PSA levels after dietary changes Lower PSA levels at 3 and 6 months were associated with higher plasma levels of trans-β-carotene relative to baseline High plasma levels of β–cryptoxanthin, cis-lutein/zeaxanthin, and all-trans-lycopene were associated with lower PSA levels after 6 months | Antwi et al. 34 |
|                 | Randomized trial | | | | |
|                 | Case cohort | | | | |
Table 1 continued

| Natural product                     | Study type                          | Intervention                                                                 | Population/location                  | Outcome                                                                 | Reference                  |
|-------------------------------------|-------------------------------------|------------------------------------------------------------------------------|--------------------------------------|------------------------------------------------------------------------|----------------------------|
| Lycopene-rich tomato products       | Randomized-controlled trial         | Daily administration of (a) tomato products containing lycopene (30 mg) per day; (b) tomato products plus (green tea (1 cup), black tea (1 cup), pomegranate juice (330 mL), grape juice (330 mL), soy-isoflavones (200 mg), 1-selenomethionin (200 µg), omega-3 fatty acids (3.13 g n-3 fatty acids); (c) control (habitual) diet | Oslo Univ. Hospital, Norway 86 cases Ethnicity: N/a (Years: 2007–2012) | Tomato intervention alone decreased median serum PSA levels (2.9%) significantly compared to controls (+6.5%) (p = 0.016) in the intermediate-risk group (post hoc analyses) | Paur et al.¹¹                |
| Lycopene/fish oil                   | Randomized, double-blinded placebo-controlled trial | Daily administration of two lycopene (15 mg), three fish oil capsules (fish oil (1 gm), eicosapentaenoic acid (EPA) (1096 mg) and docosahexaenoic acid (DHA) (549 mg) fatty acids), and one multi-vitamin or placebo; patients (young adult to older) with low burden PSA (Gleason score ≤ 6, PSA ≤ 10 ng/mL, positive cancer biopsy ≤ 33%); Time: For 90 days | Molecular Effects of Nutrition Supplements (MENS) Univ. of California San Francisco 84 patients Ethnicity: EA (78–83%) (Years: 2003–2008) | High dietary intake of tomato was strongly associated with changes in Selenoamino Acid Metabolism (p = 0.0029) and androgen/estrogen metabolism for both high intake of tomato and fish in morphologically normal prostate tissue | Magbanua et al.²⁹          |
| Lycopene-rich tomato extract        | Phase II Randomized, double-blind, placebo-controlled trial | Daily administration of 2 capsules of Lyc-O-Mato (tomato oleoresin, 15 mg of lycopene, phytoseo (1.4 mg), phytotocopherol (1.1 mg), β-carotene (0.7 mg), and α-tocopherol (4 mg)) or placebo (medium-chain triglycerides and red food coloring); patients (aged 35–75 years) diagnosed with high-grade prostatic intraepithelial neoplasia (HGPIN) (no cancer, atypical small acinar proliferation (ASAP), or antioxidant supplement use); Time: For 6 months | Northwestern Memorial Hospital and the Jesse Brown Veterans Administration Medical Center, Chicago 58 patients Ethnicity: EA [n = 42 (19 treatment, 23 placebo)]; AA [n = 15 (6 treatment, 9 placebo), Other [n = 11 (treatment)] (Years: 2006) | High expression of MCM-2 in basal epithelial cells p27 in luminal epithelial cells in benign prostate tissue | Gann et al.²⁸               |
| β-carotene/other agents             | Large population based Randomized, double-blind, placebo-controlled trial | Daily administration of a capsule containing either a placebo or a combination of vitamin C (120 mg), α-tocopherol (30 mg), β-carotene (6 mg), selenium (100 µg), and zinc (20 mg); middle-aged patients without severe health problems; Time: For 8 years | SU. VI. MAX trial France and Canada 5141 patients Ethnicity: Caucasian (Years: 1994–2002) | Reduced rate of PCa by 48% (HR = 0.52; 95% CI = 0.29–0.92; p = 0.009) among men with normal baseline PSA (<3 µg/L) Non-significant increase in PCa incidence (HR = 1.54, 95% CI = 0.87–2.72) in men with elevated PSA levels (≥3 µg/L) No effect on serum PSA and IGF levels | Meyer et al.³⁸             |
| β-carotene                          | Randomized, double-blind, placebo-controlled trial | Daily administration of α-tocopherol (50 mg), β-carotene (20 mg), both α-tocopherol and β-carotene, or placebo; patients aged 50–69 years with a smoking history; Time: For 5 years, median follow of 6.1 years, follow-up of 18 years | Alpha-Tocopherol, β-Carotene Cancer Prevention (ATBC) Study 25,563 patients in Southwestern Finland Ethnicity: Finnish (Years: 1985–1993; National Registries follow-up till 2011) | β-carotene intake increased post trial PCa mortality (RR = 1.20; 95% CI = 1.01–1.42) relative to non-recipients No significant effect of β-carotene intake on PCa incidence | Virtamo et al.²⁶          |
| Soy                                 | Randomized double-blind, placebo-controlled trial | Daily administration of soy isoflavone capsules [total isoflavones (80 mg/day), aglucon units (51 mg/day)]; patients diagnosed with localized PCa; Time: For 6 weeks prior to prostatectomy | Univ. of Kansas Hospital and Kansas City Veteran Affairs, Medical Center, 86 patients Ethnicity: EA n = 69 (38 [90%] treatment, 31 [70%] controls) AA n = 12 [3 [7%] treatment, 9 [20%] controls]; American Indian or Alaska Native n = 3 [12%] treatment, 2 [5%] controls (Years: 2006–2009) | Downregulated cell cycle and apoptosis-associated genes in prostate tumor tissue No effect on serum levels of total testosterone, free testosterone, total estrogen, estradiol, PSA, and total cholesterol | Hamilton et al.¹¹²          |
| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|-----------------|------------|--------------|---------------------|---------|-----------|
| Soy             | Phase II, Randomized trial, Case cohort | Daily consumption of soy bread (2 slices of soy isoflavones/slice) | Ohio State Univ. Medical Center, Columbus, Ohio | Decreased plasma levels of pro-inflammatory cytokines, Th1, T regulatory (CD4+CD25+Foxp3+), and monocyteic (CD33+HLADR+CD14+) myeloid-derived suppressor cells (MDSC) | Lesinski et al. 32 |
| Soy             | Phase II, Randomized trial, Case cohort | Daily consumption of soy bread (2 slices of soy isoflavones/slice) | Ohio State Univ. Medical Center, Columbus, Ohio | Increased serum PSA (PSA ≥ 4.0 ng/mL) relative to placebo (p = 0.006) | Anh Jarvis et al. 31 |
| Soy             | Phase II, Randomized placebo-controlled double-blind trial | Daily consumption of a supplement containing isoflavones (40 mg) and curcumin (100 mg) | Teikyo Univ. School of Medicine, Tokyo, Japan | Decreased serum PSA levels in patients with high PSA (PSA ≥ 10 ng/mL) (p = 0.01) | Ide et al. 36 |
| Soy protein isolate | Randomized placebo-controlled trial | Daily consumption of soy protein isolates (40 g of protein): (1) soy protein (SPI+, 107 mg of isoflavones), (2) alcohol-washed soy protein (SPI+, <6 mg of isoflavones), or (3) milk protein (MPI) | Minneapolis Veterans Administration Medical Center, Univ. of Minnesota | Decreased Bax expression in prostate tissue from SPI group relative to MPI group (p = 0.03), but no changes in EGFr, Bcl-2, Bax:Bcl-2 or Bax:PCNA ratios among treatment groups | Hamilton-Reeves et al. 36 |
| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|-----------------|------------|--------------|---------------------|---------|-----------|
| Soy             | Pilot      | Daily administration of three soy isoflavone tablet (Novasoy (genistein to daidzein (10:1:3)); 27.2 mg isoflavone aglycones) or a placebo | Stanford Univ. School of Medicine, California | Decreased COX-2, prostaglandin (PG) receptors (EP4 or FP) expression in human PCA cell lines (LNCaP; PC-3) and primary prostate epithelial cells | Swami et al.35 |
|                 | Randomized double-blind placebo-controlled trial | Patients newly diagnosed with PCa who will undergo prostatectomy and received no therapy | Ethnictiy: N/A | Outcome | Reference |
|                 |            | - Time: For 6 months | | | |
| Tea polyphenol (Green tea) | Phase 1–II | Daily administration of dietary supplement [selenium (55 mg), lycopene (35 mg), and green tea catechins (600 mg)] or placebo | Univ. of Turin, Italy | No significant change in mean serum PSA levels | Gontero et al.43 |
|                 | Randomized double-blind placebo-controlled trial | Patients diagnosed with multi-ocular high-grade prostatic intraepithelial neoplasia (mHGPIN) and/or atypical small acinar proliferation | 10 cases (Phase II), 60 cases (Phase 8) | Higher PCA diagnoses were in intervention group at re-biopsy compared to placebo (p = 0.053) | |
|                 | Case cohort | Ethnicity: Italian | (Years: 2009–2014) | Unregulated miRNAS (26b-5p, let-7i-5p, let-7d-5p, 15a-5p) and downregulated miR-494, an oncosuppressor, in PCa relative to normal tissue | |
| Tea polyphenol (Green tea) | Randomized, double-blind, placebo-controlled trial | Daily administration of Polyphenon E (800 mg) or placebo | Arizona Cancer Center, Tucson, and NCI | No significant effect on serum PSA, and insulin-like growth factor levels | Nguyen et al.41 |
|                 | Case cohort | Patients (PSA >50 ng/mL) who received no therapy and scheduled for radical prostatectomy | 48 cases | - No change in proliferative and angiogenesis biomarkers expression in prostate tissue | |
|                 |            | - Time: For 3–6 weeks prior to surgery | Ethnicity: White [n = 45], Native American [n = 1] | | |
| Tea polyphenol (Green/Black tea) | Phase II | Daily intake of brewed green tea [6 cups; EGCG (562 mg)] and/or black tea [6 cups, EGCG (28 mg), theaflavins (35 mg), Gallic acid (348 mg)] or control water | Veterans Administration Greater Los Angeles (LA) Health System; Univ. of California (LA), UCLA-Santa Monica Medical Center. | Reduced NFκB nuclear levels in radical prostatectomy tissue in green tea cohorts (p = 0.013) but not black tea (p = 0.931) | Henning et al.40 |
|                 | Open-label | Patients with localized PCa and scheduled for radical prostatectomy | 93 cases | Green tea decreased serum PSA levels and oxidative DNA damage marker in urine | |
|                 | Prospective randomized trial | Time: For 3–8 weeks prior to surgery | Ethnicity: White [n = 63], AA [n = 17], Asian [n = 4], Other [n = 8] | | |
| Tea polyphenols (EGCG) | Phase II | Daily administration of Polyphenon E [EGCG (800 mg)] with lesser amounts of (−)-epicatechin, (−)-epigallocatechin, and (−)-epigallocatechin-3-gallate] | USA | Polyphenon E decreased serum levels of PSA, HGF, VEGF, IGF-1, IGFBP-3, and the IGF-1/IGFBP-3 ratio in PCA patients | McLarty et al.38 |
|                 | Open-label, single-arm two-stage trial | Patients aged 18–75 years diagnosed with prostate cancer | 26 patients | Race had no significant effect in relation to intervention on serum biomarkers | |
retinol also known as vitamin A. Epidemiologic and clinical evidence have shown an inverse relationship between dietary lycopene (including other carotenoids), PCa development and disease progression risk. In a randomized-controlled study, a significant decrease in median PSA levels among PCa patients in the lycopene intervention group relative to control subjects was observed. Based on non-metastatic PCa patients in the Cancer Prevention Study II Nutrition Cohort, prediagnosis and postdiagnosis dietary lycopene intake did not modify PCa-specific mortality; however, lycopene intake higher than the median value was significantly associated with 59% lower hazard ratio among high-risk PCa patients. Notably, in a recent prospective study in patients from the Health Professionals Follow-Up cohort, over a 23-year follow-up, average tomato sauce intake was associated with a 46% reduction in risk of TMPRSS2-ERG-fusion positive PCa disease. High circulating levels of cis-lutein/zeaxanthin, all-trans-lycopene, and β-cryptoxanthin after 3 months corresponded to lower PSA levels at 3 and/or 6 months, respectively. High circulating levels of α-carotene, β-carotene, and total carotenoids lowered PCa susceptibility, however, circulating levels of lycopene were associated with a non-significant reduction in PCa risk.

Although carotenoids have exhibited preventive effects against PCa, other studies have reported little to no effect and/or antagonistic effect of carotenoid-associated nutraceuticals in PCa susceptibility. Epidemiologic and clinical studies have shown a lack of association between lycopene and PCa risk; however, retinol, a biosynthesis product of several carotenoids, was linked to a 13% increase in risk. Similarly, in the SU.VI.MAX study, a double-blind placebo-controlled and randomized trial, β-carotene had no effect on hazard ratios associated with PCa among 5141 men. In a cross-sectional study, high serum lycopene levels and total PSA > 2.5 ng/mL were associated with a 1.49-fold increase in PCa risk. Whereas, serum or plasma levels of retinol, β-carotene, β-cryptoxanthin, lutein and/or zeaxanthin, and total carotenoids did not significantly modify risk. Higher risk of PCa mortality was linked to β-carotene intervention in men relative to non-users in the Alpha-Tocopherol, Beta-Carotene Cancer Prevention (ATBC) Study (n = 29,133). In a prospective study, serum levels of α-carotene, β-carotene, lycopene, retinol did not show a relationship with time to disease progression, treatment, PSA kinetics (i.e., PSA, PSA velocity) and adverse histology in patients with localized disease. Likewise, another prospective study showed no association between lycopene and β-carotene serum levels and PCa risk in Australian men. In the PCa Prevention Trial (PCPT), dietary intake of carotenoid excluding lycopene had no effect on the risk of total incident symptomatic benign prostatic hyperplasia (BPH) among placebo arm participants after a 7-year follow-up (n = 4770). In the intervention arm of this study, lycopene also showed no significant association with BPH risk. Notably, carotenoid intervention and circulating levels in relation to PCa development and advanced disease have resulted in null findings in other reports.

Lycopene supplementation has shown a modest impact on cellular death, cell cycle, growth, and oxidative stress signaling mechanisms. Recent studies included in this review showed no molecular changes related to other carotenoids. In a Phase II clinical trial, 6 months of lycopene intervention marginally reduced nuclear levels of proliferative marker MCM-2 and cell cycle regulator p27 in benign prostate tissue from patients diagnosed with high-grade prostastic intraepithelial neoplasia (PIN). After a 3-month lycopene or tomato product intervention, analysis of normal prostate tissue from low-risk PCa patients showed apoptotic signaling and nuclear factor (erythroid derived-2) 2-mediated oxidative stress response as the top two ranked pathways altered by treatment compared to placebo.
| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|-----------------|------------|-------------|---------------------|---------|-----------|
| Vitamin D       | Case-control cohort | Daily oral administration vitamin D3 (4000 IU) | USA | Reduced immune and inflammation signaling in PCA transcriptome | Hardiman et al. |
| Vitamin D       | Phase I | Open-label multi-center, non-randomized dose-escalation study | Oral administration of inositol (40–800 µg) daily, or twice a day in combination with a 1-h intravenous infusion of doxetaxel (75 mg/m²), every 3 weeks and oral prednisone (5 mg twice a day) | France | Increased expression of immune and inflammation-associated genes in AAs relative to EAs | Medioni et al. |
| Vitamin D       | Phase IIa | Randomized placebo-controlled trial | Daily oral administration of cholecalciferol (vitamin D3 200,000 IU) as one dose at study entry plus genistin (1–2.5 mg/day), or placebo cholecalciferol day 1 and placebo genistin daily | Univ. of Wisconsin chemoprevention consortium | Well tolerated at 4000 µg | Jarrad et al. |
| Vitamin D       | Phase III | Multi-center, Randomized open-label study | Daily oral administration of ASCENT [high-dose calcitriol (45 µg), docetaxel (36 mg/m²), and dexamethasone (24 mg)] for 3 out of every 4 weeks or control (prednisone [5 mg] twice daily with doxetaxel [75 mg/m²], and dexamethasone [24 mg] every 3 weeks) | ASCENT Study | More deaths in ASCENT arm and trial was halted | Scher et al. |
| Vitamin D       | Case-cohort design nested within SELECT | Daily administration of selenium (200 µg of L-selenomethionine) + vitamin E (400 IU of all-rac-α-tocopheryl acetate), vitamin E and placebo, selenium and placebo, and placebo | Data from Selenium and Vitamin E Cancer Prevention Trial (SELECT) | Both low and high vitamin D concentrations were associated with increased risk of PCA, and more strongly for high-grade disease | Kristal et al. |
| Vitamin D       | Phase II | Randomized, double-blind, placebo-controlled cancer | Daily oral administration of vitamin D3 (cholecalciferol) at doses 400, 10,000, or 40,000 IU | University Health Network | Prostatic (1,25(OH)2D) concentrations showed that VDR was significantly lower in prostate tissues with the highest concentration of 1,25(OH)2D | Giangreco et al. |

Other notes:
- Vitamin D concentrations showed that VDR was significantly lower in prostate tissues with the highest concentration of 1,25(OH)2D.
- IL-6 expression was the highest in the prostate stroma, while PTGS2 (COX2) levels were lowest in the prostate cancer tissues from men in the highest tertile of prostatic 1,25(OH)2D.
- TNF-α, IL-6, and IL-8 were suppressed by 1,25(OH)2D in the primary epithelial cells, whereas TNF-α and PTGS2 were suppressed by 1,25(OH)2D in the stromal cell.
### Table 2

| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|-----------------|------------|--------------|---------------------|---------|-----------|
| **Vitamin E**   | Randomized, double-blind, placebo-controlled cancer prevention trial | Daily administration of α-tocopherol (all-rac-α-tocopheryl acetate 50 mg) or β-carotene (20 mg) or both α-tocopherol and β-carotene, or placebo | Alpha-Tocopherol, β-Carotene Cancer Prevention (ATBC) Study 25,563 patients in Southwestern Finland Ethnicity: Finnish (Years: 1985–1993; National Registries follow-up till 2011) | α-tocopherol reduced post-trial PCa mortality (RR = 0.84; 95% CI = 0.70-0.99) relative to non-recipients | Virtamo et al.66 |
| **Vitamin E**   | Randomized, double-blind, placebo-controlled trial | Daily oral administration of selenomethionine (200 μg), vitamin E (all-rac-α-tocopheryl acetate, 400 IU), or both or placebo | SELECT study 1746 PCa incident cases and sub-cohort of 3211 derived from SELECT trial 427 sites (United States, Canada, and Puerto Rico) Ethnicity: AA [n = 251(14.4%) cases, n = 735 (24.4%) ctrls] Hispanic [n = 58 (3.3%) cases, n = 139 (4.6%) ctrls] EA [n = 1406 (80.5%) cases, n = 2083 (69.2%) ctrls] Other [n = 31 (1.8%) cases, n = 51 (1.7%) ctrls] | High PCa incidence in men supplemented with high-dose α-tocopherol | Albanes et al.67 |
| **Vitamin E**   | Double-blind, placebo-controlled cancer prevention trial | Daily oral administration of α-tocopherol (all-rac-α-tocopheryl acetate 50 mg), β-carotene (20 mg), both or placebo | Patients with no PCa history, PSA ≤ 4 ng/mL and non-abnormal DRE Time: For 0–7.9 years, median follow-up of 5.5 years The trial was stopped early due to lack of efficacy of either supplement | No significant late effects (follow-up of 18 years) of α-tocopherol intake on PCa incidence | Virtamo et al.68 |
| **Vitamin E**   | Double-blind, placebo-controlled trial | Oral administration of antioxidant moiety of vitamin E [APC-100, (900–2400 mg)] | USA 20 cases Ethnicity: AA [n = 13 (65%), n = 5 (25%), Asian [n = 2 (10%)]] | Serum metabolomic response to supplementation determined | Kyriakopoulos et al.69 |
| **Vitamin E**   | Double-blind, placebo-controlled cancer prevention trial | Oral administration of antioxidant moiety of vitamin E [APC-100, (900–2400 mg)] | Data from ATBC study 200 cases (100 aggressive) and 200 controls Southwestern Finland Ethnicity: Finnish (Years: 1985–1993) | Downregulated vitamin E (γ-tocopherol, β-tocopherol) and amino acid (N6-acetyllysine, β-alanine, ornithine, and glutarylcarnitine) metabolites Upregulated vitamin E (α-tocopherol), co-factor (α-CEHC sulfate, α-CEHC glucuronide) and carbohydrate (fructose) metabolites | Mondul et al.170 |
| **Vitamin E**   | Case cohort | Daily intake/administration of vitamin E supplements (30, 100, 200, 400, 600, or 800 IU) assessed | Data from North Carolina-Louisiana PCa project 2102 cases Ethnicity: AA [n = 1023] EA [n = 1079] (Years: 2004–2009) | Dietary and supplemental α-tocopherol and PCa aggressiveness were inversely related in AAs (p-trend = 0.2, 0.15) Dietary intake of α- (p-trend = 0.006) and δ-tocopherol (p-trend = 0.007) were related inversely to PCa aggressiveness among EAs | Antwi et al.14 |
| **Vitamin E**   | Prospective study | Weekly intervention of high intake of plant-based foods (whole grains, fruits, vegetables, and legumes (soybean products) and exercise) and low intake of meat and dairy products Patients diagnosed with biochemically recurrent PCa Time: For 6 months (3 months of active intervention followed by monthly boosters) | Midlands Region of South Carolina 39 cases Ethnicity: AA [n = 28 (72%), n = 11 (28%)] | After adjusting for baseline PSA levels, plasma levels of α-tocopherol (p = 0.01) at 3 months were inversely related to serum PSA levels at 6 months Lower serum PSA levels at 3 and 6 months were associated with percent increase in plasma levels of α-tocopherol relative to baseline | Antwi et al.66 |
| **Vitamin E**   | Randomized trial | Administration of vitamin E (400 IU) every other day, vitamin C (500 mg) daily, or their | Physicians Health Study II Total of 14,641 US Physicians enrolled | Supplementation had no effect on PCA incidence | Wang et al.70 |
Table 2 continued

| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|-----------------|------------|--------------|---------------------|---------|-----------|
| **Vitamin E/lycopene** | **Prospective** | Daily administration of α-tocopherol (50 mg), β-carotene (20 mg), both α-tocopherol and β-carotene, or placebo | Physicians aged ≥ 50 years (Years: 1997–2007), follow-up till 2011 | Median follow-up of 10.3 years, post-trial follow-up of 2.8 years; Time: 10.3 years, post-trial follow-up of 2.8 years | Mondul et al.73 |
| **Vitamin E/ selenium** | **Randomized, double-blind, placebo-controlled trial** | Daily oral administration of selenomethionine (200 μg), vitamin E (all-rac-α-tocopheryl acetate, 400 IU), or both or placebo | Patients aged 50–69 years with a smoking history (Years: 1985–1993), follow-up till 2011 | Time: 5 years, median follow of 6.1 years, follow-up of 18 years | Kristal et al.68 |
| **Vitamin E/ selenium** | **Case-cohort** | Daily oral administration of selenomethionine (200 μg), vitamin E (all-rac-α-tocopheryl acetate, 400 IU), or both or placebo | Patients with no PCa history, PSA ≤ 4 ng/mL and non-abnormal DRE | Time: 7–12 years, median overall follow-up of 5.46 years | Martinez et al.72 |
| **Vitamin E/ selenium** | **Case cohort study of SELECT trial participants** | Daily oral administration of selenomethionine (200 μg), vitamin E (all-rac-α-tocopheryl acetate, 400 IU), or both or placebo | Patients with no PCa history, PSA ≤ 4 ng/mL and non-abnormal DRE | Time: 7–12 years, median overall follow-up of 5.46 years | Chan et al.94 |
| **Selenium** | **Randomized double-blind, placebo-controlled trial** | Daily oral administration of selenium (200 μg /or 400 μg) or placebo | Patients with no PCa history, PSA < 4 ng/mL and/or abnormal DRE with negative prostate biopsy | Time: Follow-up of 5 years | Lu et al.85 |
| **Selenium** | **Case-cohort study of SELECT trial participants** | Daily oral administration of selenomethionine (200 μg), vitamin E (all-rac-α-tocopheryl acetate, 400 IU), or both or placebo | Patients with no PCa history, PSA ≤ 4 ng/mL and non-abnormal DRE | Time: For 7–12 years, median overall follow-up of 5.46 years | Nguyen et al.71 |

**Outcome**

- Energy and lipid-related serum metabolite levels were associated with low risk of aggressive PCa with the exception of Ercouyl-sphingomyelin and Trimethylamine N-oxide
- Serum levels of other metabolite chemical classes were not associated to non-aggressive or overall PCa risk
- Glycerophospholipid, long-chain fatty chain, and TCA metabolites were primarily modulated compared to other metabolites
- Vitamin E supplementation increased the risk of PCa among men with low selenium status
- Overall, low-grade, and high-grade PCa risk was higher among men with lower selenium status and receiving vitamin E supplements (p = 0.008)
- Vitamin E supplementation (alone) had no effect in men with high selenium status (≥40th percentile of toenail selenium) (p = 0.02)
- Selenium supplementation did not benefit men with low selenium status but increased the risk of high-grade PCa among men with high selenium status
- Men should avoid selenium and vitamin E supplementation at doses that exceed recommended dietary intakes
| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|----------------|------------|--------------|---------------------|---------|-----------|
| Selenium       | Phase II   | Daily oral administration of 200/or 800 µg of selenium or placebo | USA | - No significant effects on PSA velocity and Gleason score | Lu et al. 85 |
|                | Randomized, double-blind, placebo-controlled trial | | | Increased PSA velocities in the highest quartile of patients receiving high-dose selenium (800 µg) relative to placebo (p = 0.018) | |
|                | Case cohort | | | Total of four deaths in treatment groups | |
|                |            | Patients diagnosed with localized non-metastatic PCa (Gleason score < 8, PSA < 50 ng/mL, age < 85 years) | | | |
|                |            | Time: For placebo (36.3 months, follow-up 38.4 months); selenium 200 µg (33.3 months, 33.3 months), 800 µg (33.3 months, 33.8 months) | | | |
| Selenium       | Double-blind, randomized, placebo-controlled trial | Daily administration of selenium as selenomethionine 200 µg/day or placebo | NGC Intergroup trial/Southwest Oncology Group (SWOG) 423 randomized men with HGPIN | Selenium supplementation had no effect on PCa risk | Marshall et al. 120 |
|                |            | Men ≥40 years of age with a confirmed diagnosis of HGPIN lesions via biopsy with no evidence of PCa | | | |
|                |            | Time: For 3 years | | | |
|                |            | The primary endpoint was progression of HGPIN to PCa over a 3-year period | | | |
| Selenium       | Phase II/II | Daily administration of dietary supplement (selenium (55 mg)), lycopene (35 mg), and green tea catechins (600 µg) or placebo | Univ. of Turin, Italy | No significant change in mean serum PSA levels | Gontero et al. 13 |
|                | Randomized double-blind placebo-controlled trial | Patients diagnosed with multi-locus high-grade prostate intraepithelial neoplasia (mHGPIN) and/or atypical small acinar proliferation (ASAP) | | Higher PCa diagnoses were in intervention group at re-biopsy compared to placebo (p = 0.033) | |
|                | Case cohort | Time: For 1 month (Phase I), 6 months (Phase II), mean follow-up of 37 months | | | |
| Selenium       | Randomized, double-blind, double-dummy trial | Patient data from ProFluss’ intake 1 tablet/day (85% of fatty acids sterols, selenium(50 µg) and lycopene (5 mg)) and control | | No detrimental or protective role of supplementation in increasing PCa risk | Morgia et al. 27 |
|                | Multi-center | Patients aged 55–80 years diagnosed with lower urinary tract symptoms (negative DRE for PCa, PSA < 4 ng/mL) | | No significant differences in the mean serum PSA levels or Gleason score | |
|                |            | Time: For 1 year, follow-up of 2 years | | No effect on PCa risk (OR = 1.07; 95% CI = 0.64–1.79; p = 0.95), incidence (HR = 1.38; 95% CI = 0.32–5.90; p = 0.67) | |
| Selenium       | Randomized, placebo-controlled trial | Daily oral administration of selenium selenized yeast, 300 µg or placebo | Netherlands 23 cases | Downregulated genes associated with cell migration, invasion, remodeling, and immunity | Kok et al. 90 |
|                |            | Patients undergoing diagnostic prostate biopsies and radical prostatectomy | Ethnicity: Dutch | Exhibited an inhibitory effect against EMT via upregulation of epithelial markers (E-cadherin and EPCAM) and downregulation of mesenchymal markers (vimentin and fibronectin) | |
|                |            | Time: For 5 weeks | | | |
| Selenium       | Randomized, double-blind, placebo-controlled trial | Daily oral administration of selenomethionine (200 µg) or vitamin E (all-rac-α-tocopherol acetate, 400 IU), or both selenomethionine and vitamin E or placebo | SELECT study data 1746 PCa incident cases and sub-cohort of 3211 derived from SELECT trial 427 sites (United States, Canada, and Puerto Rico) | Increased PCa risk (HR = 2.04; 95% CI = 1.29–3.22) in patients receiving selenomethionine alone or in combination with α-tocopherol in the highest quintile relative to the first quintile (p-trend = 0.005) | Albanes et al. 67 |
|                | Case-control trial | Patients with no PCa history, PSA ≤ 4 ng/mL and non-abnormal DRE | Ethnicity: AA [n = 251(14.4%) cases, n = 735 (24.4%) controls] Hispanic [n = 58 (3.3%) cases, n = 139 (4.6%) controls] EA [n = 1406 (80.5%) cases, n = 2083 (69.2%) controls] Other [n = 31 (1.8%) cases, n = 51 (1.7%) controls] | Positively associated with plasma levels of α-tocopherol in patients receiving selenomethionine in the fifth quintile (HR = 2.12; 95% CI = 1.32–3.40; p-trend = 0.0002) | |
|                | (sub-study of participants within SELECT) | Time: For 0–7.9 years, median follow-up of 5.5 years | | Non-significant elevation of PCa risk associated with selenomethionine in the third tertile of plasma α-tocopherol levels relative to placebo | |
|                |            | The trial was stopped early due to lack of efficacy of either supplement | | | |
| Selenium       | Randomized-controlled trial | Daily administration of a) tomato products containing lycopene 30 mg per day; (b) tomato products plus (green tea 1 cup), black tea (1 cup), and a) vitamin E acetate, 400 IU, or combination of both b) | Oslo University Hospital, Norway 86 cases | Tomato products plus therapy slightly decreased (non-significant) serum PSA levels among intermediate-risk patients post surgery | Paur et al. 11 |
|                | Case cohort | | | | |
### Table 2 continued

| Natural product | Study type | Intervention | Population/location | Outcome | Reference |
|----------------|------------|--------------|---------------------|---------|-----------|
| Selenium       | Pilot      | consumption of lycopene juice (330 mL), grape juice (300 mL), soy isoflavones (200 mg), 1-selenomethionin (200 µg), omega-3 fatty acids (1.13 g n-3 fatty acids); (c) control (habitual) diet | American Health Foundation, New York and Penn State College of Medicine, Pennsylvania 36 healthy patients 60 patients | Decreased serum PSA levels with highest increases (Years: 2007–2012) | Sinha et al.91 |
| Selenium       | Pilot      | Daily administration of selenium-enriched yeast (SY) (247 µg) or placebo (non-enriched yeast) | American Health Foundation, New York and Penn State College of Medicine, Pennsylvania 36 healthy patients 60 patients | Upregulated (clusterin isoform 1 [CLU], transthyretin, α-transthyretin, α1-antitrypsin [AAT], angiotensin precursor and albumin precursor) several proteins | Morgia et al.93 |
| Selenium/lycopene | Multi-center | Daily administration of Profluss [1:1 ratio of SeR 320 mg + lycopene (5 mg) + selenium (50 µg) (group I), control (group II), SeR 320 mg + lycopene (5 mg), selenium (50 µg), and α-blockers treatment (group III), control (group IC)] | Flogosis And Profluss in Prostatic and Genital Disease (FLOG) study 9 centers (Italy) 108 patients (Group I) 60 patients (Group II) | Decreased serum PSA levels in Group I, but no difference in Group II | Morgia et al.93 |
| Selenium/multi-vitamins | Prospective cohort | Daily intake of multi-vitamins or individual supplement (such as, selenium, β-carotene, and zinc) | National Institutes of Health (NIH)-AARP Diet and Health Study 10,241 cases (8765 localized and 1476 advanced disease) | Increased risk of advanced (RR = 1.32; 95% CI = 1.04–1.67) and fatal (RR = 1.98; 95% CI = 1.07–3.66) PCa with excessive use of multi-vitamins relative to non-users | Lawson et al.66 |
| Zinc           | Case-control surveillance cohort | Daily use of multi-vitamin containing zinc, vitamin E, β-carotene, folate, and selenium | USA hospitals located in four centers (Baltimore, Boston, New York, and Philadelphia) 1706 cases, 2404 matched ctrls | Increased risk of advanced (RR = 1.32; 95% CI = 1.04–1.67) and fatal (RR = 1.98; 95% CI = 1.07–3.66) PCa with excessive use of multi-vitamins relative to non-users | Lawson et al.66 |
| Zinc           | Randomized, placebo-controlled trial | Daily intake and supplemental use of vitamin C, vitamin D, zinc, calcium, carotenoids, lycopene, EPA* plus DHA* or multi-vitamin weekly | Prostate Cancer Prevention Trial (PCPT) subjects 4770 patients | •BPH assessed by International Prostate Symptom Score questionnaire. Diet, alcohol, and supplement use assessed by food frequency questionnaire •Supplement intake did not affect BPH risk | Kristal et al.17 |

*Note: PCa = prostate cancer; PSA = prostate-specific antigen; BPH = benign prostatic hyperplasia; SeR = selenium-rich; SY = selenium-enriched yeast; AAT = α1-antitrypsin; AARP = American Association for Retired Persons; NIH = National Institutes of Health; OR = odds ratio; CI = confidence interval; RR = relative risk; BPH = benign prostatic hyperplasia; DRE = digital rectal examination; ASAP = African American Study of the Risk of Stroke; PCPT = Prostate Cancer Prevention Trial; FLOG = Flogosis In Prostatic and Genital Disease; EPA = eicosapentaenoic acid; DHA = docosahexaenoic acid.*
Table 2

| Study type | Intervention | Population/Location | Outcome |
|------------|--------------|---------------------|---------|
| Prospective | Daily use of vitamin E, selenium, and zinc supplement, no history of PCa | VITAL study cohort (USA) | Zinc intake and cadmium exposure in relation to recommended daily allowance and cadmium exposure in those with inadequate zinc intake. Linear trend analysis for discrimination between different zinc intakes (Years: 1988–1994, follow-up to 2006) | 
| Case-control cohort | Daily zinc intake and cadmium exposure in relation to recommended daily allowance and cadmium exposure in those with inadequate zinc intake. Time: 3.5 years | Third National Health and Nutrition Examination Survey (NHANES III) | CADM-PCa-risk (Years: 1988–1994, follow-up to 2006) | 
| Case-control cohort | Daily zinc intake and cadmium exposure in relation to recommended daily allowance and cadmium exposure in those with inadequate zinc intake. Time: Average follow-up of 3.5 years | Multistage, stratified sampling design | Zinc intake and cadmium exposure in relation to recommended daily allowance and cadmium exposure in those with inadequate zinc intake. Time: Average follow-up of 3.5 years | 
| Case-control cohort | Daily zinc intake and cadmium exposure in relation to recommended daily allowance and cadmium exposure in those with inadequate zinc intake. Time: For <1.8 years, average follow-up of 3.5 years | European-American, African-American, PCA prostate cancer, PSADT-specific, antiangiogenic, anti-inflammatory, anti-cancer properties of soy. | Time: Follow-up of 12.4 years | 
| Case-control cohort | Daily zinc intake and cadmium exposure in relation to recommended daily allowance and cadmium exposure in those with inadequate zinc intake. Time: Follow-up of <1.8 years, average follow-up of 3.5 years | European-American, African-American, PCA prostate cancer, PSADT-specific, antiangiogenic, anti-inflammatory, anti-cancer properties of soy. | Time: Follow-up of 12.4 years | 

Soy isoflavones

Soy isoflavones are members of the polyphenolic flavonoid family. These compounds are found in red clover, kudzu root, and soybeans, which are commonly used in Asian and African cuisines. Genistein, daizein, aglycones, equal and glycitein are the predominant isoflavones in soybean and soy-derived food products. Clinical studies on the efficacy of soy intervention have shown some benefits against PCa via its influence on insulin and inflammatory signaling. In a 20-week phase II trial in asymptomatic PCa patients, consumption of soy-almond bread and standard soy bread contributed to a significant increase in IGFBP-3 and decrease in pro-inflammatory cytokines blood levels. Unfortunately, dietary soy intervention increased serum PSA levels and its doubling time after 126 days with a slight decrease in PSA velocity in PCa patients. Similar to the previous report, isoflavones have exhibited some immunomodulatory properties in the plasma of asymptomatic biochemically recurrent PCa patients. High plasma genistein concentrations (>640.2 nmo/L) were strongly linked to a 69% decrease in risk of developing PCa among Chinese patients. At the time of diagnosis, median levels of plasma genistein were significantly lower in PCa patients relative to controls. Short-term administration of isoflavone (80 mg) for 6 weeks in patients also showed an inhibitory effect on cell cycle and apoptotic-associated signaling in prostate tumor tissue. However, isoflavone intervention did not affect serum levels of total testosterone, free testosterone, PSA, and total cholesterol in PCa patients.

Though limited reports have evaluated genomic evidence for the anti-cancer properties of soy isoflavones in PCa clinical trials, soy supplementation has exhibited some effects on inflammatory, apoptotic, and growth signals in PCa. In a pilot randomized double-blinded clinical study, soy isoflavones intervention altered the expression of COX-2, a major molecule in prostaglandin synthesis and cyclooxygenase inhibitor p21 in PCa patient prostatectomy specimens. In the same study, genistein treatment also downregulated COX-2 in both LNCaP and PC-3 cell lines and upregulated 15-PGDH in primary PCa cells. In a 6-month clinical trial, soy protein intervention had no effect on proliferative, and apoptotic molecular markers (i.e., EGFR, Bax/Bcl-2, Bax/PCNA ratios) in high-risk and/or with low-grade PCa disease patients, but alcohol-washed soy protein intake reduced tissue levels of PCNA and Bax in patients relative to milk protein treatment.

Green and black tea extracts

Green and black tea are extracted from the plant *Camellia sinensis*. (-)-Epigallocatechin-3-gallate (EGCG) is the most abundant and well-studied bioactive polyphenolic constituent of green tea with regard to its anti-cancer properties in several malignancies including PCa. Theaflavin is the major bioactive polyphenol from black tea; however, it has not been well studied compared to EGCG. Many population studies have examined the efficacy of green tea in PCa chemoprevention. In a recent meta-analysis of 13 clinical studies, green tea catechins demonstrated protective effect against PCa risk. Concurrent consumption of green tea catechins and natural food products in an adjusted indirect comparison relative to six other natural compounds significantly reduced PCa susceptibility in men diagnosed with high-grade PIN. Daily administration of Polyphenon E (an enriched green tea polyphenol extract) containing 800 mg of EGCG prior to a radical prostatectomy significantly reduced serum levels of PSA in men with cancer-positive prostate biopsies. However, in another study, Polyphenon E intake showed no significant effects on the serum levels of PSA, insulin-like growth factor, proliferation, and angiogenesis in the prostate tissue of PCa patients after 3–6 weeks. The efficacy of green tea was also evaluated in patients with localized PCa (n = 199) and receiving active surveillance or watchful waiting as clinical management...
treatment. Oral administration of a capsule containing nutraceuticals, pomegranate, green tea, broccoli, and turmeric for 6 months significantly decreased median PSA percentage levels in patients, regardless of clinical management relative to the placebo group. 39 Unfortunately, this intervention did not alter Gleason grades in patients.

Tea polyphenols have modulated several molecular signatures in PCA patients. Green tea intake reduced nuclear NfκB in radical prostatectomy tissue and PSA levels in PCa patients compared to black tea and control treatments 3–8 weeks prior to surgical therapy. 40 Growth factor signaling mediators, HGF, VEGF, IGF, and IGFBP-3, ratio was decreased in men with PCa. 38,41–43 Although expression of cell proliferative, apoptotic, and angiogenic markers was not changed by green tea intervention after 3–6 weeks prior to surgery, serum levels of PSA, IGF, and DNA oxidative stress in leukocytes were decreased by treatment in patients. 41 Also, green and black tea reduced the proliferation of LNCaP PCa cells. 42 Green tea in a combinatorial nutraceutical intervention promoted oncogenic-related miRNAs in PCa including miR-92-3p that targets PTEN and androgen-regulated miR-125-5p. 43

Vitamin D

Vitamin D is a fat-soluble nutraceutical found in dairy, flour, and fortified food products. 44 It has five isomers, which include ergocalciferol with lumisterol (D1), ergocalciferol (D2), cholecalciferol (D3), 22-dihydroergocalciferol (D4), and sitocalciferol (D5). Its biosynthesis occurs in the skin in response to solar ultraviolet B radiation exposure. In the body, Vitamin D primarily circulates as 25-hydroxyvitamin D [25(OH)D] and is converted by 1α-hydroxylase into its active form, 1,25-dihydroxyvitamin D [1,25(OH)2D]. Calcium and bone homeostasis are closely dependent on Vitamin D bioavailability. Vitamin D deficiency is linked to elevated susceptibility of PCa development and aggressive disease. Both clinical and other population-based studies have comprehensively evaluated the role of dietary vitamin D as a preventive therapy to reduce disease development and complimentary agent to accepted clinical treatments for PCa. Overall, dietary vitamin D levels have an inverse association to PCa development and disease progression. 45,46 Vitamin D deficiency (<20 ng/mL) alone increases the risk of PCa development and aggressive PCa in certain subpopulations. 47 In two large nested case-control studies, vitamin D intervention decreased the risk of developing PCa among patients with high Gleason scores. 48 In a cross-sectional study, dietary Vitamin D intake showed an inverse association with aggressive PCa (699 cases and 958 controls). 49 Also, green and black tea reduced the proliferation of LNCaP PCa cells. 42 Green tea in a combinatorial nutraceutical intervention promoted oncogenic-related miRNAs in PCa including miR-92-3p that targets PTEN and androgen-regulated miR-125-5p. 43

Vitamin D treatment has shown some anti-inflammatory and hormone-related molecular changes in PCa. 31,33,62,63 In Taiwanese PCa patients, genetic variants, HFE rs9393682, and TUSC-3 rs1378033 were associated with time to progression in localized disease and low risk of advanced PCA for patients undergoing androgen deprivation therapy. 62 Furthermore, in vitro studies revealed 1,25-Vitamin D downregulated HFE and when silenced HFE impedes cell proliferation and wound healing. Low expression of TUSC-3 was shown to correspond with poor PCA prognosis in patients, and TUSC-3 silencing enhanced cell migration and growth. 62 However, 1,25-Vitamin D strongly induced the expression of TUSC-3 in PCa cells. In a Phase II clinical trial, PCa patients with the highest serum levels of prostatic 1,25(OH)2D also had low COX-2 levels, but exhibited high IL-6 levels in their stroma tissue. 63 Interestingly, 1,25 (OH)2D treatment suppressed TNF-α, IL-6, and IL-8 levels in primary epithelial cells, but only TNF-α and COX-2 levels were downregulated in stromal cells. Similarly, circulating levels of 25(OH)D negatively correlated with pro-inflammatory markers, serum CRP, and IL-8, but NFκB p65-positive cells were elevated in PCa patients. 53 In early stage PCa, high-dose cholecalciferol and genistein (G-2535) intervention induced AR expression in patient tumor tissue but not benign tissue relative to placebo controls. 51

Vitamin E

The protective effect of nutraceutical, Vitamin E, in PCa has been widely studied in published reports and clinical trials. 9 Tocopherols (α, γ, and δ) possess vitamin E activity and are the most studied vitamin E bioactive constituents in PCa. Vitamin E is fat soluble and found in a variety of foods, such as nuts, seeds, and vegetable oil. Both European and Western diets have a high content of tocopherols. European diets mainly include α-tocopherol, whereas γ-tocopherol is generally present in the Western diets. In a pooled study including 15 cohorts (11,239 cases, 18,541 controls), α-tocopherol consumption was associated with a decrease in risk of PCa overall and aggressive disease susceptibility. 1 However, a small clinical trial, castrate-resistant PCa patients taking APC-100, an antioxidant moiety of α-tocopherol, maintained stable disease and median progression-free survival of 2.8 months. 56 In a 6-month clinical trial, high α-tocopherol levels were inversely related to serum PSA levels in biochemical recurrent PCa patients (n = 39). 14 In contrast, other studies have shown negative effects associated with vitamin E intervention or its circulating plasma levels in PCa patients. PCa risk was slightly increased in the National Institutes of Health (NIH)-AARP Diet and Health Study due to a high frequency (>7 times per week) of dietary Vitamin E. 65 High PCa risk estimates were associated with patients who used a dosage of 800 IU per day regardless of frequency. Furthermore, patients with family history of PCa and taking frequent multi-vitamin supplementation (>7 times per week) had a 2.48 and 16.41-fold increase in susceptibility of advanced and fatal PCa disease, respectively. In another report, dietary intake of both α-tocopherol and δ-tocopherols reduced risk of PCa in European-American patients. 66
Furthermore, positive plasma α-tocopherol levels were also linked to high-grade PCA disease (Gleason grade 7–10). However, vitamin E intervention combined with low selenium status increased PCA susceptibility in men. Although vitamin E supplementation has some negative effects in PCa, some large population studies have observed null findings in relation to BPH and PCA.

Vitamin E intervention, supplementation, and associated genetic variants have been shown to modulate PCA susceptibility among men. In the multi-center Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial, rs964184 variant GG genotype located near genes, BUD13, ZNF259, and APOA5, which plays a role in Vitamin E metabolism, showed a protective role against PCA risk with the inheritance of two or more minor alleles. However, other clinical trials have identified variants such as NIXX3.1 rs11781886 linked high risk of developing advanced PCA disease in the presence of Vitamin E intervention. In the ATBC study cohort, serum levels of tricarboxylic acid cycle, long-chain fatty acid, and glycerophospholipid metabolites were strongly associated with low risk of aggressive PCa with the exception of metabolites, thyroxine, and trimethylamine oxide. However, these metabolites had no significant interaction with α-tocopherol supplementation. Vitamin E-related transcripts involved in the transport of vitamin E influence PCa susceptibility as well. In a clinical study by Bauer and colleagues (2013), circulating levels of α-tocopherol or γ-tocopherol were associated with disease recurrence; however, superoxide dismutase enzyme 3 (SOD3) rs699473 variant were linked to high-grade PCa, but SOD1 (rs17884057, rs9967983) and SOD2 (rs4880) variants were protective against disease recurrence among men. In the multi-center Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial, rs964184 variant GG genotype located near genes, BUD13, ZNF259, and APOA5, which plays a role in Vitamin E metabolism, showed a protective role against PCA risk with the inheritance of two or more minor alleles. However, other clinical trials have identified variants such as NIXX3.1 rs11781886 linked high risk of developing advanced PCA disease in the presence of Vitamin E intervention. In the ATBC study cohort, serum levels of tricarboxylic acid cycle, long-chain fatty acid, and glycerophospholipid metabolites were strongly associated with low risk of aggressive PCa with the exception of metabolites, thyroxine, and trimethylamine oxide. However, these metabolites had no significant interaction with α-tocopherol supplementation. Vitamin E-related transcripts involved in the transport of vitamin E influence PCa susceptibility as well. In a clinical study by Bauer and colleagues (2013), circulating levels of α-tocopherol or γ-tocopherol were associated with disease recurrence; however, superoxide dismutase enzyme 3 (SOD3) rs699473 variant were linked to high-grade PCa, but SOD1 (rs17884057, rs9967983) and SOD2 (rs4880) variants were protective against disease recurrence among men.

Additionally, SOD1 rs17884057 variant and circulating α-tocopherol levels were also linked to high-grade PCa, but did not remain significant in the highest quartile of α-tocopherol.

Zinc

Normal prostatic tissue exhibits the highest levels of the mineral zinc compared to any soft tissue in the body. In earlier epidemiologic studies, dietary zinc demonstrated a protective effect in individuals diagnosed with advanced PCA susceptibility. However, recent reports oppose the preventive role of dietary zinc and contribute to mix findings on the relationship of zinc consumption and PCa risk. Long-term dietary zinc from multivitamins or other supplements in PCa patients showed a non-significant 2.0-fold elevation in PCA risk in the US hospital-based Case-Control Surveillance Study (1706 cases, 2404 matched controls). Epidemiologic evidence has also revealed an interaction between dietary zinc intake and cadmium exposure in relation to PSA levels. In the 2001–2002 National Health and Nutrition Examination Survey, dietary zinc intake less than the median level (<12.67 mg/day) was linked to an increase in creatinine cadmium exposure (1 µg/g). Cadmium is a non-essential heavy metal that is one of the naturally occurring composites in zinc. This relationship induced a 35% increase in PSA levels in patients. However, this association between cadmium exposure and PSA levels disappears at higher levels of dietary zinc intake. During the 7-year follow-up period in the PCPT study, dietary and total zinc intake decreased the susceptibility to develop total symptomatic BPH incidence among the placebo population in the second, fourth, and fifth quintiles. In the VITAL study, Gonzalez and associates showed that dietary zinc intake over a 10-year period had a non-significant decrease in overall PCa and advanced disease (regional/distant) mortality risk.

Other studies have shown either null and/or possible harmful effects of dietary zinc intake and PCa and/or other associated conditions. Varying levels of dietary zinc have been linked to potential PCa risk. Two studies showed that excessive use of multi-vitamin with zinc supplements was associated with high risk of PCa. In the NIH-AARP Diet and Health Study, excessive use of multi-vitamin including zinc was associated with a 4.36-fold increase in the risk of developing fatal PCa among 10,241 men (8765 localized and 1476 advanced cases). In other studies including the Case-Control Surveillance Study and SU.VI.MAX study, non-significant associations were shown between zinc intake and/or supplementation and circulating levels with PCa susceptibility among European-American, African-American, and Mexican-American men.

Unfortunately, current clinical studies have not evaluated the molecular effects of zinc intervention in PCa. However, two studies have evaluated some aspects of zinc homeostasis affected in PCa. It is well known that zinc levels as well as the expression of zinc transporters are low in PCa and this may impair zinc absorption in the body. Zinc transporters, hZn1P and hZn2P, were under-expressed in malignant prostate tissue compared to surrounding normal tissue. In another study, miR-182 expression in prostate tissue was higher in all PCa cases, but miR-182 and miR-346 expression were inversely related to hZn1P levels in European-American men only. Overexpression of miR-183, miR-96, and miR-182 reduced intracellular zinc levels and uptake in primary prostatic epithelial cells. Despite recent reports that suggest a possible antagonistic role for zinc in PCa, additional studies are still needed to fully evaluate the efficacy of dietary zinc or supplementation in patients due to limited literature on the molecular and clinical impact of this agent on PCa.

Selenium

Changes in the physiological levels of the mineral selenium may influence biochemical and metabolic processes in PCa. Significantly low levels of selenium have been detected in malignant prostatic tissue compared to BPH specimens. Dietary selenium intervention has been linked to null findings for PCa susceptibility. In a pooled study of prospective cohorts, the daily intake of 37 supplements including selenium (49–90 µg) did not modify PCa risk in British men. In the Italian cohort for the Procomet trial (ISRCTN78639965), selenium intervention for 2 years did not significantly change PSA levels, PCa susceptibility, or exhibit a strong association with high PCa mortality risk. In a nested case control study based on the Physicians’ Health Study and Health Professionals Follow-Up Study cohorts, circulating selenium levels did not significantly modify PCa susceptibility. No association was observed between circulating levels of selenium and PCa susceptibility considering TMPRSS2-ERG-fusion positive or negative cancer stratification relative to control subjects (370 cases, 2740 controls). Over the course of 3–5 years, selenium supplementation of 200 or 400 µg did not affect PSA velocity or risk of PCa mortality in high-risk patients (n = 699) in a Phase III randomized, double-blinded placebo-controlled, multi-center trial.

Interestingly, meta-analysis of 17 western population-based studies showed serum selenium was inversely related to PCa susceptibility in individual study cohorts. Arsenic exposure and plasma selenium levels influence PCa susceptibility in patients. In a case-control study on Taiwanese patients (318 cases and 318 controls), low plasma selenium (≤28.06 µg/dL) and high urinary arsenic concentration (>29.32 µg/L) were associated with elevated PCa risk in multi-variate analyses and significantly interacted with PSA levels (≥20 ng/mL). In a meta-analysis of 15 prospective studies on PCa patients (4527 cases and 6021 controls), high selenium plasma levels were associated with a lower risk of aggressive PCa and selenium nail content was inversely related to PCa risk, but not for blood selenium. High toenail selenium levels and selenoprotein P, a major selenium transporter, variants were associated with reduced advanced PCa risk.
controlled trial, tomato product intervention consisting of selenium slightly increased PSA levels among patients with intermediate risk of PCa \((n = 86)\).\(^{11}\) During this intervention, patients with high levels of lycopene, selenium, and fatty acid C20:5 n-3 combined were linked to a large decrease in serum PSA levels. Selenium intervention has also triggered some inconsistent molecular alterations in PCa. Selenium intervention \((300 \mu g)\) triggered a reversal of epithelial-to-mesenchymal transition (EMT) via upregulation of epithelial markers and lower expression of mesenchymal-related genes in a Dutch population compared to those receiving the yeast placebo.\(^{96}\) Selenium-enriched yeast reduced levels of proteins related to anti-apoptosis \((\text{clusterin isoform 1)}\), iron transport/homeostasis \((\text{transferrin, haptoglobin})\), oncogenesis \((\alpha-1B-glycoprotein)\), inflammatory response \((\text{complement component 4B proprotein})\), oxidative stress \((\text{keratin 1)}\), NADH metabolic process \((\text{isocitrate dehydrogenase})\) in healthy African-American and European-American male subjects in a randomized, double-blinded, placebo-controlled clinical trial.\(^{71}\) However, selenium upregulated levels of α-1 antitrypsin, a protein involved in hypoxia response, angiotensin precursor, negative regulator of cell growth and proliferation, and albumin, a negative regulator of apoptosis. Seven out of the 11 proteins altered by selenium-enriched yeast play a role in cancer; α-1B-glycoprotein, transferrin, haptoglobin, transferrin, α-1 antitrypsin, and angiotensin precursor.\(^{91}\) In a nested case-control study evaluating the effect of selenium and dietary glucosinolate intake, glutathione peroxidase activity exhibited no significant association to benign hyperplasia risk among men \((n = 325)\).\(^{92}\) Proflavine, a mixture of selenium, lycopene, and saw palmetto tree berries, decreased tissue levels of total interstitial mononuclear cells, B lymphocytes, T lymphocytes, and macrophages after 6 and 3 months in patients with PIN/Atypical small acinar proliferation \((\text{ASAP})\) and BPH, respectively.\(^{93}\) Six months of selenium administration combined with lycopene and green tea extract in patients with high-grade PIN and/or ASAP resulted in the upregulation of oncopgenic microRNAs and decreased levels of miR-494, which plays a suppressive role in PCa.\(^{94}\) Inheritance of selenoprotein-associated variants were related to high-grade,\(^{95}\) susceptibility, and disease recurrence in PCa patients, but lost significance after adjustment for multiple comparisons.\(^{95,96}\)

Other studies have demonstrated a non-protective role of selenium in PCa development and progression.\(^{33,65,76,85,96}\) High selenium status in men receiving vitamin E supplements was linked to an increase in low-grade and high-grade disease development risk for PCa.\(^{85}\) Also, multi-vitamin use \((>7 \text{ times/week})\) including dietary selenium was linked to an increase in the susceptibility of PCa.\(^{46}\) Interestingly, a large elevation of 5.8-fold was related to PCa risk and dietary selenium use among fatal cases.\(^{95}\) In the SELECT trial, a strong relationship between administration of selenomethionine, a type of selenium supplement, and higher plasma levels of α-tocopherol in men potentially promoted higher PCa hazard risk ratios.\(^{67}\) Selenomethionine alone and any selenomethionine supplementation was linked to higher PCa mortality risk among men with high Gleason scores \((\geq 7)\). Moreover, high mortality risk in patients with the highest plasma levels of α-tocopherol and less than 3 year follow-up to diagnosis was associated with selenomethionine alone, α-tocopherol and selenomethionine or any selenomethionine supplementation.

**Nutraceutical efficacy in diverse populations**

To date, majority of nutraceutical chemoprevention/intervention studies have been evaluated in populations of European ancestry. Commonly, higher PCa incidence and fatalities rates have been linked to men with African ancestry. Unfortunately, only a small percentage of studies have examined the effect of nutraceutical intervention on PCa development and progression, and disease traits in diverse populations as shown in Tables 1 and 2. Thus, current clinical and epidemiological studies lack evidence on how effective nutraceutical agents are against PCa in diverse populations. Of the limited studies, majority of these reports have focused on lycopene, selenium Vitamin D, and E supplementation or their circulating plasma levels in relation to PCa risk among men of African, Asian, Latino, and Native Hawaiian descent. In a large prospective cohort study \((2015)\), selenium was related to PCa risk, but adjustment for both selenium and lycopene intake was associated with an increase in risk among Native Hawaiian, European, African, Japanese, and Latino-American men on a 1000 kcal daily diet.\(^{91}\) Selenium in African-American patients was linked to high risk of localized, advanced disease, and PCa fatality, whereas Latino men were only linked to localized and low-grade disease. Low PCa risk was associated with high legume intake \((\geq 28.2 \text{ g})\) in Latino men as well.\(^{25}\) Selenium had no effect on overall PCa risk regardless of ethnicity. However in the Selenium and Vitamin E Cancer Prevention trial, total, low-grade, and high-grade PCa incidences were higher among African-American men.\(^{97}\) Furthermore, African-American men with body mass index of 35 had a 2-fold PCa mortality risk.\(^{91}\)

The effect of carotenoid circulating patterns and intervention on PCa differs slightly in different ethnic groups. As previously mentioned, adjustment for lycopene intake combined with selenium intake in a multi-ethnic cohort was related to PCa risk, but not when taken alone.\(^{25}\) Plasma levels of cis-lutein/zeaxanthin, β-cryptoxanthin, and all-trans-lycopene inversely related to biochemical recurrence in a cohort including African-American men, but in a later study dietary lycopene and β-cryptoxanthin were inversely related to aggressive PCa disease among European and African-Americans, respectively.\(^{98}\) Moreover, higher levels of α-carotene and lycopene \((\text{cis} + \text{trans})\) were observed in European-American men compared to their African-American counterparts. African-Caribbean men had higher circulating serum levels of α-carotenes and β-carotenes and lutein/zeaxanthin, but lower lycopene/retinol levels due to their dietary intake compared to African-Americans.\(^{99}\) Moreover, high PSA levels were marginally linked to low retinol serum levels in Caribbean men. Based on the aforementioned studies, β-cryptoxanthin may have a protective role against PCa in African-American men. The data is suggestive that high α-carotene serum levels may contribute to PCa development in European-American and Caribbean men; however, due to limited epidemiological and clinical data more studies are needed.

Vitamin D deficiency in African-American men has been linked to high risk of PCa and aggressive disease.\(^{47}\) Moreover, low plasma 25(OH)D levels in African-Americans was significantly related to PCa aggressiveness, while high calcium intake elevated risk.\(^{48}\) However, high dietary Vitamin D intake has been shown to reduce aggressive PCa risk in African-Americans, but this effect was not seen in European-Americans.\(^{46}\) Levels of vitamin D metabolites, 25 (OH)D and 1,25(OH)2D, vary greatly in PCa; serum levels of 25(OH)D are lower, but prostate tissue levels of 1,25(OH)2D are higher in African-Americans compared with European-Americans.\(^{100}\) Also, vitamin D binding protein expression negatively correlated to 25(OH)D serum levels in African-Americans. Although, African-American PCa patients have low serum levels of 25(OH)D, high circulating levels of 25(OH)D \((34.27–93.20 \text{ ng/mL})\) are linked to an increase in PCa risk among Jamaican men.\(^{101}\) Taken together, the previous studies implicate a possible antagonistic effect of 25(OH)D on PCa development in Caribbean men, but protective role in African-Americans, which warrant additional studies in these populations.
Two major zinc transporters, hZIP1 and hZIP2, have been evaluated in malignant prostate tissue from European-American and African-American men.\textsuperscript{81} Very low levels of both transporters were observed in 92.8% of African-American PCa specimens. However, higher levels of zinc transporters were associated with high-grade prostate carcinoma in European-American men compared to African-Americans. In another study, miR-182 expression was higher in PCa tissue overall regardless of race, but miR-182 and miR-346 expression were inversely related to hZIP1 levels in European-American men only.\textsuperscript{62} Overexpression of miR-183, miR-96, and miR-182 reduced intracellular zinc levels and uptake in primary prostatic epithelial cells. However, no differences were observed for hZIP1 expression in African-American and European-American men, but this may be attributed to small sample size.

Furthermore, precision medicine suggests that genetic polymorphisms and socio-economic factors influence PCa susceptibility and must be taken into account for patient care and treatment of diverse populations. For example, the susceptibility of PCa patients to vitamin D supplementation has been modified by the inheritance of genetic variations in certain populations.\textsuperscript{49,101,102} In a meta-analysis of VDR genetic associations with PCa from 2006 to 2016, genetic variant VDR rs731236 was linked to an elevated risk of PCa development in relation to vitamin D supplementation in Asian-American and African-American men.\textsuperscript{101} Based on 27 case-control studies, VDR gene Fok I polymorphism was linked to higher PCa risk in men with European ancestry.\textsuperscript{49}

Also, the VDR rs11568820 variant was related to high risk of aggressive PCa in vitamin D-deficient African-Americans.\textsuperscript{29} In contrast, VDR genetic alterations in exon 4 and 8 and vitamin D intervention were associated with a protective effect against PCa risk in African-American men, whereas exons 5, 7, and 9 positively associated with disease susceptibility.\textsuperscript{53} VDR-related variants, HFE (rs9393682) and TUSC3 (rs1378033), were linked to disease progression in patients with localized tumors (post surgery) and advanced PCa (post androgen deprivation) in two independent cohorts, respectively.\textsuperscript{52} Although both circulating vitamin E plasma levels and intervention have been shown to modulate PCa risk, Vitamin E genetic polymorphisms also influence patient predisposition to develop this malignancy. During vitamin E intervention, inheritance of variants in vitamin E-related genes, SEC14L2, SOD1, and TTPA, significantly modified risk of high-grade PCa in patients.\textsuperscript{84} Two TTPA genetic variants (rs12679996, rs4606052) were linked to elevated mortality risk via inheritance of the CC genotype in high-grade PCa patients. In another study, the NKX3.1 rs11781886 variant (CC,CC + CT) genotypes combined with vitamin E or selenium treatment increased PCa risk in a cohort including African-Americans.\textsuperscript{12} Polymorphisms located near BUD13, ZNF259, and APOA5 genes play a role in vitamin E transport and metabolism and their associated variants modify PCa risk. For example, in a nested case-control study, the BUD13 rs964184 variant was linked to a decrease in PCa risk among men of European ancestry (483 cases and 542 controls).\textsuperscript{36} High lycopene levels in PCa patients (Gleason score ≤3 + 4) were linked to lower Fraction of the Genome Altered.\textsuperscript{19} Inheritance of XRCC1 rs25489 variant and high circulating α-carotene levels in the highest quartile population exerted a protective effect against high-grade PCa in men with the rs25489 GG genotype. High levels of β-carotene in carriers of the SOD3 rs699473 TC/CC genotype was associated with low risk of high-grade disease.\textsuperscript{107} Selenoprotein-associated polymorphisms were linked to PCa risk among Dana-Farber Cancer Institute patients (n = 722) with localized PCa disease.\textsuperscript{96} Dominant genetic models for TXNDR2 rs1005873 and SELENBP1 rs10788804 were significantly linked to an increase in risk of aggressive PCa. Specifically, TXNDR2 (rs1005873, rs3788310, and rs9606174) variants were linked to higher plasma selenium levels in PCa patients.

Socio-economic status among other social and geographical determinants has been also shown to be associated with elevated PCa incidence and/or mortality in several reports.\textsuperscript{104–107} An inverse relationship exists between socio-economic status and PCa incidence and mortality.\textsuperscript{104} Cancer screening and detection frequently occur earlier in European-Americans (with high socio-economic status) compared to African-Americans with the same socio-economic status. Thus, there is a delay between PCa diagnosis and treatment initiative in African-Americans relative to European-Americans. Moreover, Surveillance, Epidemiology, and End Results (SEER) data showed African-American men had a lower survival rate for surgery and radiation as well as higher incidence of metastasis. Between 2007 and 2011, the SEER registries and US census data showed health insurance had a large effect on disease outcomes for the four leading malignancies in the US.\textsuperscript{105} The most disadvantaged population (median household income $42,885) were linked to a 1.6-fold increase of distant disease and less likely to receive surgical treatment among patients diagnosed with breast, lung, and PCa relative to the high-income population (p < 0.001). These differences were observed across quintiles regardless of the insurance status.\textsuperscript{105} Cancer-specific survival was lowest among most disadvantaged patients. Socio-economic status in the VITAL cohort showed a weak association with PCa risk among men.\textsuperscript{106} However, the susceptibility of cancer-specific mortality was significantly increased among men in age-adjusted, sex-adjusted, and demographic-adjusted risk models. African-Americans residing in the Mississippi Delta suffer from higher PCa diagnoses and fatalities compared to average US incidence and mortality rates.\textsuperscript{107} This trend was observed in both rural and urban areas commonly composed of disadvantaged communities. Catchment areas, the surrounding geographic areas and populations that cancer centers service, most often include disadvantaged communities. PCa screening, diagnosis, and treatment access at major cancer centers and hospitals have been shown to differ among certain demographics. The populations serviced within or outside of catchment areas of these cancer centers are associated with lower numbers of African-Americans, Hispanics, and uninsured patients.\textsuperscript{108,109} These patterns may attribute to disadvantaged communities being medically underserved as a result of low socio-economic status or restricted accessibility to medical centers; however, additional studies are needed to validate these relationships.

CONCLUSION AND FUTURE DIRECTIONS
Recent clinical and population-based efficacy studies have shown the potential of nutraceuticals as potent anti-cancer agents; however, genetic alterations, social determinants, population/ethnic, and dosage variations modify the protective effect of these agents. Current clinical and epidemiological evidence in diverse populations has not provided sufficient data to determine whether nutraceutical intervention alone or as complement agents can address therapeutic issues associated with health disparities in PCa. Some nutraceutical agents (i.e., Vitamin D, Vitamin E, zinc, selenium) have been linked to unintended outcomes in relation to PCa susceptibility due to deficiency, and excessive use of supplements. Therefore, caution should be taken for the therapeutic application of these agents. Furthermore, some epidemiologic evidence implicate differences in PCa susceptibility in diverse populations of men based on genetic variants, socio-economic, and environmental factors. Specifically, deficient or low levels of Vitamin D and zinc may contribute to health disparities for African-Americans; however, this is still not well understood in the literature. As neo-adjuvant therapies, several nutraceutical have showed limited, but promising inhibitory effects on pathological features associated with precursor neoplastic lesions in PCa.
Most nutraceutical agents are common constituents in products or manufactured food products consumed daily by humans and have primarily exhibited non-toxic side effects. Ideally, these agents would serve as potential complement treatment options to overcome toxic side effects exerted by existing clinical therapies against aggressive and advanced PCs in diverse patient populations. Although current clinical and epidemiological evidence identified in this review does not overwhelmingly support the hypothesis of an efficacious and therapeutic role for the selected agents in humans, large efficacy studies for these agents were quite limited and lacked disproportionate populations affected by PCa to demonstrate therapeutic activity in high-risk patients. To fully assess the efficacy of these agents, comprehensive pharmacogenomic, pharmacokinetic, case-control population, and genetic studies and clinical trials are needed to determine low-response and high-response rates and to therapeutic limitations of nutraceuticals among diverse patients. Large-scale and inclusive clinical studies will aid to demonstrate whether agents are best utilized as neo-adjuvant or adjuvant therapies to address genetic and pharmacogenomic vulnerabilities in different populations and overcome health disparities. Moreover, these studies will direct the implementation and design of nutraceutical-driven precision medicine strategies to develop patient-focus therapies, reduce the burden of chemotherapy-associated toxicities, suppress disease resistance, and treat both localized and advanced-stage PCs.

ACKNOWLEDGEMENTS

Original PCa studies in our program are supported by R01 grants CA116636, CA91883, and CA102514.

AUTHOR CONTRIBUTIONS

D.R. researched, collated the literature, and composed the first draft of the manuscript including Tables 1 and 2. D.R., K.R., and R.A. contributed to the design of the manuscript and significantly edited both the manuscript and Tables 1 and 2. K.R. and R.A. cross-referenced the literature.

ADDITIONAL INFORMATION

Competing interests: The authors declare no competing interests.

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

REFERENCES

1. ACS Cancer Facts and Figures 2018. (American Cancer Society, Atlanta, GA, 2018).
2. Ting, H., Deep, G., Agarwal, C. & Agarwal, R. The strategies to control prostate cancer by chemoprevention approaches. Mutat. Res. 760, 1–15 (2014).
3. Bijak, M., Silybin, a major bioactive component of milk thistle (Silybum marianum L. Gaertn.-) chemistry, bioavailability, and metabolism. Molecules 22, 1942 (2017).
4. Flagg, T. W. et al. A study of high-dose oral silybin-phytosome followed by prostatectomy in patients with localized prostate cancer. Prostate 70, 846–855 (2010).
5. Nassiri-Asl, M., & Hosseinzadeh, H. Review of the pharmacological effects of Vitis vinifera (Grape) and its bioactive compounds. Phytother. Res. 23, 1197–1204 (2009).
6. Kaur, M., Agarwal, C. & Agarwal, R. Anticancer and cancer chemopreventive potential of grape seed extract and other grape-based products. J. Nutr. 139, 18065–18125 (2009).
7. Brasky, T. M. et al. Specialty supplements and prostate cancer risk in the Vita- mins and Lifestyle (VITAL) cohort. Nutr. Cancer 63, 573–582 (2011).
8. Capurro, C. & Vendemiale, G. The Mediterranean diet reduces the risk and mortality of prostate cancer: a narrative review. Front. Nutr. 4, 38 (2017).
9. Key, T. J. et al. Carotenoids, retinol, tocopherols, and prostate cancer risk: pooled analysis of 15 studies. Am. J. Clin. Nutr. 102, 1142–1157 (2015).
10. Nordstrom, T. et al. Associations between circulating carotenoids, genotypic instability and the risk of high-grade prostate cancer. Prostate 76, 339–348 (2016).
11. Paur, I. et al. Tomato-based randomized controlled trial in prostate cancer patients: effect on PSA. Clin. Nutr. 36, 672–679 (2017).
12. Wang, Y., Jacobs, E. J., Newton, C. C. & McCullough, M. L. Lycopene, tomato products and prostate cancer-specific mortality among men diagnosed with nonmetastatic prostate cancer in the Cancer Prevention Study II Nutrition Cohort. Int. J. Cancer 138, 2846–2855 (2016).
13. Graff, R. E. et al. Dietary lycopene intake and risk of prostate cancer defined by ERG protein expression. Am. J. Clin. Nutr. 103, 851–860 (2016).
14. Antwi, S. O. et al. Plasma carotenoids and tocopherols in relation to prostate-specific antigen (PSA) levels among men with biochemical recurrence of prostate cancer. Cancer Epidemiol. 39, 752–762 (2015).
15. Beydoun, H. A., Shroff, M. R., Mohan, R. & Beydoun, M. A. Associations of serum vitamin A and carotenoid levels with markers of prostate cancer detection among US men. Cancer Causes Control 22, 1483–1495 (2011).
16. Zhang, Y., Coogan, P., Palmer, J. R., Strom, B. L. & Rosenberg, L. Vitamin and mineral use and risk of prostate cancer: the case-control surveillance study. Cancer Causes Control 20, 691–699 (2008).
17. Kristal, A. R. et al. Dietary patterns, supplement use, and the risk of symptomatic benign prostatic hyperplasia: results from the prostate cancer prevention trial. Am. J. Epidemiol. 167, 925–934 (2008).
18. Meyer, F. et al. Antioxidant vitamin and mineral supplementation and prostate cancer prevention in the SU.VI.MAX trial. Int. J. Cancer 116, 182–186 (2005).
19. Van Hoang, D., Pham, N. M., Lee, A. H., Tran, D. N. & BInns, C. W. Dietary carotenoids and prostate cancer risk: a case-control study from Vietnam. Nutrients. https://doi.org/10.3390/nu10010070 (2018).
20. Venkitaraman, R. et al. Serum micronutrient and antioxidant levels at baseline and the natural history of men with localized prostate cancer on active surveillance. Tumour Biol. 31, 97–102 (2010).
21. Beilby, J., Ambrosini, G. L., Rossi, E., de Klerk, N. H. & Musk, A. W. Serum levels of folate, lycopene, beta-carotene, retinol and vitamin E and prostate cancer risk. Eur. J. Clin. Nutr. 64, 1235–1238 (2010).
22. Lane, J. A. et al. Prostate cancer risk related to foods, food groups, macronutrients and micronutrients derived from the UK Dietary Cohort Consortium food diaries. Eur. J. Clin. Nutr. 71, 274–283 (2017).
23. Cui, K. et al. Chemoprevention of prostate cancer in men with high-grade prostatic intraepithelial neoplasia (HPGNN): a systematic review and adjusted indirect treatment comparison. Oncotarget 8, 36674–36684 (2017).
24. Graff, R. E. et al. Circulating antioxidant levels and risk of prostate cancer by TMPRSS2:ERG. Prostate 77, 647–653 (2017).
25. Park, S. Y. et al. Racial/ethnic differences in lifestyle-related factors and prostate cancer risk: the Multiethnic Cohort Study. Cancer Causes Control 26, 1507–1515 (2015).
26. Virtamo, J. et al. Effects of alpha-tocopherol and beta-carotene supplementation on cancer incidence and mortality: 18-year postintervention follow-up of the Alpha-tocopherol, Beta-carotene Cancer Prevention Study. Int. J. Cancer 135, 178–185 (2014).
27. Morgia, G. et al. Association between selenium and lycopene supplementation and incidence of prostate cancer: results from the post-hoc analysis of the procobum trial. Phytomedicine 34, 1–5 (2017).
28. Gann, P. H. et al. A phase II randomized trial of lycopene-rich tomato extract among men with high-grade prostate intraepithelial neoplasia. Nutr. Cancer 67, 1104–1112 (2015).
29. Magbanua, M. J. et al. Gene expression and biological pathways in tissue of men with prostate cancer in a randomized clinical trial of lycopene and fish oil supplementation. PLos ONE 6, e24004 (2011).
30. Mahmoud, A. M., Yang, W. & Bosland, M. C. Soy isoflavones and prostate cancer: a review of molecular mechanisms. J. Steroid Biochem. Mol. Biol. 140, 116–132 (2014).
31. Ahn-Jarvis, J. H. et al. Isoflavone pharmacokinetics and metabolism after consumption of a standardized soy and almond bread in men with asymptomatic prostate cancer. Cancer Prev. Res. 8, 1045–1054 (2015).
32. Lesinski, G. B. et al. Consumption of soy isoflavone enriched bread in men with prostate cancer is associated with reduced proinflammatory cytokines and immunosuppressive cells. Cancer Prev. Res. 8, 1036–1044 (2015).
33. Wu, Y. et al. Plasma genistein and risk of prostate cancer in Chinese population. Int. Urol. Nephrol. 47, 965–970 (2015).
34. Zhang, H. Y. et al. Isoflavones and prostate cancer: a review of some critical issues. Chin. Med. J. 129, 341–347 (2016).
35. Szwarc, S. et al. Inhibition of prostaglandin synthesis and actions by genistein in human prostate cancer cells and by soy isoflavones in prostate cancer patients. Int. J. Cancer 124, 2050–2059 (2009).
36. Hamilton-Reeves, J. M., Rebello, S. A., Thomas, W., Kurzer, M. S. & Slaton, J. W. Effects of soy protein isolate consumption on prostate cancer biomarkers in men with HPGNN, ASAP, and low-grade prostate cancer. Nutr. Cancer 60, 7–13 (2008).
37. Sur, S. & Panda, C. K. Molecular aspects of cancer chemopreventive and therapeutic efficacies of tea and tea polyphenols. *Nutrition* 43–44, 8–15 (2017).

38. McLarty, J. et al. Tea polyphenols decrease serum levels of prostate-specific antigen, hepatocyte growth factor, and vascular endothelial growth factor in prostate cancer patients and inhibit production of hepatocyte growth factor and vascular endothelial growth factor in vitro. *Cancer Prev. Res.* 2, 673–682 (2009).

39. Thomas, R., Williams, M., Sharma, H., Chaudry, A. & Bellamy, P. A double-blind, placebo-controlled randomised trial evaluating the effect of a polyphenol-rich whole food supplement on PSA progression in men with prostate cancer—the U.K. NCNR Pomi-T study. *Prostate Cancer Prostatic Dis.* 17, 180–186 (2014).

40. Henning, S. M. et al. Randomized clinical trial of brewed green and black tea in men with prostate cancer prior to prostatectomy. *Prostate* 75, 550–559 (2015).

41. Nguyen, M. M. et al. Randomized, double-blind, placebo-controlled trial of polyphenol E in prostate cancer patients before prostatectomy: evaluation of potential chemopreventive activities. *Cancer Prev. Res.* 5, 290–298 (2012).

42. Henning, S. M. et al. Tea polyphenols and theaflavins are present in prostate tissue of humans and mice after green and black tea consumption. *J. Nutr.* 136, 1839–1843 (2006).

43. Gontero, P. et al. A randomized double-blind placebo controlled phase II study on clinical and molecular effects of dietary supplements in men with pre-cancerous prostate lesions. Chemoprevention or “chemopromotion”? *Prostate* 75, 1177–1186 (2015).

44. Bandera, M., Borcic, S., Martin-Nunez, G., Tinhones, F. J. & Macias-Gonzalez, M. The role of vitamin D and VDR in carcinogenesis: through epide-miology and basic sciences. *J. Steroid Biochem. Mol. Biol.* 167, 203–218 (2017).

45. Meyer, H. E. et al. Long term association between serum 25-hydroxyvitamin D and mortality in a cohort of 4379 men. *PLoS ONE* 11, e0151441 (2016).

46. Batai, K. et al. Race and BMI modify associations of calcium and vitamin D intake with prostate cancer. *Prostate Cancer Prostatic Dis.* 17, 81–87 (2014).

47. Henning, S. M. et al. Randomized clinical trial of brewed green and black tea in men with prostate cancer prior to prostatectomy. *Prostate* 75, 550–559 (2015).

48. Brandstedt, J., Almqvist, M., Manjer, J. & Malm, J. Vitamin D, PTH, and calcium and risk mortality: state of the science, gaps, and challenges. *Epidemiol. Rev.* 39, 28–48 (2017).

49. Pandolfi, F., Franza, L., Mandolini, C. & Conti, P. Immune modulation by vitamin D: special emphasis on its role in prevention and treatment of cancer. *Clin. Ther.* 39, 884–893 (2017).

50. Brandstedt, J., Almqvist, M., Manjer, J. & Malm, J. Vitamin D, PTH, and calcium in relation to survival following prostate cancer. *Cancer Causes Control* 27, 669–677 (2016).

51. Jannard, D. et al. Phase IIa, randomized placebo-controlled trial of single high dose cholecalciferol (vitamin D3) and daily Genistein (G-2535) versus double placebo in men with early stage prostate cancer undergoing prostatectomy. *Am. J. Clin. Exp. Urol.* 4, 17–27 (2016).

52. Medioni, J. et al. Phase I safety and pharmacodynamic of inecalcitol, a novel VDR agonist with docetaxel in metastatic castration-resistant prostate cancer patients. *Clin. Cancer Res.* 20, 4471–4477 (2014).

53. Xie, D. D. et al. Low vitamin D status is associated with in看cal risk of fatal prostate cancer in the National Cancer Institute Breast and Prostate Cancer Cohort Consortium. *Cancer Prev. Res.* 11, 1186–1197 (2018).

54. Sapota, A. et al. The bioavailability of different zinc compounds used as human dietary supplements in rat prostate: a comparative study. *Biomaterials* 27, 495–505 (2014).

55. van Wijngaarden, E., Singer, E. A. & Palapattu, G. S. Prostate-specific antigen levels in relation to calcium exposure and zinc intake: results from the 2001-2002 National Health and Nutrition Examination Survey. *Prostate* 68, 122–128 (2008).

56. Gonzalez, A., Peters, U., Lampe, J. W. & White, E. Zinc intake from supplements and diet and prostate cancer. *Nutr. Cancer* 61, 206–215 (2009).

57. Kristal, A. R. et al. Functional variant in NKK31 associated with prostate cancer risk in the Selenium and Vitamin E Cancer Prevention Trial (SELECT). *Cancer Prev. Res.* 7, 950–957 (2014).

58. Mondl, A. M. et al. Metabolomic analysis of prostate cancer risk in a prospective cohort: the alpha-tocopherol, beta-carotene cancer prevention (ATBC) study. *Int. J. Cancer* 137, 2124–2132 (2015).

59. Bauer, S. R. et al. Antioxidant and vitamin E transport genes and risk of high-grade prostate cancer and prostate cancer recurrence. *Prostate* 73, 1786–1795 (2013).

60. Hsueh, Y. M. et al. Levels of plasma selenium and urinary total arsenic interact to increase risk of prostate cancer. *Nutrients* 7, 10648–10666 (2015).

61. Major, J. M. et al. Genetic variants reflecting higher vitamin e status in men are associated with reduced risk of prostate cancer. *J. Nutr.* 144, 729–733 (2014).

62. Sapota, A. et al. The bioavailability of different zinc compounds used as human dietary supplements in rat prostate: a comparative study. *Biomaterials* 27, 495–505 (2014).

63. Singh, B. P. et al. Status and interrelationship of zinc, copper, iron, calcium and magnesium in rat prostate: a comparative study. *Appl. Immunohistochem. Mol. Morphol.* 11, 253–260 (2003).

64. Mihelic, B. L. et al. miR-183-96-182 cluster is overexpressed in prostate tissue and regulates zinc homeostasis in prostate cells. *J. Biol. Chem.* 286, 44503–44511 (2011).

65. Mandiar, D., Rossi, R. E., Pericleous, M., Whyard, T. & Caplin, M. E. Prostate cancer and the influence of dietary factors and supplements: a systematic review. *Nutr. Metab.* 11, 30 (2014).

66. Singh, B. P. et al. Status and interrelationship of zinc, copper, iron, calcium and selenium in prostate cancer. *Indian J. Clin. Biochem.* 31, 50–56 (2016).

67. Lu, J. et al. Cancer chemoprevention research with selenium in the post-SELECT era: promises and challenges. *Nutr. Cancer* 68, 1–17 (2017).

68. Cui, Z., Liu, D., Liu, C. & Liu, G. Serum selenium levels and prostate cancer risk: a MOOSE-compliant meta-analysis. *Medicine* 96, e5944 (2017).

69. Husey, Y. M. et al. Levels of plasma selenium and urinary total arsenic interact to affect the risk for prostate cancer. *Food Chem. Toxicol.* 107, 167–175 (2017).

70. Allen, N. E. et al. Selenium and prostate cancer: analysis of individual participant data from fifteen prospective studies. *J. Natl Cancer Inst.* https://doi.org/10.1093/jnci/dwj153 (2016).
20

108. Su, S. C. et al. Spatial analyses identify the geographic source of patients at a National Cancer Institute Comprehensive Cancer Center. Clin. Cancer Res. 16, 1065–1072 (2010).

109. Wang, A. & Wheeler, D. C. Catchment area analysis using bayesian regression modeling. Cancer Inform. 14, 71–79 (2015).

110. Hamilton-Reeves, J. M. et al. Short-term soy isoflavone intervention in patients with localized prostate cancer: a randomized, double-blind, placebo-controlled trial. PLoS ONE 8, e68331 (2013).

111. de Vere White, R. W. et al. Effects of a high dose, aglycone-rich soy extract on prostate-specific antigen and serum isoflavone concentrations in men with localized prostate cancer. Nutr. Cancer 62, 1036–1043 (2010).

112. Lazarevic, B. et al. Efficacy and safety of short-term genistein intervention in patients with localized prostate cancer prior to radical prostatectomy: a randomized, placebo-controlled, double-blind Phase 2 clinical trial. Nutr. Cancer 63, 889–898 (2011).

113. Ahmad, I. U. et al. Soy isoflavones in conjunction with radiation therapy in patients with prostate cancer. Nutr. Cancer 62, 996–1000 (2010).

114. Napora, J. K. et al. High-dose isoflavones do not improve metabolic and inflammatory parameters in androgen-deprived men with prostate cancer. J. Androl. 32, 40–48 (2011).

115. Kwan, W., Duncan, G., Van Patten, C., Liu, M. & Lim, J. A phase II trial of a soy beverage for subjects without clinical disease with rising prostate-specific antigen after radical radiation for prostate cancer. Nutr. Cancer 62, 198–207 (2010).

116. Ide, H. et al. Combined inhibitory effects of soy isoflavones and curcumin on the production of prostate-specific antigen. Prostate 70, 1127–1133 (2010).

117. Scher, H. I. et al. Randomized, open-label phase III trial of docetaxel plus high-dose calcitriol versus docetaxel plus prednisone for patients with castration-resistant prostate cancer. J. Clin. Oncol. 29, 2191–2198 (2011).

118. Kristal, A. R. et al. Plasma vitamin D and prostate cancer risk: results from the selenium and vitamin E prevention trial. Cancer Epidemiol. Biomark. Prev. 23, 1494–1504 (2014).

119. Mondul, A. M. et al. Serum metabolomic response to long-term supplementation with all-rac-alpha-tocopheryl acetate in a randomized controlled trial. J. Nutr. Metab. 2016, 6158436 (2016).

120. Marshall, J. R. et al. Phase III trial of selenium to prevent prostate cancer in men with high-grade prostatic intraepithelial neoplasia: SWOG S9917. Cancer Prev. Res. 4, 1761–1769 (2011).