Cardiovascular disease (CVD) has been the number one cause of death in the U.S. for 114 of the last 115 years. Lifestyle factors that promote CVD also appear to increase prostate cancer risk and those that reduce CVD risk also appear to reduce the risk of prostate cancer. The largest randomized trials utilizing dietary supplements or pharmacologic agents for prostate cancer prevention (Selenium and Vitamin E Cancer Prevention Trial [SELECT]) have also shed light on the problems and future solutions in this area. Dietary supplements that have not been found to be CVD protective, such as selenium and Vitamin E have not been found to be prostate protective. In addition, over exposure to specific anti-oxidants in nutritionally replete populations may be encouraging cancer growth. Future trials of dietary supplements to prevent prostate cancer could be problematic because by the time a definitive trial is initiated the participants will no longer be “deficient” in the nutrient being tested, which arguably occurred in the SELECT trial. It is also interesting that statins, aspirin, and/or metformin (S.A.M.) are 3 generic, low-cost, heart healthy agents derived from natural sources with separate mechanism of actions, which all appear to have the best benefit to risk ratio compared to any other agent available for prostate cancer prevention, especially aggressive disease, or as an ancillary agent (s) to conventional cancer treatment. It is time to focus on the forest over the trees and recommend proven CVD protective measures for men concerned about their risk of prostate cancer.

**Keywords:** aspirin; diet; lifestyle; metformin; prostate cancer; statins

**INTRODUCTION**

Male health issues should be triaged before recommending and construing any ideal prostate cancer prevention program. Reiterating and emphasizing the primary causes of past and current morbidity and mortality allows for an easier understanding of lifestyle, supplement or pharmacologic additions or deletions in a primary prevention setting. This advice needs to be simple, logical and practical for the patient as well as a clinician. When overall health concerns are triaged, it will be easier to understand and advocate for the ideal prostate cancer prevention program, which is supported by the phrase heart healthy equals prostate healthy.¹

Cardiovascular disease (CVD) is the number one overall cause of mortality in the United Stated (U.S.) and in other industrialized countries.²⁻³ Cancer is the second leading cause of death in the U.S. and in most developed countries, and it expected to potentially mirror the number of deaths from CVD in the near future. CVD has been the number one cause of male death in the U.S. for approximately 114 out of the last 115 years, only surpassed for a single year by the influenza pandemic in 1918. Ponder this for a moment in tangible terms, if a college football team was number 1 in the U.S. for virtually 100 years, would arguably be such awareness and attention paid to this streak in and out of the sports world because it would be so remarkable, dominant, and unprecedented. However, how aware or tangible would such a dynasty be to the public and clinicians if it involved much more than just football, but individual risk of morbidity and mortality?

Even if cancer becomes the primary cause of death, the majority of what is known concerning lifestyle and dietary change for CVD prevention directly appears to apply to cancer prevention and most other prevalent diseases, such as diabetes.⁴ For example, it should be of interest to healthcare professionals and patients that one of the most significant reductions in early morbidity and mortality rates in U.S. history for CVD and cancer was via a common behavioral/lifestyle change, smoking cessation, that simultaneously impacted both diseases.⁵⁻⁷ The opposite side of this pendulum is also true, for example, the global prevalence of tobacco use remains high and is still the largest preventable cause of death from CVD and cancer. And, obesity for example not only continues to increase the risk of CVD, but numerous cancers, diabetes, early morbidity and mortality, and it may erase the advances in the declines previously observed in early CVD and cancer mortality from smoking cessation.⁸⁻¹⁰

Men have a consistently lower life expectancy in the U.S. and in most countries around the world, and have a higher morbidity and mortality from heart disease, hypertension, cancer, and diabetes.¹¹
Yet, it must be reiterated that heart healthy changes are tantamount to overall men’s health improvements regardless of the part of the human anatomy that is receiving attention, including the penis and the prostate.13–15 Heart healthy changes need to be advocated to men concerned about prostate cancer because it places probability and the overall research into perspective. Triaging preventive medicine for men’s health is providing probability based advice via evidence-based medicine and can impact all-cause mortality as well as potentially prevent prostate cancer.  

UNAPPRECIATED LESSONS FROM PROSTATE CANCER PRIMARY PREVENTION TRIALS UTILIZING A PHARMACOLOGIC AGENT

One of the more fascinating features of large randomized clinical trials in our opinion, especially for primary prevention, is that they appear to mirror the current health status and risk issues of not only the subjects being tested but perhaps the general population. The largest, most recent, and arguably the best designed U.S. and worldwide pharmaceutical-based cancer primary prevention trials for the prevention of prostate cancer exemplify the urgency for a different perspective. For example, results of the Prostate Cancer Prevention Trial (PCPT) seem to have garnered attention and controversy regarding the use of finasteride daily versus placebo to reduce the risk of prostate cancer.13–16 The debate over finasteride abounds, but another observation from this trial has not received adequate exposure in the medical literature. Over 18,000 healthy men were included in this randomized trial, and 5 men died from prostate cancer in the finasteride arm and 5 men died of prostate cancer in the placebo arm. However, 1123 men in total died during this primary prevention trial.3 Thus, prostate cancer was responsible for approximately <1% of the deaths, while the majority of the overall causes of mortality were from CVD and other causes.13,24 Additionally, the mean body mass index (BMI), systolic blood pressure and total and high-density lipoprotein cholesterol were the following: 27–28 (50% overweight, and approximately 25% obese), 138–140 mm Hg (prehypertensive), 212 mg dL−1 and 42–43 mg dL−1 (dyslipidemia or at-risk). Despite 85% of men with no history of CVD approximately 50% of the men reported some level of erectile dysfunction.25

Interestingly, the more recent international dutasteride prevention trial known as REDuction by DUtasteride of Prostate Cancer Events (REDUCE) had somewhat similar issues to the North American PCPT in terms of overt controversies,16–21 but what was not questioned, discussed, or even debated was the BMI and several other abnormal CVD parameter issues mentioned earlier were similar in the two trials. For example, on average men in REDUCE were overweight (BMI of 27–28).23 There were 8231 men randomized and after the 4 years trial in this group of high-risk men there were 147 total deaths, primarily from cardiovascular events and none from prostate cancer. Of further note, men in the placebo arm of PCPT with low cholesterol (<200 mg dL−1) had a 59% (P = 0.02) apparent reduction in risk of being diagnosed with aggressive prostate cancer (Gleason 8–10) compared to men with high cholesterol (>200 mg dL−1),22 and men with coronary artery disease at baseline in REDUCE were found to have a significantly higher risk of a prostate cancer diagnosis, and this included low-grade (odds ratio [OR] = 1.34, P = 0.02) and high-grade cancer (OR = 1.34, P = 0.09).23 These observations do not intend to belittle prostate cancer or these trials utilizing a specific chemoprevention agent, but again it places the overall risk of morbidity and mortality in a more proper perspective. Men inquiring about a potential preventative prostate cancer prevention need to be reminded that the number 1 risk to them in general is CVD and in both clinical trials the researchers found that heart health was tantamount to prostate health.  

UNAPPRECIATED LESSONS FROM NOTABLE DIETARY SUPPLEMENT CANCER PREVENTION TRIALS

The largest male health dietary supplement clinical trial to prevent prostate cancer was the Selenium and Vitamin E Cancer Prevention Trial (SELECT).24 It randomized over 35,000 men into four groups: high-dose Vitamin E (400 IU per day), high-dose selenium (200 mcg per day), Vitamin E and selenium, or placebo. Full recruitment for the trial was achieved ahead of schedule. Thus it seemed that participants and health care professionals were equally enthusiastic to test the hypothesis that high-dose anti-oxidant supplementation could prevent prostate cancer. Yet, the trial was terminated early and recently, after a median of 5.5 years due to a lack of efficacy, although at the time a nonsignificant (P = 0.06) increase risk of nonaggressive prostate cancer in the Vitamin E arm (63% Gleason ≤ 6, 94% Gleason ≤ 7, and similar percentage of Gleason 8–10 disease vs placebo), and type 2 diabetes in the selenium group (P = 0.16) were observed.

Still, and as a credit to the SELECT research team, participant follow-up continued (54,464 added person-years), which provided more clarity of the further health impacts after the discontinuation of these agents.23 What was demonstrated recently in this follow-up period was an issue. A significant (P = 0.008; hazard ratio [HR] = 1.17) increased risk of prostate cancer was observed in the Vitamin E group, and the increased risk with this individual supplement began to emerge after only 3 years, and was found to be consistent for low- and high-grade disease types. Still, the increased risk was primarily from low-grade disease because Gleason ≥ 7, although higher in number was not significantly different from placebo. Gleason 7 or higher disease was greater for the three intervention arms compared to placebo, but did not reach statistical significance. The HR and P value for Gleason 7 and higher disease compared to placebo was 1.16 (P = 0.20), 1.21 (P = 0.11), and 1.23 (P = 0.08) for Vitamin E, selenium, and the combination.

The negative observations from SELECT cannot be simply construed by increased biopsy rates or bias, but suggest that the high-dose dietary supplements themselves were the culprits, and the confidence intervals to support this thought have continuously narrowed over time.23 Other findings from secondary endpoint analyses included other cancers and cardiovascular events, but did not find statistical differences compared with placebo. This is a modicum of good news in light of such negativity from utilizing what many would have initially perceived as potentially benign over the counter (OTC) agents. Still, what should receive more attention was the finding that CVD events and deaths represented the primary cause of morbidity and mortality overall in this trial in all 4 treatment arms. For example, there were over 4200 cardiovascular events and over 500 CVD deaths that occurred compared with 1750 prostate cancers diagnosed and 1 death from prostate cancer. There were 3363 cancers diagnosed overall (including prostate) and 476 deaths from cancer, which again emphasizes the need for future chemoprevention agents or lifestyle interventions to harbor activity against CVD and cancer, because the global burden of cancer is beginning to compete with CVD,26 again as reflected in the SELECT trial.

In our opinion, the results of SELECT could have been even more disconcerting over time if the interventions were continued. Still, even if any of these interventions would have prevented prostate cancer it is questionable whether they would have provided a tangible overall clinical advance in medicine. The issue that plagued high-dose
Vitamin E and selenium supplements from past clinical trials was the dearth of evidence or at times negative impact these supplements had on overall mortality, and on CVD.27–32 Neither type/form of Vitamin E (synthetic or natural source), or even frequency of utilization of this supplement in higher dosages would have arguably provided any difference in the SELECT trial, especially in regards to CVD and probably cancer outcomes.29,30,33–36 For example, one notable trial (HOPE TOO) actually found a significantly higher rate of heart failure with a naturally derived Vitamin E supplement.30 Another large randomized trial of Vitamin E and prostate cancer risk in healthy men, the Physicians Health Study II (PHSII), found no impact of 400 IU of Vitamin E every other day compared to placebo,35 but a significant increase risk of hemorrhagic stroke was observed.34 Furthermore, the Alpha-Tocopherol, Beta Carotene (ATBC) trial of over 29,000 men demonstrated a notable 35% risk reduction of prostate cancer risk with a Vitamin E supplement from a secondary endpoint, and provided some impetus for the design and initiation of SELECT.35 Yet, the dosage utilized in the ATBC was only 50 IU, approximately 8 times lower compared to SELECT, and a higher rate of hemorrhagic stroke was also found in ATBC. The number one and two overall cause of death during ATBC and at postintervention follow-up was ischemic heart disease and lung cancer.35,36 Men in ATBC were chronic 36 years on average smokers, and continuous tobacco users are at a higher risk of diverse nutrient deficiencies including Vitamin E.37 Less than 10% of SELECT participants were current smokers,38 and one wonders the outcome of this trial or others had a lower-dose been utilized in a more representative population of generally healthy men? If a little might be good then more is better? Isn’t this one pervasive stereotype applied to patients that utilize a multitude of non- evidence-based dietary supplements? Healthy and primarily non- or former smoking men (85% of the participants) from a unique randomized trial (SUVIMAX) utilizing far lower-doses of Vitamin E (30 IU) and several other supplement ingredients demonstrated the potential for significant overall benefit and prostate cancer prevention, but also potential harm (increase in total prostate cancer risk) for men with higher baseline prostate-specific antigen (PSA) levels.38,39

What about selenium dietary supplements? Again, the impact of high-dose selenium supplements on heart and overall health from past studies were arguably as concerning as past Vitamin E data especially in those replete with this nutrient,40,41 and included a potential significant increased risk of type 2 diabetes and nonmelanoma skin cancer recurrence.42,43 Interestingly, the increased risk of skin cancer recurrence was the final conclusion of the primary endpoint of the randomized selenium supplement U.S. trial (Nutritional Prevention of Cancer or NPC) initiated in the 1980s and completed in the 1990s.42,44 It was the NPC trial secondary endpoint results, for example, the lower rate of prostate cancer that were the impetus for the design and initiation of the SELECT trial.

Additionally, it is plausible that the SELECT researchers or even future investigators testing individual supplements for cancer or CVD prevention will not be capable of initiating a nutritionally uncontaminated clinical trial by the time of randomization. This is due to a novel situation continuously occurring with the ongoing U.S. and global popularity of functional foods and supplements45,46 and it has been referred to by one of the authors as the “over-anti-oxidation of the population.”47 In other words, currently if any nutrient appears to impact some common condition without adequate long-term research no entity exists to block the ability of nutritional commercial products in the U.S. to add more and more of these nutrients to everything from multivitamins to protein bars or energy drinks to water with added vitamins and minerals! For example, baseline serum selenium status in SELECT was actually 22 points higher (135 ng ml−1 vs 113 ng ml−1) compared to notable NPC trial completed in the 1990s in the U.S. NPC participants who were selenium deficient eventually experienced a potential reduced prostate cancer risk, but a higher rate of cancer occurred in a small group of individuals with repletion of baseline selenium levels.34,48,49 Most SELECT participants were already selenium sufficient at baseline and were recruited from all over the U.S. including some of the same geographical areas as the NPC trial participants only more than a decade later. How could selenium blood levels increase so substantially within 10–15 years between the NPC and then SELECT trial recruitment period? Arguably, we believe the increased addition of selenium (and Vitamin E) in foods, beverages, and supplements; increased overall consumption of these functional foods and calories, and the reduction in smoking since the 1990s all greatly assisted in the normalization of selenium. For example, locating a multivitamin with selenium in the 1980s or 1990s was difficult and today finding any multivitamin without selenium is almost impossible. Approximately, 30% of NPC versus 8% of SELECT participants were current smokers (smokers have lower selenium levels).

Some publications have claimed that the reason anti-oxidant trials have been neutral or negative overall in medicine is because nutritionally sufficient rather than insufficient or deficient individuals are subjects of these studies.50 However, multiple years are required to propose, fund, design, recruit and initiate any large-scale nutritional clinical trials. Thus, the initially depleted participants being tested will eventually be replete with the interventions being utilized before the trial officially commences. This will represent a challenge to any further nutrient trial in industrialized countries, and perhaps this is why other supplements such as omega-3 or Vitamin D supplements for example have not been found to have dramatic impacts in other areas of medicine from recent trials or reviews.51,52

Utilizing high-dose supplements in an already replete population could result in the nutrient in question to function as a pro-oxidant or disease initiator and promoter rather than an anti-oxidant. This is what could have occurred in the case of Vitamin E,53 and with selenium,54 or with another nutrient such as folic acid which has already been observed in multiple randomized clinical trials to be a potential prostate cancer risk factor in excessive dosages from supplements.55

MULTIVITAMINS AND OTHER DIETARY SUPPLEMENTS FOR PREVENTION

The future of dietary supplement research and cancer should arguably revolve around testing lower dosages to ensure safety first and potential efficacy, or simply test supplements for specific conditions rather than prevention itself. Interestingly, the result of the first major randomized trial of multivitamins versus placebo was recently published and the primary endpoints were total cancer incidence and cardiovascular events. The PHSII found a significant (P = 0.04) 8% cancer reduction in total cancer incidence compared to placebo in a healthy group of subjects 50 years or older (n = 14 641, 11.2 years of follow-up).56 However, a larger nonsignificant 18% reduction was found for men age 70 and over at baseline and those with a history of cancer (~27%), but no benefit was found for those with a parental history of cancer. Current smokers (~4% of the participants) appeared to receive a large benefit (~28%) compared to former and never smokers. There was no impact on prostate cancer incidence or death (HR = 0.98 and 0.91), but men with a baseline history of cancer had a 44% nonsignificant (P = 0.07) reduction in total prostate cancer risk versus placebo. Overall, it is still impressive that the low-dose multivitamin with a
similar side effect to placebo significantly and modestly reduced total cancer incidence in a group of primarily healthy men. For example, further sub-group analysis found that men consuming 7 or more fruits and vegetables per day benefited as much as those that consumed < 4 servings a day, and those with a normal BMI benefited as much as overweight or obese men. There was no increased or decreased risk of this multivitamin on cardiovascular events, which is reassuring. Yet, fatal myocardial infarction (a secondary endpoint) was reduced by 39% \( (P = 0.05) \) in the multivitamin group, but especially in those men without a baseline history of CVD \( (-44\%, \ P = 0.03) \).\textsuperscript{37} It is interesting that the original Centrum Silver utilized during this trial from 1997 to 2011 is not the OTC product offered to consumers currently because over time these nutritional formulations appear to change based on some science and marketing demand. Therefore, if one is impressed by this data, a single child's multivitamin could be recommended for an adult because this dosage appears to be similar to an older Centrum Silver or the patient should just consume the newest Centrum Silver or something close to the formula which is detailed in the clinical trial publication.\textsuperscript{38}

Regardless, it should be reiterated that more is not better in terms of multivitamins. Some of the largest past prospective epidemiologic studies are suggesting a higher rate of aggressive and fatal prostate cancer when consuming more than 1 multivitamin a day with even further increasing risk when other high-dose individual supplements are also utilized (selenium, Vitamin E and zinc).\textsuperscript{39} Men with a family history of prostate cancer experienced the largest and most significant elevated risks of this condition. Other large male observational studies have found somewhat similar results with multivitamins and some individual supplements.\textsuperscript{39-41} Multivitamins are also replete in our experience with higher-doses of B-vitamins such as B12 and folic acid, which have also recently been found to potentially have no impact on health or increase the risk of total prostate cancer incidence from the largest and most recent meta-analysis of clinical trials.\textsuperscript{42,43} Since there is no consistent suggestion of benefit with a greater intake of multivitamins or any other vitamin or mineral in supplement form, and since there is a suggestion of either no impact or serious harm it would be prudent to “first do no harm” and wait for more clarity from additional clinical studies.\textsuperscript{44}

Vitamin D in high-doses may have some similar issues to Vitamin E or selenium. The tendency for clinicians to want to recommend more Vitamin D and patients to ingest more of this supplement is concerning. In the area of prostate cancer prevention Vitamin D has not been impressive. Several epidemiologic studies have found either no impact or a potential increased risk of aggressive prostate cancer or total cancer at higher 25-OH Vitamin D blood levels.\textsuperscript{45,46} Vitamin D is important for bone health, but the amount needed has been embellished and exaggerated in our opinion. Vitamin D tends to function more like a hormone, which is why caution should be followed because the potential for a U or even J-shaped risk curve does exist for male health in general.\textsuperscript{47} One of the largest and longest randomized trials in elderly women found that excessively high blood levels of Vitamin D from high-dose supplementation compared to placebo was actually associated with an increased risk of falls and fractures.\textsuperscript{48} The normal level of Vitamin D (25-OH) could be 30–40 ng ml\(^{-1}\) based on benefit versus risk philosophy and expert opinion from a review of past clinical trials accessing multiple outcomes.\textsuperscript{49} Yet, even Vitamin D blood tests have a history of uncertainty based on the assay utilized.\textsuperscript{50,51} Monitoring Vitamin D in men, especially higher risk bone loss patients with Vitamin D deficiency, for example men on androgen deprivation therapy (ADT) for prostate cancer may be more appropriate.\textsuperscript{52} For prostate cancer prevention the Vitamin D test may provide more harm than good until more clinical endpoints are followed in healthy individuals and cost is not such an issue.\textsuperscript{53} The latest Institute of Medicine report should also be a reminder that despite the perception, the recommended intakes of Vitamin D have only increased by 200 IU (5 mcg) in most groups and Vitamin D supplements have the potential to increase the risk of hypercalcemia and nephrolithiasis.\textsuperscript{54}

Clinicians need to also remind patients that Vitamin D blood levels may simply be a marker of healthy behavior. A lean man, with a low cholesterol that consumes fish and exercises regularly is more likely to have a higher blood level of Vitamin D compared to a physically inactive overweight or obese man with a high cholesterol level and other heart unhealthy parameters.\textsuperscript{54,55} Hence, is it really the Vitamin D supplement providing the majority of the benefit for men's health, or a finding that normal Vitamin D levels could be found on average in more healthy men? Regardless, patients should be reminded that improvement in heart healthy parameters could increase Vitamin D levels without or with additional smaller increments in supplementation. In other words, this moment represents a wonderful opportunity to emphasize heart healthy lifestyle changes first before relying on increasing the pill count of the average patient.

**PRACTICAL AND REALISTIC LIFESTYLE, SUPPLEMENT AND PRESCRIPTION INTERVENTIONS FOR PROSTATE CANCER PREVENTION**

Virtually any lifestyle change that mitigates the risk of heart disease has ample evidence today that it reduces the risk of prostate cancer, and parameters that increase the risk of heart disease increase the risk of prostate cancer. Therefore, belaboring this point or reviewing mechanisms of action or lifestyle changes in extensive detail that can simultaneously reduce or increase the risk of heart disease and prostate cancer or even other men's health issues is not the purpose of this manuscript, and this detailed information is found in multiple past written resources.\textsuperscript{14,15,23,25,27}

Encouraging patients to do whatever is practical and plausible to reduce their risk of CVD to as close to zero should be the mantra. This should provide the greatest potential to not only reduce the risk of prostate cancer, but other disease morbidity and even impact all-cause mortality. It is interesting that most major behavioral risk factors for CVD morbidity and mortality today appear to be correlated with a higher risk of aggressive prostate cancer and/or fatal prostate cancer. For example, smoking is the single largest preventable cause of death and disease in the U.S. with approximately 443 000 deaths occurring per year from tobacco related disease, and approximately 20% of adults smoke, which is a number that has remained constant the past several years.\textsuperscript{74,75} Smoking has been associated with a higher risk of being diagnosed with prostate cancer in recent meta-analyses,\textsuperscript{40} a higher risk of aggressive prostate cancer in much dying from prostate cancer.\textsuperscript{41,42} Similarly, obesity is associated with a higher risk of aggressive and fatal prostate cancer,\textsuperscript{14} and this is why it is no longer surprising that a higher risk of recurrence occurs posttreatment for prostate cancer.\textsuperscript{44} It is also plausible that the obesity is associated with a lower risk of localized prostate cancer and a higher risk of advanced disease due to the artificial lowering of PSA or hemodilution impact associated with this condition.\textsuperscript{45,46}

Weight gain ancillary issues abound, for example, ongoing evidence suggests an increased risk of certain cancers with insulin resistance, and this may include aggressive prostate cancer.\textsuperscript{87-90} Increased growth factors occur with increased insulin levels, but long-term diabetes may result in insulin, insulin-like growth factor and androgen
reduction which may be correlated with a lower prostate cancer risk in the short-term (“diabetes paradox”). The dramatic increase in the diabetes epidemic, along with the known 2–4 times increased risk of CVD events in diabetics over nondiabetics, should make type 2 diabetes prevention strategies a priority for simultaneous prostate cancer prevention. Only 15 years ago 3 states in the U.S. had a diabetes prevalence of 6% or higher, but now all 50 states in the U.S. have a rate of at least 6% or higher. Six states have rates of 10% or more along with Puerto Rico, and currently 19 million people in the U.S. have diabetes, and 7 million are undiagnosed. Perhaps, prostate cancer prevention strategies can help to modestly curb this epidemic. Exercise (aerobic and resistance), dietary (caloric reduction) and other lifestyle changes have been shown to significantly prevent diabetes and metabolic syndrome in normal and high-risk individuals better than pharmacologic therapy.

Metformin also significantly reduces diabetes risk long-term and is cost-effective with a low rate of adverse events, and has the ability to also reduce the risk of CVD events and impact all-cause mortality. Metformin is also beginning to demonstrate some consistent evidence as a cancer prevention or recurrence inhibition agent in those with and without diabetes and is currently in a phase 3 trial in breast cancer patients with survival as the primary endpoint. A recent clinical trial of patients with prostate cancer on ADT for 6 months utilizing 850 mg twice a day of metformin with caloric reduction (low glycemic diet) were able to significantly reduce weight gain, BMI, waist circumference, and systolic blood pressure compared to the control group. Men on metformin were also able to control glucose and hemoglobin A1c levels. Perhaps it is time to give serious consideration for the utilization of metformin in a phase 3 PCPT. It is cost-effective, safe, reduces weight gain, diabetes risk, and arguably CVD and perhaps total cancer and prostate cancer risk. Such a combination fits our criteria for an ideal prostate cancer prevention interventional agent.

Regular vigorous exercise (3 h or more per week) is a potential strategy to significantly reduce prostate cancer death after diagnosis, and simultaneously reduce all-cause mortality to a similar degree (50%–60%) in these same patients compared to men that perform only 1 h or less exercise per week. Thus, it should not be a surprise that exercise may also contribute to a slightly lower risk of aggressive or nonaggressive prostate cancer from a review of past studies including a recent summary of 22 studies published over the past 12 years. And, one of the suggested primary mechanism providing this protection may occur through a reduction in CVD risk especially weight/waist reduction. Patients should be told that the profound reduction in blood pressure, diabetes, depression, dyslipidemia, cancer, CVD, fatigue, obesity, and multiple other conditions would arguably be enough to earn exercise a Nobel prize if it was a drug.

Over a third (35%) of Americans have dyslipidemia, and it should again be of interest that lower cholesterol levels have been associated with a lower risk of primarily aggressive prostate cancer. Heart disease may increase the risk of prostate cancer from observations derived from two major pharmacologic studies of prostate cancer prevention. Additionally, a review of past observational studies have suggested a lower risk of aggressive prostate cancer with cholesterol lowering interventions even when controlling for multiple confounding variables. It is our opinion that statins should be investigated as a prostate cancer prevention agent, and there are trials currently being initiated to determine the role of lipid lowering in the active surveillance prostate cancer population and its impact on the progression of this disease. Yet, in the prevention setting some, would argue that it is currently too difficult to conduct such a trial when a large proportion of men are already taking these medications. This is not accurate when utilizing the Justification for the Use of Statins in Prevention: an Intervention trial Evaluating Rosuvastatin (JUPITER) trial as the most recent example of the dramatic potential impact on cardiovascular health when aggressive lipid lowering is accomplished in individuals who are in no apparent need of such intervention based on their low-density lipoprotein levels, but may need more attention based on a low-cost inflammatory marker (high-sensitivity C-reactive protein). The low-cost, CVD impact, overall benefit to risk ratio in a healthy population of men, potential prostate cancer impact, and plethora of the basic science and clinical evidence suggest that like the drug metformin, statins should be a priority intervention for in clinical trials for the potential prevention of total and aggressive prostate cancer. Arguably, the positive data existed over a decade ago to potentially study this class of agents in a large trial for prostate cancer prevention.

Hypertension is a primary risk factor for CVD and stroke, and almost a third of the U.S. adult population have this condition. Hypertension increases with age to approximately 70% of individuals 65 years and older. Hypertension is a contributing factor in one out of every 7 deaths, and 70% of individuals who have a first heart attack or stroke have hypertension. Treating hypertension has been correlated with dramatic reductions in the incidence of stroke (40%), heart attacks (25%), and heart failure (>50%). However, the correlation between prostate cancer risk and hypertension and/or anti-hypertensive medications are weak. High blood pressure as part of a continuum of unhealthy parameters such as observed with metabolic syndrome (central obesity, dyslipidemia and insulin resistance) is becoming a potential risk factor for prostate cancer risk (aggressive and nonaggressive disease) and other prostate issues. It is also well known that alpha-blockers, originally discovered for blood pressure control, are now one of those most effective treatments for men with prostate issues (benign prostatic hyperplasia) and lower urinary tract symptoms despite not having consistent positive or negative impacts on prostate cancer risk. In order to maintain prostate health it is critical to prevent or control hypertension.

CONCLUSIONS AND RESOLVING THE ONGOING PROSTATE CANCER CACOPHONY VIA LIFESTYLE AND STATINS, ASPIRIN, METFORMIN (S.A.M.)

Cardiovascular disease is the number one cause of global mortality, resulting in over 17 million deaths per year, which is a number expected to rise to over 23.5 million by 2030. Approximately 1 million heart attacks and over 700,000 strokes occur every year. More than 2200 Americans die of CVD each day, over 800,000 per year, and 150,000 of these individuals are <65 years of age, which is still lower than the average age of a prostate cancer diagnosis. Therefore, while the debate over PSA screening continues, so will the urgent need to place risk in perspective and highlight less recognized observations from these same pieces of controversial data. For example, the notable PLCO U.S. PSA screening trial which was the major impetus for the U.S. Preventive Services Task Force to recently discourage PSA screening, followed an impressive 76,993 men in 10 U.S. study centers. After 10 years of follow-up there were 174 deaths from prostate cancer, 1834 total cancer deaths, 1700 deaths from ischemic heart disease, and 3323 deaths from CVD. The debate over who might benefit or not from PSA screening may be vociferous and continuous for some time, as will the positives and negatives of past failed interventions utilized for prostate cancer prevention. However, the debate over the ideal prostate cancer prevention program should be enjoying a more halcyon

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era if clinicians, patients and the overall public become more aware
and deferential of the number 1 cause of death for men for almost
100 years, and the simultaneous impact that risk factors for CVD have
on the risk of prostate cancer itself and vice versa.

Some may think this simplistic shift in thinking is just that, much
too simplistic or too folksy to garner the attention needed for clinical
and research milieu changes. However, one could argue that it is has
been a lack of recognition or motion toward the simplistic that has
caused such deviation from the forest and such gravitation toward
the individual tree. One could even argue that 100s of millions of
dollars would not have been spent over the past two decades on
adverse pharmacologic and dietary supplement interventions had
more deference been paid to more simplistic correlations or innuendo
between the CVD and prostate cancer nexus. Perhaps it is time to realize
that after an era of subscribing to a philosophy of “more is better”
including the amount of money needed to invest in a novel and costly
preventive agent for prostate cancer or high-dose dietary supplements
that do not afford CVD protection, it is now time to believe that
“less is more” and heart health is tantamount to prostate and overall
health. Coincidentally, three accepted heart healthy preventive agents
with differing primary mechanisms of action arguably appear to be
more promising than any costly interventions that might selectively
and precisely prevent prostate cancer. Cholesterol-lowering (statins)
medications, aspirin and metformin (S.A.M.) all continue to generate
individual attention for being cost effective, primarily generic as a
class of agents, generally safe in the appropriate patients and heart and
prostate healthy in the appropriate population such as middle aged
and elderly at-risk men. S.A.M. has the ability to lower the risk of
aggressive prostate cancer and/or disease recurrence, which
would be one ideal criterion for a chemoprevention agent. It is also of
interest that all three S.A.M. interventions are low-cost, generic, heart
healthy in the appropriate patients and were all derived from “natural
sources” (statins from yeast/fungus, aspirin from willow bark and
metformin from the French Lilac).

A chronic prevalent disease, at least epidemiologically speaking
has not been found to be isolated or insular in incidence or prevalence.
Areas of the world or specific populations with some of the lowest
rates of death from CVD simultaneously enjoy lower rates of
mortality from a multitude of devastating diseases including various
cancers, and ultimately this is what assists in increasing their overall
life expectancy. Yet, if CVD risk increases at a later point in
these same areas then cancer and all-cause mortality also begins to
increase. Perhaps the moment has arrived that prostate cancer or
most cancer prevention should be solidly embedded in CVD risk
reduction strategies to maximize health benefits and longevity. It
is time to prioritize and simplify preventive health recommendations
for men, women and children especially at a time where just 1%–2% of
Americans are following multiple proven heart healthy lifestyle changes

| CVD parameter/interventions | Correlation with prostate cancer risk |
|-----------------------------|---------------------------------------|
| Aspirin (low-dose)          | Determine if a man concerned about prostate cancer qualifies for aspirin based on CVD risk working with his physicians and utilizing risk scores such as Framingham or Reynolds risk score (determine if benefit > risk overall). Aspirin may be associated with a lower risk of aggressive prostate cancer |
| Caloric control or reduction/diet | Reducing caloric intake to assist in preventing weight gain may reduce total prostate cancer risk and PSA velocity, which could reduce unneeded biopsies |
| Diabetes/glucose intolerance | Associated with a higher risk of aggressive prostate cancer |
| Dyslipidemia                 | Associated with a higher risk of aggressive prostate cancer |
| Exercise                    | Associated with a lower risk of prostate cancer and possibly aggressive disease |
| Fish oil supplements         | Overall no association with prostate cancer but preliminary indirect suggestion of a higher risk of aggressive prostate cancer with excessive intakes. Supplements should only be used in those with abnormally high triglyceride levels (FDA approved) but no clarity on CVD clinical endpoints in these individuals as of yet |
| Folic acid supplements/high-dose B-vitamin supplements | Folic acid in excessive dosages has been associated with a higher risk of total prostate cancer |
| Hypertension                 | Part of the spectrum of metabolic syndrome that could increase the risk of aggressive prostate cancer |
| Metformin                   | Reduces IGF-1, diabetes risk, weight and gluconeogenesis, which could reduce the risk of aggressive prostate cancer |
| Multivitamin                | One pill a day of a low-dose and low-cost multivitamin or children’s multivitamin in adults with or without a personal history of cancer is safe and may lower overall cancer risk (no significant impact on prostate cancer), but more than 1 multivitamin a day may increase risk of aggressive prostate cancer |
| Obesity                     | Associated with an increased risk of aggressive and fatal prostate cancer and hemodilution of the PSA test (false negatives) |
| S.A.M.                      | The acronym for teaching students and patients to remember the potential for aggressive prostate cancer prevention appears to now be immersed in CVD prevention, not just for lifestyle changes but also especially in regards to pill interventions. S.A.M. are all derived originally from “natural” sources, primarily generic, low-cost and have a long history of providing CVD protection in individuals that qualify for these medications. Additionally, they continue to garner data in prostate cancer that is arguably more impressive than any dietary supplement |
| Selenium dietary supplements | High-dose selenium supplements may increase the risk of diabetes and increase the risk of aggressive prostate cancer in men already replete with selenium from dietary sources |
| Smoking/tobacco             | Increases the risk of aggressive and fatal prostate cancer and reduces the blood level of numerous anti-oxidants |
| Statins                     | Associated with a lower risk of aggressive and advanced prostate cancer |
| Vitamin D supplements       | Potentially a U- or J-shaped curve with higher blood levels showing an increased risk of aggressive prostate cancer and normalization of deficient or insufficient levels showing a reduction in total prostate cancer risk. May just be a marker of overall health since obesity, lack of exercise, poor diet, high cholesterol, inflammatory disease reduce blood levels of Vitamin D |
| Vitamin E supplements       | High-dose Vitamin E supplements (400 IU) significantly increase the risk of nonaggressive prostate cancer and may nonsignificantly increase the risk of aggressive disease |
| Zinc supplements            | High-dose individual zinc supplements have not been shown to reduce risk and may actually increase risk of aggressive prostate cancer and may increase risk of other urologic conditions |

CVD: cardiovascular disease; S.A.M.: statins, aspirin, metformin; PSA: prostate-specific antigen; FDA: Food and Drug Administration; IGF-1: insulin-like growth factor 1
and parameters that could immediately impact disease prevalence and life expectancy. The public must be constantly distracted and fatigued by a perceived infinite of incoming behavioral recommendations from countless health awareness campaigns and agendas via multi-media sources that are now open 24 h a day and 365 days a year. How else does one explain the obsession clinicians witness regularly over medical minutiae such as the latest anti-aging supplement or drug that can apparently prevent most diseases compared to long-term, evidence-based heart healthy interventions. Therefore, Table 1 is a rapid summary of more relevant potential heart and prostate healthy and unhealthy interventions that can be utilized by clinicians and patients.

In conclusion, what if heart healthy interventions or lifestyle changes ultimately do not prevent prostate cancer from some notable future randomized trial? Attempting to reduce or compress the impact of the number one cause of morbidity and mortality in the worst case scenario is still at least a worst case scenario with a positive outcome. Hippocrates would be proud because isn't this the real translation of "first do know harm" as it relates to preventive medicine?

AUTHOR CONTRIBUTIONS
MAM reviewed past and current literature on the subject and wrote the manuscript. How do you conclude your randomized trial? Attempting to reduce or compress the impact of the number one cause of morbidity and mortality in the worst case scenario is still at least a worst case scenario with a positive outcome. Hippocrates would be proud because isn't this the real translation of "first do know harm" as it relates to preventive medicine?

COMPETING INTERESTS

The authors declare that they have no competing interests.

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