Effects of androgen-deprivation therapy on hypercoagulability in prostate cancer patients: A prospective, longitudinal study

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Abstract

Introduction: Androgen-deprivation therapy (ADT) is the cornerstone of systemic therapy for men with advanced prostate cancer (PCa) and is achieved either surgically or chemically using either gonadotropin-releasing hormone (GnRH) agonists or antagonists.1 Furthermore, antiandrogens that competitively inhibit the binding of testosterone to the androgen receptor (AR), as well as newer generation AR-axis-targeted therapies, are increasingly used in PCa management.2 Given the prolonged treatment history of men with advanced PCa, there has been renewed focus on the myriad systemic and metabolic side effects of nearly complete eradication of androgen function on target organs, specifically those of bone health, dyslipidemia, obesity, and insulin resistance,3-5 as well as effects on cognitive and psychological functioning.6-8 Furthermore, there is mounting evidence of a deleterious role of ADT on cardiovascular (CV) health.9-13 Although the literature investigating this association is still controversial secondary to differences in study design, outcomes, and at-risk populations studied, several meta-analyses of observational studies have suggested a link between ADT and CV disease and mortality.14,15 This increased risk of CV events has been observed largely during the first year after initiation of testosterone therapy.11-13 There have been several proposed mechanisms explaining these more acute physiological changes as compared to the development of metabolic syndrome with prolonged ADT use.14

One of these proposed mechanisms includes the induction of a hypercoagulable state with ADT, the basis of which was theorized after the observation that lower testosterone levels in men lead to reduced fibrinolytic activity.16,17 Furthermore, it is well-recognized that cancer patients in general are at significantly higher risk of developing thromboembolic events (TE) than healthy individuals.18 Several studies have also suggested significant perturbations in hemostasis and thrombotic risks in PCa patients.19-21 More recently, our group has shown in a small cross-sectional series that PCa patients on ADT exhibit increased hypercoagulability, with a subsequent association with venous/arterial TE events.22 The aim of the current study was to more closely examine a cohort of advanced...
PCa patients in a longitudinal study to better determine the contribution of ADT to coagulation status over time.

**Methods**

This was a prospective, longitudinal, pilot study of men with advanced PCa initiating ADT due to a diagnosis of metastatic disease or progression of their non-metastatic cancer. An age-matched control group of men with biochemical failure after local therapy who were managed on watchful waiting were also enrolled into the study. A cohort of healthy males with a recent negative prostate biopsy were used as a control of coagulation status at baseline. All men enrolled had a non-significant recent past history of either CV or TE events.

The study received ethics approval in accordance with Queens University Ethics Review Board and consent forms from each patient were obtained prior to the study. Baseline clinical and laboratory data was obtained prior to initiating ADT, including clinical disease status and basic coagulation tests (such as prothrombin time [PT] and activated partial thromboplastin time [APTT] and platelet counts). Patients were assessed clinically every three months.

Global hemostasis was evaluated at baseline and early (3–6 months), as well as late (12–15 months), after ADT initiation using the sensitive global hemostasis assay, thromboelastography (TEG). TEG was performed using a TEG® 5000 Hemostasis System and TEG® Hemostasis Analyzer software version 4.2 (Haemoscope Corporation, Skokie, IL, U.S.) and was performed according to the manufacturer’s instructions. Citrated whole blood obtained from the patients (340 μL) was recalcified with 20 μL of 0.2 M CaCl₂ and placed into the TEG cup. Data were collected for 75–90 minutes and the following major parameters were evaluated: R time, alpha (α) angle, maximum amplitude (MA) and clotting index (CI), time to clot formation (R), speed of clot propagation (α), rate of clot formation (K), strength/stability of clot (MA), and clotting index (CI); a value that is based on the four parameters above. As previously described, hypercoagulability in any individual patients was defined as two or more TEG parameters with values beyond one standard deviation of the control. To assess the global difference in coagulability among the different groups, the averages of each TEG parameter among these groups were compared.

**Statistical analysis**

Analyses were performed using GraphPad Prism Software 6 (GraphPad Software). The comparative analysis of TEG parameters between controls and baseline of PCa patients was performed using Student’s t test, as well as one-way ANOVA with Tukey’s post-hoc test. A p values less than 0.05 was considered significant.

**Results**

Eighteen patients with advanced PCa initiating ADT were enrolled into the study with a mean age of 75.5 years (range 57–84) and were followed for a minimum of 12 months. Mean prostate-specific antigen (PSA) at initiation of ADT in the cohort was 33.6 ng/ml (range 0.6–177.9). Five of 18 patients (28%) had imaging evidence of metastatic disease. Eight patients were prescribed the GnRH antagonist degarelix, eight received a GnRH agonist, and two patients were exposed to a combination of agonist and antagonist. Testosterone was maintained at castrate levels throughout the trial. Ten patients with a mean age 73.5 years (range 69–80) on watchful waiting after biochemical failure were enrolled as non-ADT controls. Mean PSA of this control group on watchful waiting was 2.64 ng/ml (range 0.4–7.4).

Compared to eight age-matched healthy men, TEG data demonstrated that 14 of 18 men (78%) of men with advanced PCa were already hypercoagulable at baseline, even before initiating ADT. The analysis of TEG parameters using one-way ANOVA showed significant differences between the three cohorts (advanced cancer-ADT, watchful waiting, and healthy controls) in three TEG parameters; alpha angle (p=0.01), MA (p= 0.04) and CI (p=0.02), but not in R time (p=0.06). Post-hoc tests showed a significant increase in MA and CI in the advanced cancer-ADT group as compared to the healthy controls. The alpha angle parameter was significantly different in the advanced cancer-ADT group vs. the watchful waiting group (Fig. 1).

During the course of ADT treatment, there were no statistically significant differences in alpha angle (p=0.65), MA (p=0.795), and CI (p=0.348) before and after ADT treatment in the advanced cancer-ADT group (Fig. 2). Interestingly, there was a significant increase in R time (reduced hypercoagulability) after 12–15 months of ADT (p=0.0004).

Individual patient analysis showed that 14/18 (78%) of the advanced PCa patients were hypercoagulable prior to ADT. After 3–6 months of treatment, only 11/18 (61%) of these patients were hypercoagulable and after 12–15 months of ADT treatment, only 10/18 (56%) patients demonstrated hypercoagulable TEG tracings. The cohort of men on watchful waiting demonstrated no significant changes in any of the TEG parameters, R time (p=0.371), angle (p=0.784), MA (p=0.649), and CI (p=0.187), over the 12–15 months of followup period (Fig. 3). It is to be noted that we did not identify any significant changes in PT (p=0.06), APTT (p=0.14), platelet count (p=0.35), or white blood cell count (p=0.96) over the study period. These data suggest that ADT may not increase hypercoagulability over time, at least within the first year of treatment.

Although as a whole cohort there was no significant changes in TEG parameters and defined hypercoagulability associated with ADT use, on an individual basis, there were
3/18 (17%) PCa patients with normal TEG parameters at baseline who subsequently became hypercoagulable. Two of the three patients had TEG tracing changes on their first assessment within 3–6 months and remained hypercoagulable on the 12–15-month assessment. The third patient was found to be hypercoagulable on the 12–15-month assessment.

We found no significant correlation between PSA levels (as a marker of disease extent and subsequent treatment effect) and degree of coagulability in these patients (baseline: r=0.0042, p=0.989; 3–6 months: r=0.055, p=0.841; 12–15 months: r=–0.204, p=0.466). In some patients with very high serum PSA levels and large volume of disease, no evidence of hypercoagulability was detected, suggesting a complex relationship between ADT, extent of disease, and coagulability in PCa patients.

**Discussion**

This prospective, longitudinal, pilot study was designed to systematically examine the relationship between ADT and coagulability in PCa patients with the objective to determine whether hypercoagulability is a measurable adverse effect of ADT, thus contributing to its observed TE/CV risks. Our results demonstrate that this cohort of men with advanced PCa were hypercoagulable before starting ADT as compared to the healthy controls, indicating hypercoagulability and a significant increase in angle as compared to the early cancer group.

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**Discussion**

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Metabolic syndrome, including obesity, elevated triglycerides, and reduced high-density lipoprotein (HDL), are established side effects of ADT.\textsuperscript{23,24} All of these well-described
physiological changes in men on ADT would conceivably lead to higher risk of eventual CV/TE events and mortality. Hypercoagulability is a direct pathophysiological factor contributing to symptomatic TE and CV complications. However, increased intravascular hypercoagulability, a proposed mechanism of early TE events in cancer patients, has not been examined systematically in association with ADT.

CV and TE events are common during PCa management, particularly for men with advanced disease requiring systemic therapy. Previous studies have documented a potential association between ADT and increased venous and arterial TE risk in PCa patients. These studies, however, failed to explore the direct impact of ADT on coagulation and the independent role of disease burden. In this current study, we provide evidence that advanced cancer alone may contribute to increased coagulability, as the majority of our cohort demonstrated hypercoagulable TEG findings before initiation of ADT and this did not appear to increase over time. The cohort of PCa patients on watchful waiting did not demonstrate any significant difference in hypercoagulability over 12–15 months’ followup. Based on our observations, it is possible that the reduced tumour burden as a result of ADT use in some patients may have improved cancer-induced hypercoagulability over time.

Our data may appear to contradict those of Ziaran et al, who studied 97 patients with locally advanced PCa and showed that after 12 months of ADT, patients had significantly higher fibrinogen, indicating hypercoagulability over time and suggesting increased CV risk in men on ADT. However, this study did not use TEG as a global, sensitive measure of hypercoagulability and did not include PCa patients on watchful waiting as controls. The findings in the current study may complicate consideration of management of the potential hypercoagulable state in PCa patients relative to previous studies. However, a comprehensive guidance document was recently published and provides specific guidelines on the prevention and treatment of cancer-associated thromboembolism. Based on this prospective, longitudinal study, hypercoagulability may not be exacerbated by ADT therapy over time, at least not in a majority of patients. Identifying patients at risk, perhaps those with previous history of CV/TE events, may be appropriate to allow more careful monitoring of their coagulation status, and determine those who may benefit from adjuvant anticoagulant prophylaxis. Although one of the strengths of this pilot study is the careful, longitudinal review of coagulation status in men initiating ADT, the cohort size limits the generalizability.

Fig. 2. Thromboelastography parameters before and after androgen-deprivation therapy (ADT) treatment in advanced prostate cancer (PCa)-ADT group over time. One-way ANOVA showed no significant difference in angle, maximum amplitude (MA), or overall clotting index over 12–15 months of treatment (indicating ADT is not associated with hypercoagulability) with a significant increase in R time at 12–15 months after ADT treatment (indicating reduced hypercoagulability).
of results for men with different TE risk factors. A larger, prospective evaluation with longer followup would allow a better understanding of more subtle effects of ADT on hypercoagulability and may facilitate an ability to predict those who may suffer TE events.

Competing interests: Dr. Siemens has participated in clinical trials for AbbVie, Astellas, BNIT, Janssen, and Sanofi. The remaining authors report no competing personal or financial interests.

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References


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