Quantifying the Evidence for the Risk of Metabolic Syndrome and Its Components following Androgen Deprivation Therapy for Prostate Cancer: A Meta-Analysis

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Abstract

Background

No meta-analysis is yet available for the risk of metabolic syndrome (MetS) following androgen deprivation therapy (ADT) for men with prostate cancer. To summarize the evidence for the link between ADT and MetS or its components quantitatively with a meta-analysis including all studies published to date.

Methods

PubMed and Embase were searched using predefined inclusion criteria to perform meta-analyses on the association between metabolic syndrome, hyperglycemia, diabetes, hypertension, dyslipidemia or obesity and androgen deprivation therapy in patients with prostate cancer. Random effects methods were used to estimate pooled relative risks (RRs) and 95% confidence intervals (CI).

Results

A total of nine studies was included. There was a positive association between ADT and risk of MetS (RR: 1.75 (95% CI: 1.27–2.41)). Diabetes was the only MetS component present in more than 3 studies, and also showed an increased risk following ADT (RR: 1.36 (95% CI: 1.17–1.58)).

Conclusion

This is the first quantitative summary addressing the potential risk of MetS following ADT in men with PCa. The positive RRs indicate that there is a need to further elucidate how type and duration of ADT affect these increased risks of MetS and diabetes as the number of men with PCa treated with ADT is increasing.
Introduction

Androgen deprivation therapy (ADT), which interrupts testosterone regulation of the prostate tumour, has been the cornerstone treatment for men with locally advanced or metastatic prostate cancer (PCa) since the 1940’s [1]. It is a highly effective treatment that inhibits testosterone production rendering patients medically castrated. A number of side effects have been reported including osteoporosis, sexual dysfunction, anaemia, cardiovascular and thromboembolic disease, and metabolic changes such as weight gain, diabetes, insulin resistance, and dyslipidemia [2–10]. The latter symptoms are all part of the metabolic syndrome (MetS), which is suggested to be a common side effect for PCa men treated with ADT [11–13]. Metabolic syndrome is defined by a combination of metabolic risks: increased waist circumference (visceral obesity), raised triglycerides or its specific treatment, reduced high-density lipoprotein cholesterol or its specific treatment, raised blood pressure or treatment of previously diagnosed hypertension, and raised fasting plasma glucose or its specific treatment. The joint statement of major international associations [14] defines everybody with three of the above listed metabolic risks as having MetS.

However, to our knowledge there is no meta-analysis to date quantifying the potential association between ADT and MetS in men with PCa [11,13,15,16]. Moreover, the underlying mechanisms are not well understood. Some studies demonstrated that a decrease in testosterone levels is associated with a decrease of 2.7–3.8% in lean body mass and an increase of 9.4–11.0% in fat mass [17–19]. The same studies indicated that ADT increases fasting plasma insulin levels [8,17,19]. Nonetheless, it is important to note that most studies of these metabolic risk factors were conducted in very small study populations over a very short follow-up period of six to twelve months [18,19]. So even though the results were statistically significant, larger prospective studies are required to confirm these findings.

In addition, there are some studies that suggest a different pathway between ADT and metabolic risk factors. For example, a small study showed that ADT preferentially increase subcutaneous rather than visceral abdominal fat and increase rather than decrease HDL cholesterol, which is in contrast with the traditionally described MetS [18]. Additionally, it was shown that ADT does not alter levels of C-reactive protein or other markers of inflammation, which suggests that ADT causes a pattern of metabolic changes that is distinct from the classically defined MetS [17].

Thus, it remains unclear to what extent ADT is associated with an increased risk of MetS or its components in men with PCa. In contrast to previous systematic reviews, this study aims to summarize the evidence for the link between ADT and MetS or its components quantitatively with a meta-analysis including all studies published to date.

Methods

Literature Search Strategy

We used computerized literature search databases (Pubmed search followed by an Embase and Cochrane Library search) to identify full-text and abstracts published to date. Our searches included “metabolic syndrome”, “hypertension”, “dyslipidemias”, “hyperglycemia”, “diabetes mellitus”, and “obesity” as search/Mesh terms for the exposure variable of interest. In addition, “prostatic neoplasms” and “androgen deprivation therapy” or “Antineoplastic Agents, Hormonal/adverse effects” were used as search/Mesh terms for the outcome variable of interest.
Our search strategy was limited to publications with a focus on humans. By not restricting the search to research papers, we made it possible to include grey literature, such as letters and abstracts presented in relevant conference meetings to address the effects of ADT on MetS. In addition, all references of selected articles were checked, including hand searches, which are effective practical ways to cross-check the completeness of the electronic searches.

Inclusion Criteria
The selected articles were chosen based on the following set of inclusion criteria: the publication pertained to an observational epidemiologic study which measured exposure to ADT; the comparison group was clearly defined as a different PCa population not on ADT; MetS or one of its components was assessed as an outcome; men with PCa were the main study population. The definition of a comparison group was important as several studies to date investigated the link between ADT and MetS by observing MetS before and after ADT in the same patients [8,17,19]. These findings may not necessarily reflect the effects of ADT as changes in the tumour itself can also have an effect on metabolic changes. These studies, often with a smaller sample size, were not included in the current meta-analysis as they did not contain control patients free of ADT. Mixing these studies with other observational studies using different control groups, would have resulted in a meta-analysis of less comparable studies.

Body composition changes were not part of the current meta-analysis as a recent meta-analysis already showed that ADT has an immediate influence on body weight, BMI, percentage fat mass and percentage lean mass [20]. Initially, titles of articles were reviewed in order to ascertain whether they might potentially fit the inclusion criteria. If, after assessing the abstract, there was any doubt regarding whether it met the relevant criteria, it was kept for more thorough, subsequent assessment. The list of potential articles was further shortened by performing detailed evaluations of the methods and results of each remaining paper. Fig. 1 provides more detailed information regarding the progressive ‘flow’ of the study exclusion process. PRISMA checklist for systematic reviews and meta-analyses is provided in the S1 Checklist. S2 Checklist shows how the STROBE criteria were used to evaluate the quality of included observational studies [21].

Data Extraction
The following details were recorded for each study: author, year of publication, ADT exposure (binary), study type (case-control or cohort), outcome, and number of cases and total subjects for each level of ADT. The outcome was defined as MetS or any of its components: hyperglycemia, diabetes, hypertension, dyslipidemia, or obesity. Despite different definitions available, it is generally accepted that MetS is defined by a combination of metabolic risks: increased waist circumference (visceral obesity), raised triglycerides or its specific treatment, reduced high-density lipoprotein cholesterol or its specific treatment, raised blood pressure or treatment of previously diagnosed hypertension, and raised fasting plasma glucose or its specific treatment. The joint statement of major international associations defines everybody with three of the above listed metabolic risks as having MetS [14]. Hence, the current meta-analysis dichotomized the different outcomes irrespective of the definitions used.

Meta-Analysis Statistical Techniques
The effect of ADT on risk of MetS or its components among men with PCa was evaluated by calculating the random effects summary relative risk. Forest plots were created as they display the relative risk estimates of MetS risk comparing both levels of ADT for each study. Potential heterogeneity of the study results was statistically evaluated using the I² statistic. Potential
publication bias was assessed using Begg’s Test. All analyses were performed using STATA (version 12).

**Results**

The initial search for ADT and MetS or its components resulted in 79 articles via PubMed and 170 via Embase. After extracting information from the abstracts, 28 articles were selected for further investigation. Finally, nine studies were selected for primary data-analysis from which six studies were conducted in the United States, two in Spain, and one study in China. One additional study, conducted in Canada, was identified via hand searches (**Fig. 1 and Table 1**).

Based on the above defined inclusion criteria we excluded 20 studies (**Fig. 1**). Amongst these, eight were excluded due to lack of control populations, three were systematic reviews, three presented data that overlapped with studies that were already included, and another six were omitted due to lack of information or methodological errors.

The random effects analysis, comparing ADT and risk of MetS, indicated a pooled effects relative risk of 1.75 (95%CI: 1.27–2.41). This pooled analysis included four studies of which three used the National Education Cholesterol Programme definition for MetS and one used the definition from the International Diabetes Federation [22,23]. The $I^2$ statistic suggested no heterogeneity ($I^2 = 0.00\%$), which can also be observed in the corresponding Forest plot.
Table 1. Overview of studies included in meta-analyses.

| Author | Country | Study Type | ADT Type | Outcome | Number of patients | Main findings |
|--------|---------|------------|----------|---------|-------------------|---------------|
| Keating et al. [37] | USA | Cohort | GnRH agonist, combined androgen blockage, orchiectomy, anti-androgens | Diabetes | 14,597 ADT. 22,846 no ADT | No ADT: Ref. GnRH agonists: 1.28 (95%CI: 1.19–1.38) Orchiectomy: 1.16 (95%CI: 0.87–1.54) Combined androgen blockage: 1.17 (95%CI: 0.96–1.42). Oral anti-androgens: 1.02 (95%CI: 0.72–1.45) |
| Lage et al. [38] | USA | Retrospective claims database | Any ADT | Diabetes | 1,231 on ADT. 7,250 no ADT | While controlling for other factors, the estimated relative risk of incident diabetes associated with the receipt of ADT was 1.36 (95% CI: 1.07–1.74) |
| Keating et al. [6] | USA | Cohort | GnRH agonist, orchiectomy | Diabetes | 26,570 on ADT. 46,626 on ADT | No ADT: Ref. GnRH agonists: 1.44 (95%CI: 1.34–1.55) Orchiectomy: 1.34 (95%CI: 1.20–1.50) |
| Braga-Basaria et al.[39] | USA | Cross-sectional | ADT | Mets\(^1\) Obesity Hyperglycaemia Hypertriglyceridemia Low HDL Hypertension | 20 on ADT 18 no ADT | Prevalence of MetS: 55% vs 22% (ADT vs no ADT). Prevalence of obesity: 75% vs 33% Prevalence of hyperglycaemia: 65% vs 16%. Prevalence of hypertriglyceridemia: 55% vs 44%. Prevalence of low HDL: 35% vs 50%. Prevalence of hypertension: 45% vs 28% |
| Basaria et al.[40] | USA | Cross-sectional | ADT | Hyperglycaemia | 18 on ADT 17 no ADT | Men on ADT had significantly higher levels of fasting serum glucose (131.0 mg/dL) compared with men not on ADT (103.0 mg/dL; P: 0.01) |
| Bo et al. [41] | China | Cross-sectional | ADT Orchiectomy | Metabolic changes\(^2\) | 46 orchiectomy/ ADT 37 prostatectomy no ADT. 50 controls. | After 3 months ADT group had increased levels of fasting serum insulin and LDL compared to the other 2 groups (P< 0.05) After 12 months ADT group had increased levels of waist circumference, fasting serum insulin and glucose, total cholesterol, HDL and LDL compared to the other 2 groups (P<0.05) |
| Garcia et al.[42] | Spain | Cross-sectional | ADT | Mets\(^1\). Osteoporosis | 216 on ADT. 50 no ADT | Prevalence of Mets in no ADT: 19% Prevalence Mets ADT: 6 months treatment: 21%. 12–18 months treatment: 36%. >24 months treatment: 24% |
| Valverde et al.[43] | Spain | Cross-sectional | ADT | Mets\(^1\) | 53 on ADT. 104 no ADT (52 PCa 52 no PCa) | Mets in patients on ADT: 51.9%. Mets in patients without ADT: 35.8% |
| Alibhai et al. [44] | Canada | Cohort | ADT Orchiectomy | Diabetes Acute myocardial infarction Sudden death | 19,079 on ADT /Orchiectomy 19,079 men with PCa no ADT | Increased risk of diabetes HR 1.16 (95%CI: 1.11–1.21). No increased risk of AMI HR 0.91 (95%CI: 0.84–1.00) or of sudden death HR 0.96 (95%CI: 0.83–1.10) |

\(^1\) MetS Definition of the National Cholesterol Education Programme—Adult treatment panel III

\(^2\) Mets Definition of the International Diabetes Federation.
The only MetS component with more than 3 studies was diabetes. The random effects analysis showed a pooled relative risk of 1.36 (95% CI: 1.17–1.58) for the association between ADT and risk of diabetes. The I² statistic suggested heterogeneity (I² = 84.7%), but this was rather limited as can be seen in the corresponding Forest plot (Fig. 3). For both analyses, Begg’s test did not indicate publication bias (P: 0.34 and 0.23, respectively), which is also evident from the funnel plots as there is a symmetric distribution observed among studies (Fig. 4).

**Discussion**

Despite the plethora of reviews on risk of MetS or its components following ADT for PCa [11–13,24], this is the first meta-analysis of evidence published to date. The results suggest a 75% increased risk of MetS and a 36% increased risk of diabetes following ADT for men with PCa.

In 2010, the Food and Drug Administration required labeling on gonadotropin-releasing hormone (GnRH) agonists, warning men about an increased risk of diabetes when receiving these medications for PCa treatment [25]. Supporting evidence came from large North American cohorts [6,26,27]. To our knowledge no large European studies have yet quantified the risk of diabetes following ADT, nor have most cohorts made a distinction between different types and duration of ADT while comparing with PCa-free men. This information would be of
interest to patients and clinicians as the association between ADT and PCa may be affected by differences in lifestyle and treatment practices, but most importantly by type and duration of ADT.

The current meta-analysis thus aimed to quantitatively summarize how ADT increases risk of metabolic abnormalities. The findings corroborated the above-mentioned FDA statement as we found a positive association for ADT with both MetS and diabetes. A recent systematic review, not including any summary statistics, also concluded that most studies indicate that ADT is positively associated with risk of insulin resistance and MetS [28]. However, based on the limitations of some studies, such as lack of sample size calculations or control groups, this review concluded that more studies are needed to further disentangle the association between ADT and metabolic abnormalities. The latter would require studies focused on specific subtypes of ADT, but most studies to date did not show results for different subtypes of ADT. The study by Keating et al. [6] was the only making a distinction between orchiectomy and GnRH agonists and found a slightly higher risk for those on GnRH agonists (HR: 1.44) than those who underwent orchiectomy (HR: 1.34). In a more recent study, the same authors showed that treatment with GnRH agonists was associated with increased risk of diabetes (HR: 1.28), but no statistically significant association was found for anti-androgens, combined androgen blockage, or orchiectomy [26]. Thus, even though most studies found a positive association

| Study                                      | RR (95% CI)         | Weight |
|-------------------------------------------|---------------------|--------|
| Keating et al 2010                        | 1.71 (1.51, 1.94)   | 20.37  |
| Lage et al. 2007                          | 1.28 (1.05, 1.56)   | 16.77  |
| Keating et al 2006 (GnRH agonists)        | 1.39 (1.26, 1.53)   | 21.50  |
| Keating et al 2006 (orchiectomy)          | 1.17 (0.98, 1.41)   | 17.32  |
| Braga-Basaria et al. 2006                 | 3.90 (1.32, 11.51)  | 7.8    |
| Allibhai et al. 2009                      | 1.18 (1.09, 1.27)   | 22.26  |
| Overall (I-squared = 84.7%, p = 0.000)    | 1.36 (1.17, 1.58)   | 100.00 |

Fig 3. Forest plot for association between ADT and diabetes.
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Fig 4. Begg’s funnel plots to test for publication bias for the associations between ADT and diabetes (a) and ADT and diabetes (b).

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between ADT and metabolic abnormalities, more experimental and epidemiological studies are needed to differentiate the metabolic effects of different types of ADT.

In addition to an increase in epidemiological evidence, also experimental biological findings are increasingly supporting an association between ADT and MetS or its components. Given that ADT lowers testosterone levels, evidence suggests that pathogenic mechanisms linking these low testosterone levels with MetS or its components are complex and bi-directional. Visceral obesity has been shown to be a cause of hypogonadism given that adipose tissue is a key source of estrogens due to the presence of an enzyme that converts testosterone into estrogens [29–32]. On the other hand, several studies have shown that testosterone increases lean body mass and that low levels of this hormone promote fat deposition [12,18,33]. Obesity has also been shown to be a major risk determinant for insulin resistance and type 2 diabetes [34]. Furthermore, low levels of testosterone have been linked to alterations in lipoprotein lipase enzyme activity and increases in triglyceride turnover, leading to abnormal levels of LDL and triglycerides, both components the MetS [35]. Additionally, testosterone has been suggested to regulate cell lineage determination by promoting the myogenic and inhibiting the adipogenic lineage [36].

Nonetheless, a limitation of observational studies is the possibility of bias introduced by selection of men to receive ADT. Men who are treated with ADT may differ from men who are not in ways that are also associated with risk for metabolic abnormalities. Although most of the studies included in this meta-analysis conducted analyses adjusting for observed confounders, we could only rely on crude event rates, as most studies did not provide sufficient data to allow us to account for potential confounders in our analyses. In future studies it would be of interest to add sensitivity analyses focused on specific subgroups of patients such as, for instance, those with or without a history of cardiovascular disease.

Our literature search methods were performed to include all relevant publications available to date through various sources, including grey literature, and two main online databases (Pubmed and Embase). Furthermore, objective inclusion and exclusion criteria were defined a priori. All included studies fulfilled these criteria and had a clearly defined study design and statistical analysis plan. There were however not enough studies available to examine the association between ADT and each component of MetS individually. Combining different definitions of MetS was needed given the small number of studies published to date and the recently published joint statement of major international associations defining everybody with three of the above listed metabolic risks as having MetS. However, all definitions were clearly stated and complied with this joint statement [14]. Furthermore, most studies investigated GnRH agonists as the main type of ADT, so that it was not possible to make a distinction between different types of ADT. Another limitation is that we could not make a distinction between patients with and without a history of cardiovascular disease. This difference could have shown whether ADT increases even more the risk of metabolic abnormalities in a subgroup of patients with cardiovascular history.

Conclusion

This meta-analysis quantified the positive association between ADT for PCa and risk of developing MetS and diabetes. It also highlights the need to further disentangle this association by investigating different types and duration of ADT in relation to risk of these metabolic disturbances. The latter may provide insight in potential underlying mechanisms and may advocate the potential value of applying appropriate lifestyle changes.
Supporting Information

S1 Checklist. PRISMA Preferred Reporting Items for Systematic Reviews and Meta-Analyses checklist.
(PDF)

S2 Checklist. STROBE STrengthening the Reporting of OBServational studies in Epidemiology checklist.
(PDF)

Author Contributions

Conceived and designed the experiments: MVH JA. Performed the experiments: CB MVH. Analyzed the data: MVH CB. Contributed reagents/materials/analysis tools: MVH JA. Wrote the paper: CB DC JA SR MVH.

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