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Association of polymorphisms in CYP19A1 and CYP3A4 genes with lower urinary tract symptoms, prostate volume, uroflow and PSA in a population-based sample

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Abstract

Purpose The known importance of testosterone for the development of benign prostatic hyperplasia (BPH) prompted us to test the hypothesis whether polymorphisms of two genes (CYP19A1 and CYP3A4) involved in testosterone metabolism are associated with clinical BPH-parameters.

Methods A random sample of the population-based Herne lower urinary tract symptoms cohort was analysed. All these men underwent a detailed urological work-up. Two polymorphisms in the CYP19A1 gene [rs700518 in exon 4 (A57G); rs10046 at the 3’UTR(C268T)] and one in the 3’UTR of CYP3A4 [rs2740574 (A392G)] were determined by TaqMan assay from genomic DNA of peripheral blood. These polymorphisms were correlated to clinical and laboratory BPH-parameters.

Results A total of 392 men (65.4 ± 7.0 years; 52–79 years) were analysed. Mean International Prostate Symptom Score (IPSS; 7.5), Qmax (15.4 ml/s), prostate volume (31 ml) and prostate specific antigen (PSA) (1.8 ng/ml) indicated a typical elderly population. Both polymorphisms in the CYP19A1 gene were not correlated to age, IPSS, Qmax, prostate volume and post-void residual volume. Serum PSA was higher in men carrying the heterozygous rs10046 genotype (2.0 ± 0.1 ng/ml) than in those with the CC-genotype (1.7 ± 0.2 ng/ml, P = 0.012). Men carrying one a mutated allele of the CYP3A4 gene had smaller prostates (27.0 ± 2.0 vs. 32 ± 0.8 ml, P = 0.02) and lower PSA levels (1.6 ± 0.3 vs. 1.9 ± 0.1 ng/ml).

Conclusions The inconsistent associations observed herein and for other gene polymorphisms warrant further studies. In general, the data regarding the association of gene polymorphism to BPH-parameters suggest that this disease is caused by multiple rather than a single genetic variant. A rigorous patient selection based on anatomopathological and hormonal profile may possible reduce the number of confounders for future studies thus enabling a more detailed assessment of the association between genetic factors and BPH-parameters.

Keywords Prostate · Gene polymorphism · Marker · BPH

Introduction

Benign prostatic hyperplasia (BPH), benign prostatic enlargement (BPE) and benign prostatic obstruction
(BPO) are progressive disorders that can cause lower urinary tract symptoms (LUTS) and ultimately lead to acute urinary retention (AUR) or the need for prostatic surgery [1]. Predictors for disease progression are age, symptom status, post-void residual volume, maximum flow rate ($Q_{\text{max}}$) and—currently most widely used—prostate volume with prostate specific antigen (PSA) as a proxy parameter [2]. Progression risk assessment is used in contemporary medical BPH management treatment algorithms [3].

Long-term treatment with 5α-reductase inhibitors (5ARIs) results in a 50% risk reduction for AUR or the need for surgery indicating that the disease progression can be altered by medical therapy [4, 5]. The magnitude of this effect is dependent on various parameters, such as prostate volume. The prostate cancer (PCa) prevention trial (PCPT) has also shown that long-term treatment with finasterid in healthy men without LUTS leads to a favourable outcome regarding several BPH-parameters (AUR, need for surgery, urinary tract infection) thus supporting the concept of a preventive strategy [6].

The development of LUTS in men is a multifactorial process with multiple factors involved, such as BPH, BPE, BPO, inflammatory processes in the prostate, changes in detrusor function, alterations of the peripheral and central nervous system and—potentially—genetic factors. A genetic marker predictive for the development of BPE, BPO or LUTS would be a major improvement for preventive strategies and patient management. In contrast to clinical parameters such as prostate volume or PSA, genetic markers would allow a much earlier risk assessment with earlier and potentially more effective intervention, e.g. by 5ARIs.

Polymorphisms of genes involved in the testosterone metabolism are currently under intense evaluation as biomarkers for PCa risk [7, 8]. The CYP19A1 gene is located on chromosome 15 (15q21.1) and encodes the enzyme aromatase catalysing the irreversible conversion of androstenedione to estrone and testosterone to estradiol [8]. Aromatase is present in the gonads and in the extragonadal tissue, including the prostate and adipose tissue [8]. Aromatase mRNA and protein have both been detected in BPH and PCA tissue. Polymorphisms in the CYP19A1-gene showed borderline increased risks for PCA [8]. CYP3A4 is located on chromosome 7 (7q21.1) and involved in the oxidation of testosterone to $2\beta$-, $6\beta$- or $15\beta$-hydroxytestosterone, which is biologically less active than testosterone or dihydroxytestosterone [8]. A variant in the 5′ untranslated region of CYP3A4 (rs2740574) has been associated with PCA risk in some studies and also in men with a history of BPH [9].

The known importance of androgens for the pathogenesis of BPH/BPE, the potential role of CYP19A1 and CYP3A4 genes in the androgen metabolism and association of both polymorphisms to PCa risk prompted us to investigate the association of genetic polymorphisms in the CYP19A1 and CYP3A4 genes, to several clinical and laboratory BPH-parameters in a population-based sample (Herne LUTS study). Neither of these polymorphisms has been previously tested for their association to clinical BPH-parameters.

Materials and methods

The design of the Herne LUTS study has been outlined elsewhere [10]. In brief, this is a population-based study that was initiated in 1999 in the city of Herne, Germany. Of the 8,973 men on the mailing list, 60.2% ($n = 5,404$) completed the initial questionnaire and underwent a detailed urological examination. For the current analysis all men meeting the following criteria were eligible: serum PSA <10 ng/ml, age 50–79 years and a prostate volume <100 ml.

Of a subset of men who entered the Herne LUTS study whole blood serum samples were stored suitable for DNA extraction. This sample of 392 men entered the current study. At baseline all participants underwent a detailed clinical investigation with assessment of the following parameters: International Prostate Symptom Score (IPSS), free uroflow study, post-void residual volume (PVR), prostate volume by transrectal ultrasound (TRUS) and a serum PSA analysis. All PSA analyses were performed by the same PSA-test.

The presence of PCa was carefully excluded by digital rectal examination and serum PSA analysis. All men with a suspicious finding on digital rectal examination (DRE) and/or serum PSA >4.0 ng/ml underwent TRUS guided biopsy and were only included if the biopsy was negative. Serum androgen levels were not determined.

Genotyping

Genomic DNA was extracted from peripheral blood using QIAamp Blood Midi Kit (QIAGen, Hilden, Germany). The polymorphisms were determined by 5′-nuclease TaqMan assays with ABI PRISM 7500 sequence detection system. The reaction contained 20 ng genomic DNA, TaqMan $2 \times$ PCR Master Mix, forward and reverse primers and probes for the wild-type and the mutant allele in a total volume of 10 µl. Universal reaction conditions were 2 min at 50°C, 10 min at 95°C, 40 cycles with 15 s at 92°C and annealing/extension at 60°C for 1 min. Allelic discrimination was carried out by measurement of fluorescence yields of the two different dyes at 60°C.
Statistical analyses

Linear regression models were estimated for parameters IPSS, prostate volume, $Q_{\text{max}}$ and PSA, respectively. Stepwise backward regression was applied to select polymorphisms that were used to generate the final model. At each step, Akaike’s information criterion (AIC) was calculated to decide if the polymorphism with the smallest effect was dropped from the model. Age was included as confounder in all four models. Polymorphisms considered as regressors were rs700518, rs1004 and CYP3A. Statistical procedures were calculated by the computer software SPSS (Chicago, IL) for Windows (version 6.0.1) and Epi Info (version 6.04c, Centers for Disease Control and Prevention, Atlanta, GA). Descriptive analyses included Student’s $t$ test of means and standard deviations. Results were considered to be statistically significant when $P < 0.05$; all $P$ values were checked for multiple testing according the Bonferroni method.

Results

Patient characteristics

Principal characteristics of the study population ($n = 392$) are listed in Table 1. These data suggest a typical population of elderly men generated via population-based sampling.

Cyp19A1 gene polymorphism

All polymorphisms were distributed in the Hardy–Weinberg equilibrium.

Two polymorphisms in the CYP19A1 gene [rs700518 in exon 4 (A57G); rs10046 in the 3’UTR (C286T)] were determined (Table 2). The rs700518-AA genotype was present in 96 (24.5%), the heterozygous in 194 (49.5%) and the homozygous GG genotype in 102 men (26%). IPSS, $Q_{\text{max}}$, prostate volume and PSA were similar across all rs700518 genotypes (Table 2).

Cyp3A4 gene polymorphism

The AA genotype was present in 360 men (91.8%) and the heterozygous genotype (AG) in 32 men (8.2%); no one had the homozygous GG genotype (Table 3). There was no association between the CYP3A4 gene polymorphism and IPSS and the $Q_{\text{max}}$ (Table 3). Prostate volume was larger in men carrying the wild-type genotype [32.1 ± 0.8 (range 25–87 ml) vs. 27.0 ± 2.0 (range 26–78 ml), $P = 0.02$]. In parallel, men carrying the AA genotype had a significantly higher serum PSA (1.9 ± 0.1 ng/ml) than those with one G allele (1.6 ± 0.3 ng/ml, $P = 0.046$). This level of significance was lost after adjustment for multiple testing ($\alpha$ value 0.017).

Discussion

Polymorphisms of genes involved in the androgen pathway hold promise as biomarkers for PCa, for review see [7, 8]. However, the association between androgen pathway gene polymorphisms and PCa risk is complex and characterized by contradictory results [7, 8]. The cause of this conflict in any particular association of genotype and phenotype is difficult to identify and it can be attributable to biological, statistical and technical causes.

Relatively little is known regarding the potential role of gene polymorphisms for the development of BPH/BPE. Although cellular origins of BPH/BPE and PCa are different, both diseases are under comparable endocrine and may be under similar genetic control.

Several reasons render genetic markers for BPE clinically relevant: (1) high prevalence of the disease, (2) demographic trends towards ageing in upcoming decades, (3) availability of medical therapies (5ARI) known to alter the natural history to the disease and (4) BPE is an important prognosticator for LUTS-related complications. Men with prostates >30 ml have a threefold increase in the risk of AUR and a fourfold increase in the risk of undergoing BPH-related treatment compared to men with smaller

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Principal cohort characteristics ($n = 392$)</th>
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<tbody>
<tr>
<td></td>
<td>Mean</td>
</tr>
<tr>
<td>Age (years)</td>
<td>65.4</td>
</tr>
<tr>
<td>IPSS</td>
<td>7.5</td>
</tr>
<tr>
<td>$Q_{\text{max}}$ (ml/s)</td>
<td>15.4</td>
</tr>
<tr>
<td>Prostate volume (ml)</td>
<td>31.1</td>
</tr>
<tr>
<td>PSA (ng/ml)</td>
<td>1.8</td>
</tr>
</tbody>
</table>
age has been implicated in the induction and progression of prostate cancer [1]. The oestrogen-dominant status in men after middle age, may lead to an increase in oestrogen production via the aromatase enzyme. CYP19A1 codes for aromatase that converts C19 androgens into aromatase C18 oestrogenic steroids [8]. Hence, up-regulation of CYP19A1 may lead to an increase in oestrogen production. The oestrogen-dominant status in men after middle age has been implicated in the induction and progression of prostate cancer [11]. The recent data suggest that aromatase expression in prostatic stromal cells is induced by prostaglandin E2 via a paracrine mechanism [11]. However, functional tests in the CYP19A1 gene showed no significant changes in protein activity or level compared with the WT enzyme after transient expression in COS-1 cells [8]. Studies correlating several polymorphisms in the CYP19A1 gene to PCa yielded conflicting results with the majority being negative [7, 8]. This is in line with the current study: the rs700518 polymorphisms showed to correlation to clinical BPH-parameters, the rs10046 revealed a weak correlation to PSA. While the latter remained significant after adjustment for multiple testing, it is noteworthy that there was no “gene dose-response”, i.e. that homozygous TT carriers had higher mean annual growth rate than smaller ones [1, 2].

Our study has two strengths: first, we have investigated a population-based, representative sample of elderly men, hence our cohort is not biased by referral patterns; secondly, all men in this study underwent a detailed urological investigation with exclusion of PCa. Although it well established that around 20% of men have PCa in the <4.0 ng/ml PSA-range based on the PCPT, the cut-off value of 4.0 ng/ml is clinically established. It is unrealistic to propose routine prostate biopsies in a population-based sample. Population-based samples, which are not carefully screened with respect, carry the risk of being contaminated with PCa cases. A limitation is the rather small sample size; the down-side of using a population-based cohort is that the number of men with severe LUTS and significantly impaired flow rates is limited.

This is the first study that has investigated the association of polymorphisms within CYP19A1 and CYP3A4 genes to BPH-parameters. The CYP19A1 gene on 15q21.1 codes for aromatase that converts C19 androgens into aromatic C18 oestrogenic steroids [8]. Hence, up-regulation of this enzyme may lead to an increase in oestrogen production. The oestrogen-dominant status in men after middle age has been implicated in the induction and progression of prostate cancer [1]. In addition larger prostates have a higher mean annual growth rate than smaller ones [1, 2].

Table 2: Association between two polymorphisms in the Cyp19A1-gene and BPH-parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>RS 700518</th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>A/A (n = 96)</td>
<td>A/G (n = 194)</td>
</tr>
<tr>
<td></td>
<td>IPSS 8.0 ± 0.6</td>
<td>7.4 ± 0.4</td>
</tr>
<tr>
<td></td>
<td>Q_{max} (ml/s)</td>
<td>15.7 ± 0.9</td>
</tr>
<tr>
<td></td>
<td>Prostate vol. (ml)</td>
<td>31.6 ± 1.5</td>
</tr>
<tr>
<td></td>
<td>PSA (ng/ml)</td>
<td>1.7 ± 0.2</td>
</tr>
</tbody>
</table>

Table 3: Association between a CYP3A4 gene polymorphism and BPH-parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>RS 10046</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C/C (n = 91)</td>
<td>C/T (n = 190)</td>
</tr>
<tr>
<td></td>
<td>IPSS 8.0 ± 0.07</td>
<td>7.5 ± 0.4</td>
</tr>
<tr>
<td></td>
<td>Q_{max} (ml/s)</td>
<td>15.7 ± 0.9</td>
</tr>
<tr>
<td></td>
<td>Prostate vol. (ml)</td>
<td>31.6 ± 1.6</td>
</tr>
<tr>
<td></td>
<td>PSA (ng/ml)</td>
<td>1.7 ± 0.2</td>
</tr>
</tbody>
</table>

Numbers indicate mean + standard error of the mean

* P = 0.012
no statistically association (after adjustment for multiple testing) of a CYP3A4-polymorphism to clinical BPH-parameters.

Several other polymorphisms in the androgen pathway have been correlated to clinical BPH-parameters. Four studies have determined the association of the number of CAG-repeats within the androgen receptor (AR) to prostate volume [12–15]. Three studies, including a previous one from our group analysing a typical BPH-population (men referred for TURP), were negative [12–15]. Giovannucci et al. [16] reported on an association of the number of CAG-repeats within the AR to prostate volume and risk for prostate surgery.

Four studies addressed the potential role a polymorphism within the 17 hydroxylase (CYP17) gene, one of the key enzymes in testosterone biosynthesis to several BPH-parameters [17–20]. While two studies observed no association to hormone levels, our group observed a lower stroma/epithelial cell ratio in individuals carrying the mutated allele [17, 18]. In a Turkish cohort, Gunes et al. [19] observed an association between this polymorphism and BPH. By analysing a population-based Chinese cohort, Madigan et al. [20] failed to identify a clear association between CYP17 polymorphism and BPH-parameters.

The potential role of polymorphisms within the 5α reductase gene (SRD5A2) in the development of BPH/BPE has also been addressed [13, 21, 22]. An association to serum testosterone levels has been observed in both studies analysing the SRD5A2 V89L polymorphism; larger prostates and higher PSA levels were correlated to the SRD5A2 A49T genotype [13]. Salam et al. [22] reported on a non-significant association between the SRD5A2 V89L polymorphism and BPH, yet noticed a significant association for Hispanics.

In conclusion, all these studies—including the current one—yielded conflicting data which is in part due to different study designs (cross-sectional, longitudinal), different study populations (surgical cases, outpatients, population-based cohorts, screening populations) and BPH-definitions (prostate volume, $Q_{\text{max}}$, need for surgery). Many human traits and their underlying polymorphic genes show independent patterns of racial, ethnic and geographic variation. Understanding the importance of polymorphisms in the androgen pathway genes still requires more studies. A predisposition caused by a combination of genetic variants, of which each carries a small relative risk, may be of much greater significance. Hence, this understanding can only emerge if research extends beyond single gene studies to encompass gene–gene and gene–environment interaction studies. The inconsistent associations observed herein warrant further studies. In general, the data regarding the association of gene polymorphism to LUTS, prostate volume, uroflowmetry and PSA suggest that this disease is caused by multiple rather than a single genetic variant. If this is true, it becomes unlikely that any single gene polymorphisms will exhibit a strong enough correlation with BPH-parameters and/or progression risk to be clinically useful.

**Conflict of interest statement** There is no conflict of interest.

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