AKT2 Gene Polymorphisms, Srankl/OPG And Hormone Measurements in Polycystic Ovarian Syndrome (PCOS) Women

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Abbreviations

BMI: Body Mass Index; CI: confidence intervals; HWE: Hardy Weinberg Equilibrium; GVS: Genome Variation Server; MAF: Minor Allele Frequency; OPG: osteoprotegerin; OR: Odds ratio; PCOS: Polycystic Ovarian Syndrome; PCR: Polymerase Chain Reaction; POM: Polycystic ovarian morphology; qPCR: quantitative Polymerase Chain Reaction; sRANKL: soluble receptor activator of nuclear factor (NF)-κB ligand; SNP: Single Nucleotide Polymorphism; SD: Standard Deviation; Tm: Melting Point.

Introduction

Polycystic ovarian syndrome (PCOS) is a common endocrinologic disorder diagnosed in 6-10% of the female population at reproductive age [1-2]. Although many studies have investigated PCOS prevalence, there are discrepancies in their results, partly due to the use of various definitions of the syndrome and its sub-phenotypes, differences between study cohorts, ethnicities, and types of recruitment and sampling. PCOS is characterized by hyperandrogenism, oligo-ovulation, and polycystic ovarian morphology [3-4].

In 1990 the National Institute of Health (NIH) proposed the first PCOS diagnostic criteria that required the combination of chronic oligo/anovulation and clinical or biochemical evidence of hyperandrogenism, with the exclusion of related disease such as non-classical congenital adrenal hypertrophy, Cushing syndrome,
dependent kinase (PIDK), which is activated by the product of PI3K, participates in insulin signaling. There is reduced activation of AKT2 gene by phosphatidylinositol 3,4,5-trisphosphate and participates in insulin metabolic arm through the activation of phosphatidylinositol 3-kinase (PI3K) and a mitogenic arm acting via mitogen-activated protein kinase [9]. The expression and activity of PI3K downstream targets have been investigated in PCOS tissues, with distinct molecular defects in post–insulin receptor signaling identified in fibroblasts and adipose tissues of PCOS women [10]. AKT gene activates 3-phosphoinositide dependent kinase (PI3K), which is activated by the product of PI3K, phosphatidylinositol 3,4,5-trisphosphate and participates in insulin signaling. Three highly conserved genes encode the forms of AKT1, AKT2 and AKT3 [7,11]. AKT2 (AKT serine/threonine kinase 2) is a highly conserved genes encode the forms of AKT1, AKT2 and AKT3 [7,11]. AKT2 (AKT serine/threonine kinase 2) is located at 19q13.2 locus (22 exons) and is a widely expressed gene that is involved in metabolism of insulin, mitogenic signaling and apoptosis [11]. In type 2 diabetes, there is reduced activation of AKT2 gene by insulin in adipocytes. Stimulin-stimulated AKT phosphorylation is reduced 40-60% in PCOS skeletal muscle compared with matched control muscle [9].

It has traditionally been suggested that prolactin excess, enzymatic steroidogenetic abnormalities and thyroid disorders need to be excluded before a PCOS diagnosis is confirmed. However, there is paucity of data regarding the prevalence of PCOS phenotype in some of these disorders, whereas other endocrine diseases that exhibit PCOS-like features may evade diagnosis and proper management, if not considered. Hyperinflammation and impaired epithelial function were reported to a larger extent in PCOS women and could be particularly associated with hyperandrogenism [12], obesity and insulin resistance. Available data from register-based and data linkage studies support that metabolic-vascular and thyroid diseases, asthma, migraine, depression, and cancer are diagnosed more frequently in PCOS, whereas fracture risk is decreased [13]. Drug prescriptions are significantly more common in PCOS than controls within all diagnostic categories including antibiotics use. The causal relationship between PCOS and autoimmune disease represents an interesting new area of research. PCOS is a lifelong condition and long-term morbidity that could be worsened by obesity, sedentary lifestyle, western-type diet and smoking, whereas lifestyle intervention including weight loss may partly or fully resolve the PCOS symptoms and could improve the long-term prognosis. As many as 20% of women with infertility problems (including fecundability and early pregnancy loss) have been diagnosed with PCOS. It is often called the most common cause of anovulatory infertility in women.

PCOS is also associated with cardiovascular problems, neurological and psychological effects on quality of life (including anxiety and depression), and breast and endometrial cancers [14]. Recently there was a connection between cardiovascular disease and osteoprotegerin (OOG, an osteoclast inhibitory factor) and sRANKL (soluble receptor activator of nuclear factor (NF)-κB ligand) biomarkers [15]. There is no known PCOS cause; however, there has been evidence that shows that both environmental as well as genetic factors play a role in the etiology [16].

The goal of this study was to investigate the association among AKT2 gene SNPs (Single Nucleotide Polymorphisms), OPG/sRANKL as cardiovascular biomarkers, hormone measurements and clinical characteristics as well in PCOS patients in the Greek female population.

Materials and Methods

Subjects

We studied a total of 60 Caucasian patients with PCOS aged 18-35 and 30 Caucasian control women that were age-and BMI-matched. Weight and height were measured and obesity was assessed by estimating Body Mass Index (BMI). All patients were recruited from the reproductive endocrinology clinic of Attikon University General Hospital (physician in charge: E.T.). Participation in the research protocol was offered to patients meeting the PCOS Rotterdam diagnostic criteria and the following inclusion criteria: being premenopausal, not pregnant and not submitted to hormonal therapy (including oral contraceptives) for at least 3 months prior to testing. Control subjects were healthy women working at Attikon University General Hospital with regular menstrual cycles and without any family history of PCOS or hirsutism. They had no evidence of hirsutism, acne, alopecia, endocrine dysfunction and were not submitted to hormonal therapy (including oral contraceptives) for at least 3 months prior to testing. A standard questionnaire was filled, including menstrual periods and irregularity, hirsutism and acne, reproductive history, gynecological history and use of medications including oral contraceptive pills. The presence of hirsutism was scored in all subjects using the Lorenzo Scale [17]. Subjects with diabetes were excluded. Transvaginal ultrasonography was performed in all subjects. The ethics committee of the hospital approved the study and a signed informed consent was obtained from each participant.

Methods

Serum measurements

Whole blood was obtained on day 2 or 3 of the menstrual cycle or during amenorrhea, centrifuged for 10 min at 3500 rpm and serum was collected and stored in -80°C until assayed. Hormone measurements included DHEAS, SHBG, testosterone, E2, LH, FSH, T3, T4, TSH, FT3, FT4, insulin, prolactin and were performed on an automated routine immunochemical analyzer COBAS e-411 (Roche Diagnostics, Manheim, Germany) that employs electrochemiluminescence.

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Serum OPG and sRANKL were measured by commercial ELISA kits (Biomedica Gruppe, Vienna, Austria). The limit of detection for OPG was 0.14 pmol/L and that of sRANKL 0.02 pmol/L. Both kits show inter-assay precisions of less than 8% in their performance characteristics.

Genomic DNA isolation

From an additional blood sample collected in EDTA tubes, genomic DNA was isolated from 200 μL by the QIAamp DNA Blood Mini kit (Qiagen, Germany) according to the manufacturer’s instructions. DNA purity and quantity were determined by fluorescence readings with the use of Quant-iTdsDNA-BR kit in the Qubit 1.0 fluorimeter (Life Technologies Invitrogen, USA). Extracted DNA was stored in -20°C until assayed.

Genotyping

Four intronic SNPs (rs2304188, rs11671439, rs3730051 and rs8100018) spanning the entire 51.5-kb genomic length of \textit{AKT2} gene were selected for evaluation of their association with the disease (Table 1). We developed novel real-time qPCR methods for their genotyping by using either the dual or single hybridization probe format. The sequences of the primers and probes of the proposed real time PCR assays synthesized by TIB MOLBIOL (Berlin, Germany) are shown in Table 1.

| SNP      | Location | PCR primers | Probes                  | Tanneal |
|----------|----------|-------------|-------------------------|---------|
| rs2304188| Intron 10| GTGGGGGTTAGGAACGTG | CCCCGAGCCCTGTGTTT GGA | 56      |
|          |          | GAGGCTGGCATTACAGTG | GTTTTCTGCCACACAC     |         |
| rs11671439| Intron 1 | CTCGCAATTAGCCTGTGATGAA | GGGCTCTTTCTGCCCACAC-FL | 59      |
|          |          | CTCCTAGACCTTCTCAAGGATGAC | ACCTTGGTTAACCCTATGC |         |
| rs8100018| Intron 4 | GAAAGAGCCGTCATGGA | AAAAAAXIAGACGGGGCGGGGAC | 53      |
|          |          | ACTTGTCTAACCCTATGC |                          |         |
| rs3730051| Intron 8 | TTGTGATGAGGATGACGCACAGC | ACGAGGCGCCCTGCCTACAGG-FL | 59      |
|          |          | TGGGCAAGCCACTTAACCT | LC705-GAGGGGACGATGACACTGA |         |

Table 1: Primers and probe sequences used in \textit{AKT2} genotyping analysis. The SNP position is underlined. Probes were labeled either with fluorescein (FL) or with LC 640 or 705 dyes. ‘XI’ denotes FL.

Real-time PCR was performed in glass capillary tubes in the LightCycler 1.5 instrument (Roche Applied Science, Switzerland). The amplification mixture of a total volume of 10 μL included 1 μL of genomic DNA extract as template (~30 ng), 2 pmol/μL forward primer, 2pmol/μL reverse primer, either 0.1 mM Simple probe or a combination of 0.15 μM FL-Sensor probe with fluorescein and 0.15 μM Anchor probe labeled with LC640 or LC705 dye (in the case of dual probe format) and 1 μL of 10× FastStart DNA Master Hybprobe mix (Roche Applied Science, Switzerland). We added MgCl2 to a final optimal 3 mM concentration. Sterile H2O was used to supplement up to 10 μL. The cycling protocol consists of pre-incubation at 95°C for 10 min for hot-start activation, followed by 40 cycles of denaturation at 95°C for 10 secs, annealing at a specified temperature (Table 1 for Tanneal) for 20 sec and extension at 72°C at 30 sec. Emitted fluorescence was measured at the end of each annealing step at a selected fluorescence channel (F1 for simple probes, F2 for LC640 or F3 for LC705 anchor probe). Immediately after amplification, melting curve analysis was performed on the LightCycler. The melting curve protocol included raising the temperature at 95°C for 10 sec, cooling at 40°C for 120 sec and slow heating to 85°C at a rate of 0.3°C/sec, during which time fluorescence measurements were continuously collected in the selected fluorescence channel. The first derivative plot (-d(F)/dT vs. T) is then used for easy identification of wild-type and mutant alleles by their different peaks, as the mismatched probe will dehybridize at an earlier stage.

DNA Sequencing

For the verification of the real-time qPCR results, the Gold Standard method of DNA Sequencing was used for method comparison. After the purification of the amplicons prepared from conventional PCR reactions (High Pure PCR Cleanup Micro kit, Roche Applied Science, Switzerland), cycle sequencing reaction were performed with the Big Dye 1.1 reagent in both directions with the use of either the forward or the reverse primer (ThermoFisher Applied Biosystems, USA). 10 μL of the purified cycle sequencing reactions (by NucleoSeq columns, Macherey-Nagel, Germany) were heated at 95 for 2 min and cooled immediately at 4 for 2 min with 10 μL formamide and then run in capillaries of the ABI Prism 310 Genetic Analyzer. For the analysis of DNA sequencing electropherograms the Chromas 2.01 software was used (Technelysium Pty Ltd, Australia) and results were compared with the expected gene sequences with the NCBI BLAST tool (https://blast.ncbi.nlm.nih.gov).

Statistics

Genotyping statistical analysis was performed through the SNPStats Internet platform (http://bioinfo.iconcologia.net/snpstats/start.html). The variants were tested for Hardy-Weinberg equilibrium (HWE) in either POCS patients or controls for each studied SNP, tables with genotype and allele frequencies were constructed and Odds Ratios (OR) and 95% confidence intervals (CI) were calculated.

Further analysis was performed with SPSS v.22 (IBM, USA). Descriptives were calculated for all variables. The distribution of continuous variables was assessed with Kolmogorov-Smirnov test.
Comparisons between continuous variables non-normally distributed were performed with Mann-Whitney U. Dichotomous comparisons were performed using Fisher's exact test. The level of significance was set at p ≤ 0.05.

Results

Cases and controls were matched for age and BMI (Table 2, p>0.05, Mann Whitney). Almost all PCOS clinical characteristics except amenorrhea were significantly more frequent for cases than controls as calculated with Fisher’s exact test (Table 2). Cases had significantly higher values for sRANKL (p<0.000), DHEAS (p<0.000), 17-OH-progesterone (p<0.000), and testosterone (p=0.021), and significantly lower values for SHBG (p=0.003), E2 (p<0.000) and PRL (p=0.012) in comparison to controls (Table 3).

Table 2: Differences in clinical characteristics between PCOS cases and controls.

| Cases          | Controls         | p     |
|----------------|------------------|-------|
| Mean (SD)      | Mean (SD)        |       |
| Age            | 28.18 (3.53)     | 29.07 (3.53) | 0.289 |
| BMI            | 24.63 (4.76)     | 24.71 (3.81) | 0.585 |
| %              |                  |       |
| Amenorrhea     | 15               | 3.33  | 0.000 |
| Oligomenorrhea | 40               | 3.33  | <0.000 |
| Dysmenorrhea   | 60               | 0     | <0.000 |
| Acne           | 60               | 0     | <0.000 |
| Hirsutism      | 63.3             | 0     | <0.000 |
| Oily skin      | 40               | 6.66  | <0.000 |

All novel AKT2 gene genotyping methods were optimized and developed satisfactorily and were validated for their accuracy with the Gold Standard DNA sequencing (100% concordance). Characteristic figures with amplification plot, melting curves with Tm (melting point) allele determination and DNA Sequencing electrophore grams e.g., for SNP rs3730051 are shown in Supplementary Figures 1,2,3 respectively. Each real-time PCR reaction included a blank and either a homozygote or a heterozygote positive control.

Table 3: Differences in serum biochemical parameters between PCOS cases and controls. * denotes statistical significance.

|                | Controls | Cases | p     |
|----------------|----------|-------|-------|
| Mean (SD)      | Mean (SD)|       |       |
| sRANKL         | 0.01     | 0.47  | p<0.000* |
| OPG            | 2.99     | 2.76  | p=0.205 |
| DHEAS          | 212.83   | 325.20| p<0.000* |
| SHBG           | 66.63    | 55.90 | p=0.003* |
| Testosterone   | 0.65     | 3.07  | p=0.021* |
| E2             | 58.79    | 46.20 | p<0.000* |
| LH             | 6.24     | 6.97  | p=0.583 |
| FSH            | 5.26     | 5.71  | p=0.127 |
| Prolactin      | 14.42    | 17.31 | p=0.012* |
| Insulin        | 8.62     | 10.66 | p=0.728 |
| 17-OH Progesterone | 0.30 | 0.96  | p<0.000* |

All SNP genotyping results were in HWE (p>0.05). Minor Allele frequencies (MAF%) were separately calculated for cases and controls and also compared to those given by the Genome Variation Server (GVS) from data collected by analyzing a minimum of 450 genomes per SNP. Only one of the SNPs showed statistical significant difference with PCOS: rs2304188 T minor allele increased from 9% in controls to 29% in patients providing an OR 4.04 (C.I. 1.12-14.54) for disease risk to the T allele carrier versus the wild-type carrier (Figure 1 and Table 4). Differences in MAF% between the GVS and the current study were significant for rs2304188 (controls only) and rs3730051 (both cases and controls).

Table 4: AKT2 gene SNPs descriptive statistics for PCOS case and control participants versus GVS data. * Fisher’s exact test between the study controls’ MAF% and GVS MAF%. ** Fisher’s exact test between the study cases’ MAF% and GVS MAF%. *** Fisher’s exact test between the study cases’ MAF% and controls’ MAF%.

| SNP            | Alleles | GVS MAF % | Control MAF % | Case MAF % | p* | p** | p*** | OR (C.I.) |
|----------------|---------|-----------|---------------|------------|----|-----|------|-----------|
| rs2304188      | C/T     | 24        | 9.4           | 26.7       | 0.014 | 0.870 | 0.016 | 4.04 (1.12-14.54) |
| rs8100018      | C/G     | 29        | 37.5          | 41.7       | 0.231 | 0.076 | 0.540 | 1.29 (0.56-2.99) |
| rs11671439     | C/T     | 18        | 18.8          | 23.3       | 1.000 | 0.484 | 0.480 | 1.37 (0.55-3.42) |
| rs3730051      | A/G     | 21        | 37.5          | 41.7       | 0.013 | 0.002 | 0.970 | 0.99 (0.50-1.95) |

Table 4: AKT2 gene SNPs descriptive statistics for PCOS case and control participants versus GVS data. * Fisher’s exact test between the study controls’ MAF% and GVS MAF%. ** Fisher’s exact test between the study cases’ MAF% and GVS MAF%. *** Fisher’s exact test between the study cases’ MAF% and controls’ MAF%.
Additionally, there were significantly more cases of hirsutism among those bearing the rs2304188 polymorphism (92.3%) than the wild-type carriers (63.4%, p=0.044). Regarding the entire population, participants that bear the rs2304188 polymorphism had significantly higher DHEAS (316.8 ± 100.1, p=0.051) and higher 17-OH progesterone (1.02 ± 0.68, p=0.009). Those with the rs11671439 polymorphism had significantly lower E2 values in the whole population (44.4 ± 13.0, p=0.040) but also when examining the patient group only (39.1 ± 10.6, p=0.025, Table 5).

Table 5: Statistically significant differences in serum hormone values for participants bearing various polymorphism of AKT2 gene.

| Wild-type | Polymorphism | p     |
|-----------|--------------|-------|
| Mean (SD) | Mean (SD)    |       |
| For the whole population |
| Differences for rs2304188 |
| DHEAS      | 278.1 (126.7) | 316.8 (100.1) | 0.051 |
| 17-OH-Progesterone | 0.64 (0.43) | 1.02 (0.68) | 0.009 |
| Differences for rs11671439 |
| E2         | 52.5 (15.0)  | 44.4 (13.0)   | 0.040 |
| Only for the patient group |
| Differences for rs11671439 |
| E2         | 48.8 (16.1)  | 39.1 (10.6)   | 0.025 |

Discussion

Although PCOS is believed to be one of the most common disorders of women, there are few data available regarding its prevalence [18-20]. In the present study, we evaluated the clinical, biochemical and genetic characteristics of women with polycystic ovary syndrome in the Greek population. A strong point of this study is the fact that all ultrasound examinations were performed at one institution, with the same transvaginal ultrasound device and by the same physician who strictly kept to the Rotterdam criteria for diagnosis of PCOS. Our data demonstrate significantly higher serum levels of DHEAS, total testosterone and 17-OH progesterone in the PCOS group compared to the control group, as expected from our well-ascertained patient population. Hyperandrogenemia is the biochemical hallmark of PCOS and women with such a disorder are expected to have elevated levels of DHEAS [21] and testosterone [22-23].

Also, our PCOS group had higher serum levels for sRANKL and significantly lower serum levels for SHBG, E2 and prolactin in comparison to controls. High serum levels of sRANKL have been shown to be a significant predictor of cardiovascular disease [24-25]. Many clinical studies have shown the connection between PCOS and cardiovascular risk due to a lipid/glucose altered metabolism, hypertension, systemic inflammatory condition and vascular injuries [26]. However, there was no significant difference between our two groups regarding OPG values. OPG is a dimeric glycoprotein of the
TNF receptor family with a molecular weight of 60kDa and 120 kDa respectively [15]. sRANKL is a member of the tumor necrosis factor (TNF) family, is the main stimulatory factor for the formation of mature osteoclasts and essential for their survival [23]. sRANKL produced by osteoblastic lineage cells and activated T lymphocytes activates its specific receptor RANK that is located on osteoclasts and dendritic cells [23]. OPG acts as a soluble secreted receptor for RANK and prevents the binding of RANKL to RANK [22,27]. Thus, OPG inhibits RANK activation on the osteoclast surface. The high serum level of sRANKL is a significant predictor of cardiovascular disease [24,25]. OPG has a number of biological functions such as anti-inflammatory properties and also a protective role of the cardiovascular system. All these functions of OPG are associated with PCOS. Different studies have previously evaluated OPG levels in PCOS patients; however, there is no agreement whether PCOS increases OPG circulating concentration or diminishes it [15]. In concordance with our study, other studies also reported no difference in OPG levels between the two groups [22]. However, there are studies that show either lower OPG levels in PCOS compared to controls [15, 23, 27] or higher OPG levels between the two groups [28]. Although there are many studies that focus on OPG circulating concentration in PCOS, the results vary and this may be due to lack of consensus regarding units, type of sample and antibodies used that target different epitopes in the different ELISA kits used. Since OPG exists both as a monomer and a dimer, its final concentration could differ significantly depending on the kit used.

PCOS seems to run in families. Although genetic factors play an important role in PCOS, the genetic basis of the syndrome remains controversial [29-31]. Our data shows that AKT2 gene polymorphisms could be associated with PCOS or with some of its features. In particular, rs2304188 SNP was found to be more frequent in PCOS patients compared to controls [OR 4.04 (C.I. 1.12-14.54)], whereas concerning the entire population, studied individuals that bear the rs2304188 SNP have higher values of DHEAS and 17-OH progesterone, both biomarkers of PCOS. The rest of the studied SNPs didn't show any correlation in contrast with the only other study –to the best of our knowledge- that has examined AKT2 polymorphisms in PCOS: they showed association with rs3730051 and rs8100018 with 8.7% of total) had the polymorphism of rs8100018 in combination with either rs11671439 or rs2304188 polymorphism and all of these eight participants were PCOS patients (five were rs8100018 SNP- rs11671439 SNP and three were rs2304188 SNP-rs8100018 SNP). However, since a sample of eight people cannot be considered sufficient for drawing reliable conclusions, further studies performing valid haplotype analysis in larger populations are necessary in order to confirm the present results and to firmly establish this genetic association.

Finally, it was found that eight participants (13.33% of cases and 8.7% of total) had the polymorphism of rs8100018 in combination with either rs11671439 or rs2304188 polymorphism and all of these eight participants were PCOS patients (five were rs8100018 SNP- rs11671439 SNP and three were rs2304188 SNP-rs8100018 SNP). However, since a sample of eight people cannot be considered sufficient for drawing reliable conclusions, further studies performing valid haplotype analysis in larger populations are necessary in order to confirm the present results and to firmly establish this genetic association.

Therefore, in our study, rs2304188 SNP shows statistical significant correlation with PCOS and some of its features and AKT2 gene polymorphisms seem worthy of further investigation in larger populations of PCOS. The studied SNPs could be tags for other coding SNP, could affect level of AKT2 mRNA transcript or its splicing; at least 12 different AKT2 splice variants have been detected. The developed AKT2 gene genotyping methods are fast, reproducible and accurate since method comparison with the Gold Standard DNA sequencing has shown 100% concordance. Also, the aforementioned methods are cost effective and since they use a closed-tube format, they are less prone to contamination.

The genetic etiology of PCOS is still unknown although it is known that a family history of PCOS is common among PCOS patients. This suggests an important genetic component although the mode of inheritance is not yet known. However, since our knowledge of the human genome expands rapidly and new polymorphisms are identified with the advent of novel massive parallel sequencing methods that can easily provide genetic information in panels of multiple genes, this could lead to detailed mapping of all disease loci for PCOS in the near future. This is very important since it will allow family screening and early diagnosis of women with PCOS as well as estimation of the potential risk of developing complications.

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