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Short title
African polygenic hazard score for prostate cancer

Keywords
Prostate cancer; health disparities; genome wide association study; polygenic risk; genomics; genotypic ancestry; African

Abbreviations

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<tr>
<td>HR</td>
<td>Hazard ratio</td>
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<tr>
<td>PHS</td>
<td>Polygenic hazard score</td>
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<td>PHS46</td>
<td>Previously published polygenic hazard score model using 46 SNPs</td>
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<tr>
<td>PHS46+African</td>
<td>Updated polygenic hazard score model for men with African genetic ancestry</td>
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<td>PRACTICAL</td>
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We identified three SNPs (rs76229939, rs74421890, and rs5013678), using cross-validation, that significantly improved performance of a previously published polygenic hazard model (PHS46) in men of African genetic ancestry. The performance of the novel PHS46+African score improved by as much as 79% after incorporation of rs76229939, rs74421890, and rs5013678. PHS46+African significantly improved association between polygenic risk with age at diagnosis of prostate cancer in Africans, a population generally under-served in genome-wide studies, to levels comparable with Europeans.
Abstract

Polygenic hazard score (PHS) models are associated with age at diagnosis of prostate cancer. Our model developed in Europeans (PHS46), showed reduced performance in men with African genetic ancestry. We used a cross-validated search to identify SNPs that might improve performance in this population. Anonymized genotypic data were obtained from the PRACTICAL consortium for 6,253 men with African genetic ancestry. Ten iterations of a ten-fold cross-validation search were conducted, to select SNPs that would be included in the final PHS46+African model. The coefficients of PHS46+African were estimated in a Cox proportional hazards framework using age at diagnosis as the dependent variable and PHS46, and selected SNPs as predictors. The performance of PHS46 and PHS46+African were compared using the same cross-validated approach. Three SNPs (rs76229939, rs74421890, and rs5013678) were selected for inclusion in PHS46+African. All three SNPs are located on chromosome 8q24. PHS46+African showed substantial improvements in all performance metrics measured, including a 75% increase in the relative hazard of those in the upper 20% compared to the bottom 20% (2.47 to 4.34) and a 20% reduction in the relative hazard of those in the bottom 20% compared to the middle 40% (0.65 to 0.53). In conclusion, we identified three SNPs that substantially improved the association of PHS46 with age at diagnosis of prostate cancer in men with African genetic ancestry to levels comparable to Europeans.
Introduction

Polygenic models can provide personalized estimates of the risk of developing prostate cancer. In the context of survival analysis, these models can provide insight into age at diagnosis of prostate cancer, and thus could be used to guide decisions on whether and when to offer screening. Studies of polygenic models have often included only individuals of European genetic ancestry, owing to greater availability of data from that population. As a consequence, these models have been tailored to identify and estimate coefficients of genetic common variants for that particular population, while potentially missing variants that may hold value in other populations. There is concern that using these European-focused models could actually exacerbate health disparities.

As an example, our group recently published on the performance of a polygenic hazard score (PHS) originally developed using a European dataset, in a multi-ethnic dataset consisting of individuals of European, African, and Asian genetic ancestry. The model (called here PHS46, referred to in the referenced manuscript as PHS2), includes 46 single nucleotide polymorphisms (SNPs) in its calculation and was strongly associated with age at diagnosis in all three genetic populations (p<10\(^{-16}\)). However, the hazard ratio for prostate cancer between individuals in the upper 20\(^{th}\) percentile to those in the lower 20\(^{th}\) percentile of PHS46 was approximately half as large for those with African genetic ancestry (2.6) as it was for those with European (5.6) or Asian (4.6)
ancestry. A similar pattern was observed for clinically significant prostate cancer and for death from prostate cancer.

In the current study, we attempt to bridge the apparent gap in model performance of PHS46 for individuals with African genetic ancestry. To this end, we used a machine learning approach to systematically search for SNPs that add statistical value to a base model of PHS46 among African men (PHS46+African). By including PHS46 as a covariate in our SNP search, we sought to identify those SNPs that may hold particular value for individuals with African genetic ancestry.

Material and Methods

Study dataset

We obtained genotype and phenotype data from the Prostate Cancer Association Group to Investigate Cancer Associated Alterations in the Genome (PRACTICAL) consortium for this study. Genotyping was performed using the OncoArray platform and had undergone quality assurance steps, as described previously. The study dataset contains no overlap with that used to estimate model coefficients of PHS46. It is a subset of another dataset wherein the performance discrepancy of PHS46 between different genotypic ancestries was first observed. All 46 SNPs of PHS46 were directly genotyped on the OncoArray platform.

The genotypic ancestry of each individual was also determined previously. In total, the African dataset consisted of data from 6,253 men with African genotypic ancestry. Missing SNP calls were replaced with the mean of the genotyped data for that
SNP in the African dataset. The percentage of individuals with missing SNP calls ranged from 0 to 5.9% across the SNPs, while the percentage of SNPs with missing calls ranged from 0.5% to 5.1% across the individuals. Individuals without prostate cancer were censored at age at last follow-up in the Cox proportional hazards models. A description of the PRACTICAL study groups that contributed data towards this analysis are described in Supplementary Table 1. PHS46 risk score for each individual in the African dataset was estimated as the sum of SNP allele counts (X) multiplied by their respective coefficients ($\beta$):

$$PHS46 = \sum_{i=1}^{46} X_i \beta_i$$

**SNP-scan**

A multi-step approach was used to select SNPs, from those directly genotyped on the OncoArray platform, that would improve the performance of PHS46 in the African dataset. Training and testing sets were generated using 10 iterations of a 10-fold cross-validation structure resulting in 100 total permutations. For each permutation, a multivariable logistic regression model using case/control status as the dependent variable was estimated using each genotyped SNP in turn, adjusting for PHS46 and four principal components based on genetic ancestry, determined previously. SNPs with p-values less than $1 \times 10^{-6}$ were considered for further analysis. In order of increasing p-value, each SNP was tested in a multiple Cox proportional hazards model, after adjusting for PHS46, four ancestral principal components, and previously selected...
SNPs. The Cox model in the SNP-scan used age at diagnosis of prostate cancer as the dependent variable. If the p-value of the coefficient of the tested SNP was less than $1 \times 10^{-6}$, it was considered for the final model in that permutation. SNPs that reached this p-value threshold in more than 50% of the permutations were selected to construct the PHS46+African model, consisting of PHS46 and the newly identified SNPs.

Comparing performance between PHS46 and PHS46+African – Hazard Ratio

For each permutation of the previously described cross-validation structure, an PHS46+African Cox proportional hazards model was estimated in the training set using PHS46 and the selected SNPs as independent predictors. The PHS46+African risk score for each individual is then estimated using the corresponding PHS46 score, selected SNP allele counts ($Y$) and their respective coefficients ($\alpha$):

$$PHS46 + African = PHS46 + \sum_{i=1}^{SNPs} Y_i \alpha_i$$

The performance of the PHS46+African and PHS46 models was then determined in the cross-validation testing set, and the resulting hazard ratios (HR) were obtained, as previously described\(^1\). For each model, the PHS risk scores within the cross-validation testing set are assigned to quantile groups identified using the corresponding training set control values. The hazard ratio between two quantile groups, such as those in the top 20% to those in the bottom 20%, is estimated as the exponential of the difference in mean PHS values for each group. In this calculation, the PHS values are linearly scaled by a sample-weight correction factor, to account for case-control sampling\(^1,5,10\). Three
HR were calculated: HR80/20 (top 20% to bottom 20%), HR98/50 (top 2% to middle 40%) and HR20/50 (bottom 20% to middle 40%). The average HR across permutations for both PHS46+African and PHS46 are reported.

To allow for comparisons with previously published results, the performance metrics for PHS46 and PHS46+African were also estimated for age at diagnosis of clinically significant prostate cancer. When estimating performance for clinically significant prostate cancer, controls and non-clinically significant cancers were censored at age of last follow-up and age of diagnosis, respectively. The previously used criteria for clinically significant cancer were any of: Gleason score $\geq 7$, stage T3-T4, PSA concentration $\geq 10$ng/mL, pelvic lymph nodal metastasis, or distant metastasis. Paired t-tests were used to test for statistically significant differences ($\alpha = 0.05$) in HR between PHS46+African and PHS46.

Additionally, in each permutation, the performance of a Cox model consisting of PHS46 and SNPs that were considered in that permutation was also estimated. These results are provided within the Supplementary Table 2 and provide performance estimates that are not prone to information leakage from training to testing set.

**Comparing performance between PHS46 and PHS46+African – C-index**

In addition to the hazard ratio, the performances of PHS46 and PHS46+African were compared using Harrell’s c-index. For each permutation of the aforementioned cross-validation structure, the c-index of PHS46 and PHS46+African scores were estimated in the testing fold using the “coxph” function in the R “survival” package.
Paired t-tests were used to test for statistically significant differences ($\alpha = 0.05$) between the two models.

**Characterization of PHS46+African**

Coefficients of the PHS46+African model, consisting of PHS46 and the SNPs selected in the SNP-scan, were estimated using 1000 bootstrapped samples of the African dataset.

**Clinical utility of PHS46+African**

As an example of the clinical utility of the PHS46+African risk score, the risk-equivalent age was estimated for those individuals in the upper 2 percentile of the distribution of PHS46+African risk scores. The risk-equivalent age, as defined previously, is when an individual from a given PHS percentile has prostate cancer risk equivalent to the average 60-year-old man. The age-specific general cumulative incidence curve was generated using data from SEER*Explorer incidence rates by age at diagnosis, 2003-2017 for Black Americans. The corresponding risk-adjusted incidence curve was estimated by multiplying the general cumulative incidence curve by the mean value of HR98/50 for PHS46+African obtained from the analysis of the age-of-diagnosis of prostate cancer. The risk-equivalent age was then calculated as the age at which the risk-adjusted cumulative incidence curve had the same value as the general cumulative incidence curve at age 60.
Results

Individual and OncoArray characteristics

In total, there were 3,013 men with (cases) and 3,240 men without (controls) prostate cancer in the African dataset. The mean [95% CI] ages of cases and controls were 62.4 [62.1, 62.7] and 61.8 [61.4, 62.1] years respectively. The OncoArray genotypic data, after the quality assurance process, included 444,323 SNPs.

Single Nucleotide Polymorphism (SNP)-scan

Across the 100 permutations of the cross-validation iterations, a total of twelve SNPs were considered for final selection (Supplementary Table 3). Three SNPs were selected in more than 50% of the permutations and included in the final PHS46+African model. By cross-referencing the chromosomal positions against dbSNP\textsuperscript{14}, these variants were identified as rs76229939\textsuperscript{15}, rs74421890\textsuperscript{16}, and rs5013678\textsuperscript{17}. All three SNPs (Table 1) are located on chromosome 8q24, a region of the chromosome previously identified as containing common variants associated with prostate cancer\textsuperscript{18,19}. An examination of the $R^2$ (Supplementary Table 4) showed little association, ranging from 0.0027 to 0.0057, among genotype data from the three SNPs in the African dataset.

Reference threshold (Supplementary Table 5) and mean (Supplementary Table 6) values for PHS46+African in the African dataset are presented in the Supplemental Data.
HR performance of PHS46+African

Figure 1 demonstrates the difference in HRs between PHS46+African and PHS46 within the African dataset using age at diagnosis of any prostate cancer (Supplementary Table 7). Overall, we observed an improvement in all the metrics calculated: a 75% increase in HR98/50 from 2.10 to 3.67; a 79% increase in HR80/20 from 2.47 to 4.42; and a 23% decrease in HR20/50 from 0.65 to 0.51. We also observed improvements in all performance metrics when using age at diagnosis of clinically significant prostate cancer: 103% increase in HR98/50 from 1.91 to 3.88, 113% improvement in HR80/20 from 2.21 to 4.71, and 29% improvement in HR20/50 from 0.70 to 0.50. All observed changes in HR were statistically significant (p < 1x10^-16).

C-index of PHS46+African

The mean c-indices of PHS46 and PHS46+African across the cross-validation folds were estimated as 0.55 and 0.58 (p < 1x10^-16), respectively.

Risk-equivalent age for PHS46+African

The risk-equivalent age for those individuals in the top 2 percentiles of the distribution of PHS46+African scores was estimated as 50 years old, suggesting that a man with a PHS46+African score in the top 2 percentile reached a prostate cancer detection risk equivalent to that of a standard 60-year-old roughly 10 years earlier, at an age of 50 years. The corresponding risk-equivalent age when using PHS46 scores was 54 years.
Discussion

Using a cross-validated search of a dataset made up entirely of men with African genetic ancestry, we were able to identify three SNPs that substantially improved the performance of PHS46 in this population to levels that are comparable to those observed in Europeans and Asians. Performance improvements were observed in hazard ratios tracking risk between PHS groups, concordance-indices tracking the overall utility of PHS as a continuous variable, and risk-equivalent age tracking the potential clinical utility of PHS. The three SNPs, rs76229939, rs74421890, and rs5013678, are all located on chromosome 8q24 – a region of the genome where variants have been associated with prostate cancer in both the general population and specifically in men with African genetic ancestry\(^\text{19,20}\). Despite the relative proximity of the three SNPs on chromosome 8, their genetic data was not strongly associated in our dataset, suggesting that each SNP provides non-redundant information for an individual’s genetic score.

Each of the three SNPs have been previously identified in the literature to be associated with prostate cancer: rs76229939 is an intron variant of the prostate-cancer-associated transcript 2 (PCAT2) gene, while rs74421890 and rs5013678 are both non-coding transcript variants of the prostate-cancer-associated non-coding RNA 1 (PRNCR1) gene. The minor allele frequencies of rs76229939 and rs74421890 in Europeans, as reported by dbSNP\(^\text{14}\), are approximately zero to three decimal places, which may explain why they were not selected in the original formulation of PHS46.
This study is not meant to be an exhaustive search for all possible SNPs that are associated with the age of diagnosis of prostate cancer in individuals with African genetic ancestry. Our study is also limited by the small number of available observations relative to those often found in many genome-wide association studies, which can have tens or hundreds of thousands of individuals. However, we were able to extract information that is likely robust by employing a cross-validated search for those SNPs that specifically add value to the performance of PHS46, and not simply independently associated with prostate cancer. Future analysis will include a more detailed analysis of the 8q24 region, including SNPs that are imputed using TOPMed reference panels. We also note that no SNP score, including PHS46 and PHS46+African, has been shown to discriminate men at risk of aggressive prostate cancer from those at risk of indolent prostate cancer. Finally, the performance metrics reported in this study may be biased by the leakage of information across cross-validated folds of the data when identifying those SNPs to include in the final African-PHS model. This bias is expected to be similar for all SNPs and should not have influenced selection of the three SNPs included in the final model over those not selected.

In conclusion, we identified three SNPs (rs76229939, rs74421890, and rs5013678) on 8q24 that substantially improved the performance of PHS46 in a dataset of men with African genetic ancestry. The addition of these SNPs to the polygenic risk score substantially improved its association with age at diagnosis of prostate cancer in Africans, to levels comparable with those found in Europeans.
Ethics Statement

All contributing studies were approved by the relevant ethics committees; written informed consent was obtained from the study participants\textsuperscript{21}. The present analyses used de-identified data from the PRACTICAL consortium and have been approved by the review board at the corresponding authors’ institution.
Conflict of Interest:

All authors declare no personal or financial conflicts of interest for the submitted work except as follows. CCF is a scientific consultant for CorTechs Labs, Inc. RE reports honorarium as a speaker for GU-ASCO meeting in San Francisco Jan 2016, support from Janssen, and honorarium as speaker for RMH-FR meeting Nov 2017. She reports honorarium as a speaker at the University of Chicago invited talk May 2018, and an educational honorarium by Bayer & Ipsen to attend GU Connect “Treatment sequencing for mCRPC patients within the changing landscape of mHSPC” at ESMO Barcelona, Sep 2019. She reports member of external Expert Committee on the Prostate Dx Advisory Panel. OAA received speaker’s honorarium from Lundbeck, and is a consultant for Healthlytix. AMD reports that he was a founder and holds equity in CorTechs Labs Inc., and serves on its Scientific Advisory Board. He is a member of the Scientific Advisory Board of Human Longevity, Inc., and the Mohn Medical Imaging and Visualization Centre. He received funding through research grants from GE Healthcare to UCSD. The terms of these arrangements have been reviewed by and approved by UCSD in accordance with its conflict of interest policies. TMS reports honoraria, outside of the present work, from: University of Rochester, Varian Medical Systems, Multimodal Imaging Servcies Corporation; and WebMD. He reports research funding from NIH/NIBIB, U.S. Department of Defense, Radiological Society of North America, American Society for Radiation Oncology, and Varian Medical Systems.
Data Availability Statement

The data used in this work were obtained from the Prostate Cancer Association Group to Investigate Cancer Associated Alterations in the Genome (PRACTICAL) consortium. Readers who are interested in accessing the data must first submit a proposal to the Data Access Committee. If the reader is not a member of the consortium, their concept form must be sponsored by a principal investigator (PI) of one of the PRACTICAL consortium member studies. If approved by the Data Access Committee, PIs within the consortium, each of whom retains ownership of their data submitted to the consortium, can then choose to participate in the specific proposal. In addition, portions of the data are available for request from dbGaP (database of Genotypes and Phenotypes) which is maintained by the National Center for Biotechnology Information (NCBI):
Anyone can apply to join the consortium. The eligibility requirements are listed here: http://practical.icr.ac.uk/blog/?page_id=9. Joining the consortium would not guarantee access, as a proposal for access would still be submitted to the Data Access Committee, but there would be no need for a separate member sponsor. Readers may find information about application by using the contact information below:

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References


Table Legends

**Table 1. Characteristics of PHS46+African SNPs.** RS-ID, chromosome and base-pair position (based on version 37), effect and reference alleles, bootstrap-estimated beta, and effect allele frequencies in aggregated Africans from 1000 Genomes (referenced from dbSNP) of the three SNPs selected for addition to PHS46
Figure Legends

Figure 1 Comparison between PHS46 and PHS46+African. Mean hazard ratio metrics plotted for PHS46 and PHS46+African models in the African dataset. Improvements were observed in all performance metrics investigated. Error bars represent 95% confidence interval.
Appendix 1. Members of the PRACTICAL Consortium

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Appendix 2. Funding sources for the PRACTICAL consortium

CRUK and PRACTICAL consortium

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Information of the PRACTICAL consortium can be found at http://practical.icr.ac.uk/

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CPDR
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MOFFITT

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Table 1. Characteristics of PHS46+African SNPs. RS-ID, chromosome and base-pair position (based on version 37), effect and reference alleles, bootstrap-estimated beta, and effect allele frequencies in aggregated Africans from 1000Genomes (referenced from dbSNP) of the three SNPs selected for addition to PHS46.

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