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Prostate Cancer

# Rare Germline Variants Are Associated with Rapid Biochemical Recurrence After Radical Prostate Cancer Treatment: A Pan Prostate Cancer Group Study

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# Abstract

**Background:** Germline variants explain more than a third of prostate cancer (PrCa) risk, but very few associations have been identified between heritable factors and clinical progression.

**Objective:** To find rare germline variants that predict time to biochemical recurrence (BCR) after radical treatment in men with PrCa and understand the genetic factors associated with such progression.

*Design, setting, and participants:* Whole-genome sequencing data from blood DNA were analysed for 850 PrCa patients with radical treatment from the Pan Prostate Cancer Group (PPCG) consortium from the UK, Canada, Germany, Australia, and France. Findings were validated using 383 patients from The Cancer Genome Atlas (TCGA) dataset.

**Outcome measurements and statistical analysis:** A total of 15,822 rare (MAF <1%) predicted-deleterious coding germline mutations were identified. Optimal multifactor and univariate Cox regression models were built to predict time to BCR after radical treatment, using germline variants grouped by functionally annotated gene sets. Models were tested for robustness using bootstrap resampling.

**Results and limitations:** Optimal Cox regression multifactor models showed that rare predicted-deleterious germline variants in "Hallmark" gene sets were consistently associated with altered time to BCR. Three gene sets had a statistically significant association with risk-elevated outcome when modelling all samples: PI3K/AKT/mTOR, Inflammatory response, and KRAS signalling (up). PI3K/AKT/mTOR and KRAS signalling (up) were also associated among patients with higher-grade cancer, as were Pancreas-beta cells, TNFA signalling via NKFB, and Hypoxia, the latter of which was validated in the independent TCGA dataset.

**Conclusions:** We demonstrate for the first time that rare deleterious coding germline variants robustly associate with time to BCR after radical treatment, including cohort-independent validation. Our findings suggest that germline testing at diagnosis could aid clinical decisions by stratifying patients for differential clinical management.

**Patient summary:** Prostate cancer patients with particular genetic mutations have a higher chance of relapsing after initial radical treatment, potentially providing opportunities to identify patients who might need additional treatments earlier.

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# 1. Introduction

Prostate cancer (PrCa) is the most common cancer in men in the developed world. Although the majority of PrCa cases are diagnosed with low- or intermediate-risk disease, approximately 10% of patients develop metastatic disease with poor survival rates [1,2]. Genetic predisposition to the overall disease risk of PrCa of any severity is well researched; however, an understanding of potential heritable genetic factors contributing to tumour progression is limited [3].

Biochemical recurrence (BCR) is often used as a prostatespecific antigen (PSA)-based predictor of progression to poor prognosis phenotype and is observed in approximately 25% of patients after radical prostatectomy (RP) [4]. Identification of men at a high risk for progression to lethal disease and who are likely to relapse after primary treatment would present the possibility to triage treatment intensification using current or novel systemic therapies. Most research into BCR to date has focused on gene expression or mutational signatures in prostate tumour tissue or specific candidate genes only [5]. In this study, we investigate for the first time whether rare germline variants across the full exome are predictive of poor prognosis after radical treatment. This information could aid clinical management of the disease, particularly at diagnosis, pre- or posttreatment staging, and prognostication.

#### 2. Patients and methods

#### 2.1. Sequencing of DNA from PrCa patients

Whole-genome sequencing (WGS) data derived from whole blood samples were collated for PrCa patients from member countries of the Pan Prostate Cancer Group (PPCG, http://panprostate.org; Australia n = 133, Canada n = 288, France n = 15, Germany n = 230, and UK n = 184; Table 1, further characteristics in Supplementary Table 1). The study presented here combines data from patients following RP and a small subset of samples with radical radiotherapy (RT; 8%) from the Canadian study group. We refer to the samples collectively as having radical treatment.

Samples were collected according to the criteria outlined in the Supplementary material. Collection was subject to the International Cancer Genome Consortium (ICGC) standards of ethical consent. Collection and

Table 1 – Number of samples, genes, and variants contributed, by study, also showing the number of samples with high-Gleason score (>3 +
Gleason grade group 3-5), the numbers of samples in each set with biochemical recurrence (BCR), numbers associated with mutations that a
predicted deleterious, and how many of those are known deleterious/loss of function (LoF) mutations

Study	European Genome- phenome Archive ID	Samples used in study after QC (with BCR)	Samples with high Gleason score (with BCR)	Number of genes with predicted- deleterious mutations (LoF)	Number of predicted- deleterious mutations included in analysis (LoF)
Melbourne, Australian Research Group	EGAD00001004182	133 (79)	110 (70)	2917 (1884)	3728 (2473)
Canadian Prostate Cancer Genome Network	EGAD00001004170	288 (92)	63 (22)	4579 (2637)	5900 (3154)
French ICGC Prostate Cancer Group	EGAD00001003835	15 (10)	15 (10)	409 (255)	393 (243)
Germany ICGC Prostate Cancer Group – Early Onset	EGAD00001005997	230 (68)	85 (45)	3787 (2160)	4761 (2404)
CRUK-ICGC Prostate Group, UK	EGAC00001000852	184 (36)	63 (22)	3365 (2073)	4071 (2401)
Total		850 (285)	336 (169)	8455 (5792)	15,822 (9006)
QC = quality control.					

analysis of the Australian samples received institutional review board approval (Epworth Health 34506; Melbourne Health 2019.058). WGS was performed using Illumina technology to  $\geq 30 \times$  depth. The Burrows-Wheeler Aligner [6] was used to align sequencing data to the GRCh37 human genome (human\_g1k\_v37) with polymerase chain reaction duplicates removed [7]. Sequencing data have been deposited at the European Genome-phenome Archive (https://ega-archive.org; dataset IDs in Table 1) and are available upon request.

### 2.2. Variant calling

Variant calling was performed with the Genome Analysis Toolkit pipeline (GATK v4.0) [8] following GATK best practice recommendations for germline single nucleotide variant and indel calling (Supplementary material) [9,10], apart from for the German samples that were called using FreeBayes v1.1.0 [11] and processed as described by Gerhauser et al [12], normalised with vt v0.5 (Supplementary material) [13]. This analysis was restricted to variants within protein-coding transcript sequences according to GENCODE v29 [14].

### 2.3. Quality control, variant annotation, and prioritisation

Low-quality variants and samples were removed based on established quality control (QC) protocols [15-17]. We excluded samples from related individuals (using R package SNPRelate method identity by descent [18]) or those with non-European ancestry (using Principal Component Analysis relative to 2504 samples from the 1000 Genomes Project [19]). We used Picard tools v2.23.8 [20] to remove samples with a mean insert size of <250 bp, AT or GC dropout >5%, <95% aligned reads, >5% mismatch rate, <80% with  $\ge 20 \times$  coverage, or >5% missing call rate. Using verifyBamID v1.1.2 [21], we removed samples with >3% sample contamination. We excluded variants with a missing call rate in >5% of the samples, those with monomorphic loci, those in repetitive regions (simple repeats, segmental duplications, and centromeric regions), and those for which the ExAC minor allele frequency in any population was >1%. Of the submitted samples, 3% were excluded based on ancestry, whilst 2% were removed because of sequencing quality. One sample was removed due to relatedness.

Post-QC variants were annotated using the Variant Effect Predictor (VEP v101) and loss-of-function transcript effect estimator (LOFTEE) package [22]. For downstream analyses, we retained only variants categorised as deleterious/loss of function, comprising those with protein-truncating mutations (nonsense, frameshift, and splice site variants) occurring in the first 95% of the protein, as well as predicted-deleterious (PD) missense variants with a CADD PHRED score of >30 (Table 1) [23].

# 2.4. Pathways and gene sets

For a pathway-level analysis, all 50 "Hallmark" gene sets from GSEA MsigDB were considered (downloaded April 2017) [24], along with the BROCA extended panel of 66 genes and 175 curated DNA repair genes (DRGs; Supplementary Table 2) [16,17].

#### 2.5. Statistical analysis

#### 2.5.1. Software and libraries

All statistical analyses were applied using Python v3.8 [25]. Data in VCF format were converted using PyVCF v0.6.7 [26] and processed using pandas v1.3.0 [27], SciPy v1.4.1 [28], NumPy v1.18.3 [29], IPython v7.14 [30], and Scikits.bootstrap v1.1 [31]. Survival analysis for Cox's proportional hazard (PH) model and Kaplan-Meier estimates were performed using the Lifelines v0.25 package [32]. Tables and graphs were output using Matplotlib v3.3.4 [33], to\_precision [34], and Maftools v2.6.5 [35].

#### 2.5.2. Multifactor Cox regression

Analyses were performed on the combined post-QC dataset (Table 1) and a subset of patients with high Gleason score tumours, with models stratified by study to compensate for differing baseline hazards. Geneset predictors of the Cox PH model were generated by recording the presence of any gene with PD mutations in the selected gene sets across all samples. Pathological T stage had a baseline of stage 1-2 and a second group for stage 3-4. Clinical T stage was used for patients receiving RT. Preoperative PSA and age at the time of surgery were continuous variables. Gleason score had a baseline of  $\leq 3 + 4$  (Gleason grade group 1– 2) and a group for  $\geq$ 4 + 3 (Gleason grade group 3–5). Time was measured from radical treatment until BCR, which for samples with RP was defined as two consecutive post-RP PSA measurements of >0.2 ng/ml on the last known follow-up date [36]. For the 72 Canadian samples with RT, BCR was defined as a rise in PSA concentration of >2.0 ng/ml above the nadir, backdated to first PSA >0.2 ng/ml if PSA continues to rise [37]. We performed a sensitivity analysis on a subset that excluded RT samples, which did not affect the significant risk-elevating gene sets observed (Supplementary Table 3).

Variables included in the final models were selected by performing Cox regression with penalisation based on the least absolute shrinkage and selection operator (LASSO) [38]. The optimal penalty factor (lambda) was determined as within 1 standard error of the optimum from the mean of 100 ten-fold cross-validation models. Only features with a nonzero coefficient were retained. The final prediction models were then built using Cox regression without penalisation.

	HR (95% CI)	p value	Bootstrap HR (95% CI)	Bootstrap p value
Gleason (≥4 + 3 : <4 + 3)	1.98 (1.47-2.67)	<0.001	2.01 (1.99-2.04)	<0.001
Stage (T3–T4 : T1–T2)	1.69 (1.29-2.21)	<0.001	1.75 (1.74–1.77)	<0.001
PI3K/AKT/mTOR signalling	1.55 (1.06-2.25)	0.023	1.58 (1.56-1.60)	0.012
Age	1.53 (1.20-1.96)	<0.001	1.03 (1.03-1.03)	0.001
Inflammatory response	1.35 (1.00-1.82)	0.048	1.37 (1.35-1.38)	0.028
KRAS signalling (up)	1.35 (1.01–1.79)	0.041	1.37 (1.36-1.38)	0.020
Fatty acid metabolism	1.29 (0.96-1.71)	0.087	1.32 (1.30-1.33)	0.040
G2-M checkpoint	1.25 (0.94-1.66)	0.13	1.27 (1.26-1.28)	0.074
Myc targets v2	1.23 (0.84-1.81)	0.3	1.26 (1.24-1.27)	0.16
Mitotic spindle	1.21 (0.94–1.56)	0.14	1.22 (1.21-1.23)	0.10
DRG	1.16 (0.90-1.51)	0.3	1.18 (1.17-1.19)	0.15
p53 pathway	1.16 (0.85-1.60)	0.4	1.18 (1.16-1.19)	0.2
IL-2/STAT5 signalling	1.06 (0.77-1.46)	0.7	1.07 (1.06-1.09)	0.4
Preop PSA	1.04 (1.01-1.06)	0.006	1.01 (1.01-1.01)	0.004
Coagulation	1.01 (0.76-1.36)	0.9	1.01 (1.00-1.02)	0.5
Glycolysis	0.81 (0.61-1.08)	0.16	0.82 (0.81-0.82)	0.080
UV response (dn)	0.71 (0.51-0.99)	0.042	0.72 (0.71-0.73)	0.038
Cholesterol homoeostasis	0.58 (0.34-1.00)	0.048	0.59 (0.58-0.60)	0.013

Table 2 – Multifactor Cox model results for predicted-deleterious mutations in 850 germline samples, grouped into 52 gene sets

BCR = biochemical recurrence; CI = confidence interval; dn = down; DRG = DNA repair gene; HR = hazard ratio; PSA = prostate-specific antigen. Shown are *p* values and hazard ratios of LASSO-selected gene sets as well as clinical variables reported at the time of BCR or last check-up, impacting the predicted time until BCR.

# 2.5.3. Univariate Cox regression

Each gene set was modelled individually along with clinical covariates (preoperative PSA, pathological T stage, Gleason score, and age) and p values were adjusted for multiple testing using false discovery rate (FDR).

#### 2.5.4. Validation

We performed harmonised variant filtering for PD mutations on germline PrCa samples from The Cancer Genome Atlas (TCGA) PRAD project. From the original 500 TCGA PRAD samples, those from contributing institutions with <15 samples were excluded, and models were stratified by institution, resulting in 383 samples used in the analysis. Of these, 233 were included in the high-Gleason subset analysis. We applied the variants to the predictors selected from the Cox model built using the combined PPCG samples, to compare the hazard ratios (HRs) in both sets.

# 2.5.5. Kaplan-Meier analysis

A Kaplan-Meier plot measuring time to BCR in the event of a relapse was used to visualise the impact of mutations within significant gene sets on the risk of BCR. This was applied separately to the whole dataset and the high-Gleason subset, and reported alongside log-rank test *p* values.

We performed a combined analysis, considering mutations in any of the gene sets significant for the corresponding analyses and subdivided to ascertain potential additive effects upon a patient's time to relapse.

#### 2.5.6. Bootstrapping validation

To test model robustness, we produced new datasets of the same sample size by randomly choosing samples with replacement, without stratification, and building a Cox regression model from the resulting dataset. This was repeated 1000 times to derive a distribution of coefficients. The p values were computed for each predictor as a percentage of the iterations where the coefficient was in a different direction than expected.

# 3. Results

We analysed germline WGS data from 850 patients across five studies in the PPCG consortium (Table 1 and Supplementary Tables 1 and 2) for germline predictors of PrCa progression measured by BCR after radical treatment. This analysis was restricted to variants within protein-coding transcript sequences, resulting in 15, 822 rare variants identified as deleterious or likely deleterious, jointly categorised as PD. No individual variants or genes demonstrated a significant association with time to BCR (Cox regression analysis; p > 0.05), although the available sample size of 850 cases is underpowered for such an analysis. Therefore, we focused on finding gene sets or pathways with significant associations to identify potential biological mechanisms linked with progression. To this end, we determined whether there was at least one PD germline alteration in 52 gene sets, including 50 Hallmark gene sets from the MsigDB database [24], containing over 4000 genes with sets varying in size from 30 to 200, the DRG panel containing 175 DNA repair genes [16], and the extended BROCA gene panel containing 65 genes (Supplementary Table 4) [17].

After variable selection by LASSO, the optimal model for predicting time to BCR contained 14 gene sets, three of which were significantly associated with time to BCR (Cox PH model for all samples; p-value threshold <0.05; Table 2 and Fig. 1A). Clinical variables at the time of radical treatment (preoperative PSA, pathological T stage, age, and Gleason score) were added to the model as covariates. The significant risk-elevating Hallmarks were PI3K/AKT/mTOR (HR = 1.55; 95% confidence interval [CI] 1.06-2.25; p = 0.023), Inflammatory response (HR = 1.35; 95% CI 1.00-1.82; p = 0.048), and KRAS signalling (up) (HR = 1.35; 95% CI 1.01–1.79; p = 0.041). These gene sets are associated with shortened average time to BCR. The UV response (down) (HR = 0.71; 95% CI 0.51–0.99; p = 0.042) and cholesterol homoeostasis (HR = 0.58; 95%) CI 0.34–1.00; p = 0.048) gene sets were borderline significantly protective. Application of this model to multiple bootstrap resamplings showed that these results are robust, with all risk-elevating gene sets having HR >1 in >97% of resamples and *p* values indicating the same coefficient direction.

The clinical covariate-only model built using all the samples determined that Gleason score, preoperative PSA, age,





	HR (95% CI)	p value	Bootstrap HR (95% CI)	Bootstrap <i>p</i> value
Pancreas-beta cells	2.52 (1.01-6.29)	0.047	3.58 (3.43-3.73)	0.034
PI3K/AKT/mTOR signalling	1.95 (1.21-3.15)	0.006	2.13 (2.09-2.17)	0.007
TNFA signalling via NFKB	1.79 (1.19-2.68)	0.005	1.86 (1.83-1.89)	0.005
Hypoxia	1.73 (1.14-2.63)	0.010	1.82 (1.79–1.85)	0.011
Stage (T3-T4 : T1-T2)	1.70 (1.13-2.56)	0.012	1.86 (1.84-1.89)	0.003
KRAS signalling (up)	1.58 (1.08-2.32)	0.019	1.65 (1.63-1.67)	0.016
Myc targets v2	1.54 (0.92-2.60)	0.10	1.60 (1.57-1.63)	0.081
DRG	1.33 (0.92–1.91)	0.13	1.38 (1.36-1.39)	0.071
G2-M checkpoint	1.31 (0.89-1.93)	0.17	1.41 (1.39-1.43)	0.092
Age	1.17 (0.90-1.52)	0.2	1.01 (1.01-1.01)	0.17
IL-6/JAK/STAT3 signalling	1.17 (0.69-1.98)	0.6	1.22 (1.19-1.24)	0.3
Preop PSA	1.06 (1.00-1.11)	0.039	1.00 (1.00-1.00)	0.028
Coagulation	1.05 (0.71-1.55)	0.8	1.08 (1.06-1.09)	0.4
mTORC1 signalling	0.79 (0.50-1.25)	0.3	0.80 (0.79-0.82)	0.17
Androgen response	0.71 (0.41-1.22)	0.2	0.73 (0.72-0.74)	0.12
Glycolysis	0.60 (0.40-0.91)	0.017	0.61 (0.60-0.62)	0.012
Cholesterol homoeostasis	0.564 (0.270-1.18)	0.13	0.586 (0.571-0.602)	0.063
CI = confidence interval: DRG = DNA repair gene: HR = hazard ratio: PSA = prostate-specific antigen.				

Table 3 – Multifactor Cox model results for predicted-deleterious mutations in 336 high-Gleason germline samples, grouped into 52 gene sets

Shown are p values and hazard ratios of LASSO-selected gene sets impacting the predicted time until biochemical recurrence.

Table 4 - Multifactor Cox model results for predicted-deleterious mutations in 233 high-Gleason TCGA germline samples, stratified by location and grouped into 52 gene sets

	HR (95% CI)	p value	Bootstrap HR (95% CI)	Bootstrap <i>p</i> value
Stage (T3-T4 : T1-T2)	7.85 (1.65-37.3)	0.01	$6.24 \times 10^{12}  (1.73 \times 10^8  3.73 \times 10^{13})$	0.001
Coagulation	3.53 (1.30-9.59)	0.014	11.3 (7.47–28.5)	0.022
Hypoxia	3.18 (1.04-9.74)	0.043	$7.88 imes 10^{6}(1.14 imes 10^{6}  extrm{}3.40 imes 10^{7})$	0.097
Myc targets v2	2.90 (1.00-8.40)	0.049	5.63 (5.29-6.07)	0.044
TNFA signalling via NFKB	2.12 (0.78-5.79)	0.14	3.95 (3.51-4.97)	0.11
G2-M checkpoint	2.00 (0.79-5.05)	0.14	2.89 (2.75-3.11)	0.10
Androgen response	1.43 (0.52-3.97)	0.5	1.81 (1.70-2.00)	0.3
IL-6/JAK/STAT3 signalling	1.32 (0.36-4.77)	0.7	$2.86 imes 10^8~(5.67{-}1.71 imes 10^9)$	0.3
KRAS signalling (up)	0.97 (0.39-2.43)	>0.9	1.36 (1.29–1.46)	0.5
PI3K/AKT/mTOR signalling	0.70 (0.08-5.77)	0.7	$1.52 \times 10^6  (0.972  7.60 \times 10^6)$	0.3
DRG	0.68 (0.31-1.49)	0.3	0.72 (0.70-0.75)	0.18
mTORC1 signalling	0.46 (0.14-1.50)	0.2	0.46 (0.43-0.49)	0.075
Glycolysis	0.27 (0.07-1.09)	0.067	0.34 (0.31-0.36)	0.047

CI = confidence interval; DRG = DNA repair gene; HR = hazard ratio; TCGA = The Cancer Genome Atlas. Shown are p values and hazard ratios of the same predictors identified by the Pan Prostate Cancer Group Cox model (pancreas-beta cells and cholesterol homoeostasis were removed, as most samples had a mutation or no mutation in the gene set, respectively, which caused convergence errors).

and pathological T stage significantly associate with time to BCR (Cox PH; *p* value threshold <0.05; Supplementary Table 5). This model is significantly improved by the addition of the selected gene sets (likelihood ratio test p = 0.048; c-index 0.68 vs 0.66).

Within the PPCG set, patients presenting with highergrade localised PrCa (a subset of 336 patients where Gleason score was  $\geq 4 + 3$ ; Gleason grade group 3–5) had a higher proportion of BCR events (50.2% compared with 33.5% for all samples; Table 1). We developed an optimal multifactor Cox regression model (Cox PH; p value threshold <0.05; Table 3 and Fig. 1B) for this subset of high-Gleason samples with poorer prognosis disease. After feature selection by LASSO, we identified five significant riskelevating gene sets: Pancreas-beta cells (HR = 2.52; 95% CI 1.01-6.29; p = 0.047), PI3K/AKT/mTOR signalling (HR = 1.95; 95% CI 1.21–3.15; *p* = 0.006), TNFA signalling via NFKB (HR = 1.79; 95% CI 1.19–2.68; p = 0.005), Hypoxia (HR = 1.73; 95% CI 1.14-2.63; p = 0.010), and KRAS signalling (up) (HR = 1.58; 95% CI 1.08-2.32; p = 0.019). PI3K/AKT/mTOR has a higher HR and a lower p value than in the all-sample model. The Glycolysis gene set is shown here as significantly protective (HR = 0.60; 95% CI 0.400.91; p = 0.017). The bootstrap resamplings for the significant gene sets have the same coefficient direction in >96% of resamples.

After examining each gene set in individual univariate models with all samples, none was found to have a significant association with outcome after multiple testing correction (FDR; p value threshold <0.1; Supplementary Table 6). PI3K/AKT/mTOR signalling (q = 0.14), KRAS signalling (up) (q = 0.20), and TNFA signalling via NKFB (q = 0.16) had p values close to the significance threshold and achieve the threshold of significance in the high-Gleason subset (Table 3). In the high-Gleason subset, performing a logrank test on each gene set revealed four gene sets that had a significant association with time to BCR: TNFA signalling via NFKB (p = 0.027), PI3K/AKT/mTOR signalling (p = 0.025), KRAS signalling (up) (p = 0.013), and Pancreas-beta cells (p = 0.023). In the multifactor high-Gleason Cox model, these four gene sets are also statistically significant (Table 3), alongside Hypoxia.

Application of the all sample Cox multifactor model to the TCGA validation set results in two significant gene-set predictors that are not reflected in the PPCG data: Myc targets v2 (HR = 4.46; 95% CI 1.73–11.5; p = 0.002) and Coag-



Fig. 2 – Kaplan-Meier plot showing survival probability against time in months until biochemical recurrence for (A) all samples and (B) the 336 samples in the high-Gleason subset (Gleason score >3 + 4; Gleason grade group 3–5). The impact of mutations in significant sets are subdivided by samples with mutations in multiple gene sets. Log-rank tests for each category: in Figure A–1 (p = 0.63) and  $\geq 2$  ( $p = 2.88 \times 10^{-3}$ ); and Figure B–1 (p = 0.27), 2 ( $p = 8.55 \times 10^{-3}$ ), and  $\geq 3$  ( $p = 3.29 \times 10^{-3}$ ).

ulation (HR = 3.49; 95% CI 1.47–8.30; p = 0.005; Cox PH; p value threshold <0.05; Supplementary Table 7). Performing the same high-Gleason filtering on TCGA samples and applying that set to the high-Gleason PPCG model, three significant risk-elevating predictors are identified: Myc targets v2 (HR = 2.90; 95% CI 1.00–8.40; p = 0.049), Coagulation (HR = 3.53; 95% CI 1.30–9.59; p = 0.014), and additionally Hypoxia (HR = 3.18; 95% CI 1.04–9.74; p = 0.043; Cox PH;

*p* value threshold <0.05; Table 4 and Fig. 1C). The consistent significance and same direction of coefficient of Hypoxia in patients with more advanced disease are compelling evidence that germline variations in genes within this pathway contribute to clinical progression.

We used Kaplan-Meier plots to visualise the additive effect of mutations in the corresponding risk-elevating gene sets for the all-sample and high-Gleason sets (Fig. 2). In both plots, we show a significant difference in survival when multiple gene sets carry PD mutations. In the allsample analysis, 285 of 850 patients had a mutation in one significant gene set and 58 had mutations in two or more gene sets, whilst in the high-Gleason subset analysis, 129 of 336 patients had a mutation in one significant gene set, 36 had mutations in two gene sets, and 12 had mutations in three or more gene sets, which was the clearest detrimental impact (Fig. 2B).

To search for individual genes mutated more frequently in patients with BCR, we calculated the odds ratio (OR) between the BCR-positive and BCR-negative groups (Supplementary Table 8). Twelve genes within the significant gene sets for all samples (PIKFYVE, MYD88, CAB39, RPS6KA1, IRAK2, IL2RB, MSR1, ITGB8, PIK3R5, MMP10, HKDC1, and *RBM4*) and 17 genes within the significant gene sets in the high-Gleason subset (GAPDHS, GRHPR, PGM1, SELENBP1, NAGK, SLC6A6, PIKFYVE, MYD88, CAB39, RPS6KA1, DDX58, KYNU, NR4A1, DENND5A, MMP10, HKDC1, and RBM4) had an OR at least two-fold higher and a mutation count difference of >2 between samples with a mutation and BCR and those with a mutation and no BCR (Supplementary Table 9). The overwhelming majority (92.7%) of the PD mutations identified in these combined 22 risk-elevating genes are missense variants (Supplementary Fig. 2), although patients with BCR exhibited more protein-truncating variants (Supplementary Fig. 3) than those without BCR (Supplementary Fig. 4).

# 4. Discussion

The primary aim of genetic profiling of germline or tumour DNA is to aid clinical decisions, such as targeted screening of asymptomatic individuals and treatment options for cancer patients. Germline signatures in particular would have the advantage of helping stratify patients in both pre- and post-operative settings. Follow-up strategies and decisions on further treatments could be aided by predicting which individuals are likely to develop prostate tumours, progress to clinically significant disease, or relapse. To our knowledge, this study is the first to evaluate the association of rare germline mutations across the full exome as opposed to specific plausible candidate genes and provides evidence that germline mutation status is predictive for BCR after radical treatment for PrCa. Our multifactor Cox model identified that rare PD variants in three Hallmark gene sets are associated with time to BCR after radical treatment (PI3K/ AKT/mTOR, KRAS signalling (up), and Inflammatory response) and five gene sets associated with BCR in a subset of cases with more aggressive phenotype at diagnosis (PI3K/ AKT/mTOR, KRAS signalling (up), Hypoxia, TNFA signalling via NFKB, and Pancreas-beta cells). Importantly, we also show that these gene sets remained independent predictive biomarkers of time to BCR, over and above the inclusion of clinical variables. We further demonstrate that the Hypoxia gene set was replicated in an independent cohort of high-Gleason tumour cases from TCGA. With additional confirmation and refinement, these signatures could inform prognosis and clinical decision-making.

Among the gene sets associated with a greater risk of BCR in PrCa patients, genes involved in PI3K/AKT/mTOR and KRAS signalling (up) remained significant across all PPCG samples as well as when restricted to patients with high-Gleason tumours. In somatic analyses, AKT expression and phosphorylation have previously been linked to the risk of BCR after RP [39,40] and poorer survival in patients with metastatic castrate-resistant PrCa [41]. Somatic loss of *PTEN*, a tumour suppressor that downregulates the AKT signalling pathway, is also associated with poorer prognosis PrCa [5] and disease recurrence [42,43]. The fact that these gene sets were not significant in the TCGA replication set could result from power limitations owing to the lower sample size (383 vs 850 samples), but these signatures will require validation in independent cohorts.

In the analysis of patients with high-Gleason tumours, the Hypoxia gene set was established at statistical significance in the PPCG cohort and also replicated in the independent TCGA validation cohort. This provides strong evidence that germline mutations within this gene set contribute to recurrence in patients with more aggressive disease. Hypoxia has previously been reported to contribute to progression when analysing tumour samples [44,45], with a 28-gene mRNA signature for hypoxia demonstrated to predict BCR and metastases after RP or RT and provide independent prognostic value after adjustment for clinical features [46]. Our results indicate for the first time that heritable mutations in genes upregulated in response to a low oxygen environment predispose PrCa patients towards a greater likelihood of, and shorter time to, BCR.

A small number of additional gene sets also achieved significance in a single analysis only (Inflammatory response in PPCG all samples, TNFA signalling via NFKB, and Pancreas-beta cells in the PPCG high-Gleason subset, and Myc targets v2 and Coagulation in the TCGA validation cohort). Owing to the less consistent selection of these gene sets, their importance in germline susceptibility towards BCR is less compelling; however, they would nonetheless represent potential gene sets of interest for examination in future larger replication studies.

In this study, we observed significantly shorter time to BCR among the individuals carrying mutations in more than one of the risk-increasing gene sets, compared with both noncarriers and individuals carrying mutations in a single gene set only. This provides further support that mutations affecting multiple regulatory networks may co-operatively serve to negatively influence PrCa prognosis, and that for some men, intraprostatic features that determine an aggressive tumour environment may be predetermined in the germline. This has been suggested before, based on hypoxia associating with genetic instability and aggressive subpathologies as field defects in PrCa, and warrants further investigation [47]. Fifty-eight out of the 850 total patients were carriers of mutations in two or more of the three allsample gene sets and 48 out of the 336 patients carried mutations within two or more of the five high-Gleason gene sets identified through our multifactor analysis.

The limitations of this study include multicohort biases; relatively small, European ancestry–only sample size, and in turn, limited statistical power to detect associations at the individual gene or variant levels; and the imperfect status of BCR as a definitive surrogate for clinical recurrence and survival. In addition, this analysis included only coding variants with strong evidence for deleterious effect, excluding variants of uncertain significance, copy number alterations, and structural variants. It may be necessary to integrate different data types, including expression and methylation data, to fully understand the mechanisms behind our findings. Although it is very encouraging that genes curated within PI3K/AKT/mTOR signalling and KRAS signalling (up) remained significant across both the PPCG all-sample and the high-Gleason subset analyses, and the independent validation cohort confirmed evidence for genes curated as involved in Hypoxia, additional larger studies remain necessary to confirm these findings and disentangle which specific genes contribute towards an increased risk of PrCa progression and invasiveness.

# 5. Conclusions

Our findings have potentially important clinical implications. Germline DNA can be sequenced at an early stage of disease or even for healthy individuals, which could enable the prediction of PrCa progression close to, or even in advance of, the point of diagnosis. This would allow clinicians to stratify and differentiate patients who are more likely to relapse, putting them on a different clinical treatment pathway comprising more radical intervention or more frequent follow-up.

**Author contributions:** Zsofia Kote-Jarai had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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# **Peer Review Summary**

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