Intrinsic Genomic Differences Between African American and White Patients With Clear Cell Renal Cell Carcinoma

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Abstract

Importance—There are well-documented racial disparities in outcomes for African American patients with clear cell renal cell carcinoma (ccRCC). Despite a dramatic improvement in overall survival in white patients since the advent of targeted therapy, survival for African Americans with advanced ccRCC has not changed. There is little known about potential racial differences in tumor biology of ccRCC.

Objective—to determine if there are racial differences in the somatic mutation rate and gene expression of ccRCC tumors from white and African American patients.

Design, Setting, and Participants—Overall, 438 patients with ccRCC were identified through The Cancer Genome Atlas (TCGA) clear cell kidney (KIRC) dataset (419 white and 19 African American patients). The GSE25540 dataset containing 135 patients (125 white and 10 African American patients) was used for validation. Tumor samples were collected from numerous cancer centers and were examined for racial differences in somatic mutation rates and RNA expression. Racial differences in somatic mutation rates and RNA expression were examined.

Main outcomes and Measures—The comparison of somatic mutation rates and differences in RNA expression in white and African American patients with ccRCC.

Results—Overall, 419 ccRCC tumor data sets from non-Hispanic white patients and 19 from non-Hispanic African American patients were identified through the publically available TCGA
KIRC data set, and a validation set of 125 white and 10 African American ccRCC patient tumors was identified from the publicly available GSE25540 data set. African American patients were significantly less likely than white patients to have \( VHL \) mutations (2 of 12 [17\%) vs 175 of 351 [50\%], respectively; \( P = 0.04 \)) and were enriched in the ccB molecular subtype (79\% in African American vs 45\% in white patients; \( P = 0.05 \)), a molecular subtype that carries a worse prognosis. It was found that RNA expression analysis revealed relative down-regulation of hypoxia-inducible factor (HIF) and vascular endothelial growth factor (VEGF)-associated pathways in African American patients compared with white patients.

Conclusions and Relevance—African American patients have less frequent \( VHL \) inactivation, are enriched in the ccB molecular subtype, and have decreased up-regulation of HIF-associated gene signatures than white patients. These genomic differences would predict decreased responsiveness to VEGF-targeted therapy and are a biologically plausible contributing factor to the worse survival of African American patients with ccRCC, even in the targeted therapy era.

Renal cell carcinoma (RCC) is the eighth most commonly diagnosed cancer in the United States. It accounts for nearly 64,000 new cancer cases and over 13,000 deaths per year in the United States alone.1 Multiple studies from the pretargeted therapy era demonstrate that African American patients with RCC have inferior overall survival compared with white patients regardless of age, sex, stage, histologic subtype, or surgical treatment.2-4 Despite this documented survival disparity, there is no data examining genomic or transcriptomic differences of clear cell RCC (ccRCC) in African American patients vs white patients.

The last decade has produced a wealth of knowledge about the molecular drivers of RCC, including the comprehensive molecular characterization of ccRCC by The Cancer Genome Atlas (TCGA).5 A hallmark event in the development of ccRCC is loss of heterozygosity of the von Hippel-Lindau (\( VHL \)) tumor suppressor gene. \( VHL \) inactivation occurs at a rate of 52\% to 82\% in sporadic ccRCC5 and leads to stabilization of the \( \alpha \) subunit of the hypoxia-inducible factor (HIF) family of transcription factors. Stabilization of HIF\( \alpha \) subunits results in their heterodimerization with HIF\( \beta \) subunits and a transcriptionally active complex whose target genes are intimately involved in tumor angiogenesis. The vascular endothelial growth factor (VEGF) pathway in particular is a key transcriptional HIF target and is targeted by a number of the currently US Food and Drug Administration-approved therapies. In addition, several other genes, such as \( PBRM1, SETD2, \) and \( BAP1 \), are also mutated in ccRCC patient tumors.6 Herein, we investigate potential genetic differences in ccRCC that are associated with race.

Methods

The Cancer Genome Atlas and Validation Data and Analysis

RNA sequencing, somatic mutation, and copy number data from the TCGA kidney clear cell (KIRC) data set were downloaded from the publically available TCGA data portal (https://tcga-data.nci.nih.gov/tcga/), and somatic mutation data was downloaded from an independent and publicly available data set from Peña-Llopis et al6 (GSE25540 data set).

The TCGA RNaseq data set was log2 transformed and median-centered. Two-class significance analysis of microarrays (using 2-fold change and false discovery rate = 0) was
performed to generate race-specific gene lists.\textsuperscript{7} The significant genes and corresponding fold changes as determined by significance analysis of microarrays were analyzed by Ingenuity Pathway Analysis (Ingenuity Systems) for predicted pathway activation and/or inhibition. Gene Set Enrichment Analysis (Broad Institute) was performed comparing ccRCC tumors from African American patients vs white patients (ethnicity defined by the TCGA) against Molecular Signature Database c2.all.v4.0. The mutations in the 9 most commonly mutated genes—\textit{VHL}, \textit{PBRM1}, \textit{SETD2}, \textit{KDM5C}, \textit{PTEN}, \textit{BAP1}, \textit{MTOR}, \textit{TP53}, and \textit{PI3KCA}\textsuperscript{7}—were analyzed between African American patients and white patients. Patients were then classified into the previously described RNA subtypes of ccRCC—ccA and ccB—based on patterns of differential gene expression using prediction analysis of microarray.\textsuperscript{8,9}

\textbf{Statistical Analysis}

Differences in somatic mutation rates by race were compared using $\chi^2$ or Fisher exact test where applicable. Fisher exact test was used to compare prevalence of the ccA and ccB molecular subtypes between races.

\textbf{Results}

We first investigated whether there are racial differences in the mutational spectrum of ccRCC tumors comparing data from African American patients (n = 19) and white patients (n = 419) from the TCGA ccRCC KIRC data set. Overall, 175 of 351 (50\%) white patients had \textit{VHL} mutations, whereas 2 of 12 (17\%) African American patients had \textit{VHL} mutations ($P = .04$) (Figure 1A) (eTable 1 in the Supplement). In contrast, there were no racial differences in the mutational frequency of other TCGA KIRC–defined significantly mutated genes (Figure 1A) (eTable 1 in the Supplement). These results were validated for \textit{VHL}, \textit{PBRM1} and \textit{BAP1} in an independent data set verifying the lower prevalence of \textit{VHL} mutation in African American patients (Figure 1B) (eTable 1 in the Supplement).\textsuperscript{6}

Given the lower frequency of \textit{VHL} mutations in African American patients, we investigated the RNA expression of the VEGF ligands and VEGF receptors to evaluate for downstream effects of \textit{VHL} loss. In the TCGA KIRC data set, 419 white patients and 19 African American patients were available for analysis. In the TCGA data set, African American patients had significantly lower expression of the \textit{VEGFA} ligand (eFigure 1A in the Supplement), as well as the FLT-1 (\textit{VEGFR1}) and KDR (\textit{VEGFR2}) receptors (eFigure 1B in the Supplement). Furthermore, significance analysis of microarrays and subsequent ingenuity pathway analysis of upstream regulators predicted that numerous genes associated with HIF (\textit{EPAS1}, \textit{HIF1A}, \textit{ARNT}, and \textit{CREB1}) transcriptional regulation were up-regulated in white patients (eTable 2 in the Supplement). Finally, several VEGF and HIF signatures were negatively enriched in African American patients as determined by Gene Signature Enrichment Analysis (eFigure 3 and eTable 3 in the Supplement). In aggregate, these findings indicate distinct differences in the biology of ccRCC in African American patients compared with white patients and suggest that ccRCC in African American patients may be less dependent upon HIF and VEGF signaling.

Clear cell renal cell carcinoma is a heterogenous disease that can be classified into 2 distinct molecular subtypes, ccA and ccB,\textsuperscript{8} with the ccB subtype associated with a worse survival.\textsuperscript{8,9}
Classification of the TCGA KIRC tumors into ccA and ccB subtypes showed that African Americans had significant enrichment for the ccB subtype (15 of 19 [79%] African American patients have the ccB subtype compared with 190 of 419 [45%] white patients; \( P = 0.05 \)) (Figure 2A) (eFigure 3 in the Supplement) and in general had high correlation to the ccB centroid (Figure 2B). This predominance of the ccB subtype in African American patients and its association with worse survival suggests that the intrinsic biology of ccRCC in African American patients may contribute to their well-documented worse survival outcomes.

**Discussion**

This is the first report to our knowledge to examine genomic differences between ccRCC in African American patients vs white patients. Our results indicate that ccRCC tumors from African American patients exhibit a lower rate of VHL mutation and a correspondingly lower level of HIF and VEGF pathway activation. We therefore postulate that a significantly larger proportion of tumors from African American patients may have a HIF-independent and VEGF-independent propensity for aggressiveness, resulting in resistance to the commonly used VEGF-targeted therapies. While our results point toward a biologic rationale for the lack of improvement in survival of African American patients since the advent of targeted therapy,\(^{10}\) we recognize that there are also a host of other potential factors (eg, access to health care, time to diagnosis, and appropriate treatment) that likely contribute to the racial disparities seen in advanced ccRCC. Nonetheless, the results presented herein provide insight into the potential role that genomic variation plays in the disparity observed between races.

Finally, why African American patients with ccRCC have less frequent \( VHL \) mutations is unclear. One possibility is that the development of ccRCC in African Americans may have a different etiology. For example, patients with end-stage renal disease have a higher incidence of RCC than the general population,\(^{11}\) and ccRCC tumors developing in patients with end-stage renal disease have lower rates of 3p loss or \( VHL \) mutation.\(^{12,13}\) Given higher rates of end-stage renal disease in African Americans (likely a consequence of a higher incidence and earlier onset of hypertension),\(^{14}\) it is plausible that the lower rate of \( VHL \) mutation in ccRCC tumors from African Americans may be the result of increased rates of end-stage renal disease. Comprehensive population-based studies will be needed to define correlations between race and etiologic heterogeneity.

**Conclusions**

Despite an impressive improvement in overall survival in white patients since the advent of VEGF pathway–targeted therapy, the survival for African American patients with advanced ccRCC has not changed. We present evidence of distinct differences in tumor biology between African American patients and white patients that would predict for less responsiveness to VEGF-targeted therapy. While this is a plausible biologic explanation for the lack of improved survival in African American patients, it is likely that disparities in health care delivery contribute to survival differences as well.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References


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### Key Points

**Question**
Are there genomic differences between clear cell renal cell carcinomas (ccRCC) that arise in African American patients and white patients that may contribute to the disparity in survival outcomes observed between races?

**Findings**
Using data from the The Cancer Genome Atlas clear cell kidney project, we found that African American patients have a significantly lower rate of mutations of the von Hippel-Lindau tumor suppressor gene, as well as decreased hypoxia-inducible factor activation. In addition, African American patients are enriched for a previously defined RNA subtype (ccB) that is associated with worse survival outcomes.

**Meaning**
There are distinct racial differences in the biology of ccRCC tumors that may partially explain the well-documented racial disparity in survival outcomes of African American patients with advanced ccRCC.
Figure 1. Gene Mutations in Clear Cell Renal Cell Carcinoma Tumors by Data Set Source and Race
A, Mutations in significantly mutated genes according to the The Cancer Genome Atlas kidney clear cell data sets are shown by race, B, Frequency of VHL mutations in an independent data set are shown by race.
Figure 2. Incidence and Correlation to Centroid of Clear Cell Renal Cell Carcinoma Tumor Subtype

A, Visual representation of the incidence of ccA and ccB RNA subtypes of clear cell renal cell carcinoma tumor subtype by race, B, Distance and correlation to centroid of tumor subtype (ccA and ccB) by race.