Molecular Stratification of Clear Cell Renal Cell Carcinoma by Consensus Clustering Reveals Distinct Subtypes and Survival Patterns

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Abstract
Clear cell renal cell carcinoma (ccRCC) is the predominant RCC subtype, but even within this classification, the natural history is heterogeneous and difficult to predict. A sophisticated understanding of the molecular features most discriminatory for the underlying tumor heterogeneity should be predicated on identifiable and biologically meaningful patterns of gene expression. Gene expression microarray data were analyzed using software that implements iterative unsupervised consensus clustering algorithms to identify the optimal molecular subgroups, without clinical or other classifying information. ConsensusCluster analysis identified two distinct subtypes of ccRCC within the training set, designated clear cell type A (ccA) and B (ccB). Based on the core tumors, or most well-defined arrays, in each subtype, logical analysis of data (LAD) defined a small, highly predictive gene set that could then be used to classify additional tumors individually. The subclasses were corroborated in a validation data set of 177 tumors and analyzed for clinical outcome. Based on individual tumor assignment, tumors designated ccA have markedly improved disease-specific survival compared to ccB (median survival of 8.6 vs 2.0 years, P = 0.002). Analyzed by both univariate and multivariate analysis, the classification schema was independently associated with survival. Using patterns of gene expression based on a defined gene set, ccRCC was classified into two robust subclasses based on inherent molecular features that ultimately correspond to marked differences in clinical outcome. This classification schema thus provides a molecular stratification applicable to individual tumors that has implications for influence treatment decisions, define biological mechanisms involved in ccRCC tumor progression, and direct future drug discovery.

Keywords
cCRCC, microarray, gene expression profiling, molecular signatures, survival, PCA, robust clustering, logical analysis of data, LAD, ConsensusCluster

Introduction
Clear cell renal cell carcinoma (ccRCC) affects upwards of 50,000 patients annually. Most of these patients will present initially with localized disease, managed with surgery, but unfortunately, nearly a third will develop recurrence and succumb to their disease. ccRCC incidence has increased uniformly over the past 30 years, associated with stage migration toward lower stages, likely due to the increased detection of lesions incidentally. However, there has not been commensurate improvement in survival. ccRCC tumors have variable natural histories, and genetic strategies have been largely unhelpful in identifying patients with higher or lower risk for recurrence due to the overwhelming association of this cancer with von Hippel-Lindau (VHL) tumor suppressor gene inactivation.

The Fuhrman classification system stratifies ccRCC by tumor cell morphology: low-grade (grade 1), intermediate-grade (grades 2 and 3), and high-grade (grade 4) tumors, with corresponding association with RCC-related death. Prognostic scoring systems such as the UCLA Integrated Staging System (UISS) have been developed using these morphologic characteristics, tumor size, and patient performance status as well as the inherent characteristics of stage.
and nodal status.\textsuperscript{5,6} Other algorithms incorporate postoperative clinical information but have limited discriminative ability for the abundant intermediate-grade and intermediate-stage tumors, and they fail to account for molecular distinctions in tumors.\textsuperscript{7} The molecular basis of this diversity in clinical behavior is unclear and makes ccRCC a ripe target for investigating the nature of these heterogeneities.

Gene expression analyses have provided meaningful insight into the clinical heterogeneity of many solid tumors. Unsupervised clustering of gene expression data with supervised learning methods can provide powerful strategies to identify molecularly and clinically significant cancer subtypes.\textsuperscript{8-11} New unsupervised consensus ensemble clustering strategies have been developed that have successfully identified breast cancer subtypes correlated with clustering strategies have been developed that have successfully identified breast cancer subtypes correlated with distinctive clinical information but have limited discriminative power.\textsuperscript{7} The molecular basis of this diversity in clinical behavior is unclear and makes ccRCC a ripe target for investigating the nature of these heterogeneities.

In ccRCC, using traditional unsupervised gene expression analysis, we and others have demonstrated that two or more molecular subclassifications of this tumor type exist.\textsuperscript{16-20} Many prior investigations, however, have relied on preselected molecular features or clinical outcomes as the criteria to identify expression signatures and distinguish gene sets. This type of approach fails to permit the underlying tumor biology, through the molecular end products of genetic changes, to inform the formation of tumor subgroups. A robust molecular classification system that connects tumor biology with individual tumor behavior should identify a priori the inherent patterns of gene expression that classify samples into nonoverlapping sets with a high degree of accuracy.

To investigate the molecular features that best define subsets of renal cell carcinoma, we applied unsupervised consensus clustering to the gene expression data of ccRCC tumors, without applying biologic or clinical information. Two robust subtypes (we have designated ccA and ccB) with differentiating biological signatures could be distinguished using a small gene set defined by logical analysis of data (LAD). This gene set allows for assignment of individual tumors within the ccA/ccB classification scheme and is easily translatable to reverse transcription PCR (RT-PCR) technology. Validation in an independent data set demonstrated that ccA tumors have a markedly better prognosis than ccB and that the molecular subtype was significantly associated with survival in both univariate and multivariate analysis. The identification of two robust ccRCC subclasses, which can be assigned by a small but highly significant panel of gene features, will provide a biological resource for future ccRCC investigation, allow better prognostication of ccRCC, and supply a wealth of information for therapeutic decisions.

\section*{Results}

\textit{Identification of ccRCC subtypes.} Gene expression data were obtained for 48 ccRCC samples and 3 independent replicate sample preparations. A flowchart diagram depicting the analyses performed is presented in Figure 1.

First, we performed ConsensusCluster, an unsupervised ensemble clustering algorithm, on the ccRCC samples (Supplementary Table S1), yielding two subsets, designated ccA and ccB (Fig. 2A). Removing the independent replicates produced an identical clustering assignment of tumors (data not shown), further confirming the stability of these clusters. Neither cluster was caused by inclusion of normal tissue in the RNA extraction as normal kidney assorts independently of either cluster (Supplementary Fig. S2).

Representative samples within each cluster were used for the development of characteristic gene signatures and the decipherment of biological pathways. Samples whose membership shifted through multiple bootstrapped iterations were set aside for later classification. These “core” clusters included 39 of the original 51 samples and permitted tumors with best-patterned features to define the cluster. As Figure 2B shows, the core cluster samples split into two robust subtypes of ccRCC that are stable when $k$ (degrees of freedom) increases to $k = 3$ or $k = 4$ (Fig. 2C and D), suggesting that the optimal number of robust clusters in this data set is 2. These analyses demonstrate that ccRCC can be optimally clustered into two distinct subtypes (ccA and ccB), defined purely by molecular characteristics of the tumors.

\textit{Analysis of pathway differences between two core clusters.} The identification of subtypes provides an opportunity to identify biological differences within the spectrum of ccRCC. SAM (Significance Analysis of Microarrays) analysis identified 2,701 and 3,512 probes overexpressed in ccA and ccB, respectively (Fig. 3A and Supplementary Table S3). This result confirms the gene expression profile heterogeneity observed in previous studies.\textsuperscript{17-19,21} The functional classification program, DAVID, was used to functionally categorize the probes identified in our analysis. A demonstration of the gene ontologies and pathways found to be differentially regulated between ccA and ccB is provided in supplementary material (Supplementary Table S3). In addition, SAM Gene Set Analysis, a more statistically robust way of identifying correlated gene groups, was performed using curated gene sets, providing similar results (Supplementary Table S4). The most notable genes, gene sets, and gene ontologies associated with cluster ccA were involved in angiogenesis (Fig. 3B), the beta-oxidation pathway (Fig. 3C), organic acid metabolism, fatty acid metabolism (Fig. 3D), and pyruvate metabolism. In contrast, core cluster ccB tumors overexpressed genes associated with cell differentiation, epithelial to mesenchymal transition (EMT) (Fig. 3E), the mitotic cell cycle, transforming growth factor beta (TGF\beta; Fig. 3F), response to wounding, and Wnt targets (Fig. 3G).

\textit{Delineation of a gene set to stratify ccRCC into ccA and ccB.} To identify a feature panel that could accurately identify ccA and ccB tumors, we used LAD, which uses pattern
recognition and supervised learning to identify key discriminating elements and has been successfully implemented in several biomedical studies.\textsuperscript{13,14,22} Using the core ccA and ccB tumors, LAD patterns were identified and validated. Using these patterns, we identified 120 probes, consisting of 110 genes, valuable for cluster assignment (Fig. 4A, Table 1, and Supplementary Table S5). The LAD model (Supplementary Table S6) was applied to the 12 noncore samples from the original analysis and predicted cluster membership for 11 samples, 8 ccA and 3 ccB (Supplementary Table S7).

To confirm that the genes identified by LAD are differentially expressed ccA and ccB ccRCC subtypes within individual tumors, we tested primers for ccA overexpressed genes FLT1, FZD1, GIPC2, MAP7, and NPR3 on available tumor samples using semi-quantitative RT-PCR. Figure 4B demonstrates that each of these products can predict tumor classification for individual tumors. These results collectively indicate the potential for a limited gene set to correctly distinguish between the two ccRCC subtypes using RT-PCR, a platform immediately transferable to formalin-fixed, paraffin-embedded tissues.

Validation of ccRCC subtypes. To validate the presence of two ccRCC subtypes in a second, independent data set, we applied ConsensusCluster and the LAD probe set to 177 ccRCC microarrays generated using a different gene expression profiling technique.\textsuperscript{17} Figure 5 shows the same two strong clusters in the data, which remained stable when \( k \) was increased (data not shown). The clusters were assigned to ccA or ccB by comparison of gene expression patterns to those in the primary data set.

Assignment of individual tumors. Assignment of tumors to a subtype with Cluster3.0 (traditional heat maps) or ConsensusCluster requires the presence of other tumors.
Therefore, we used LAD score to separately assign each individual tumor in the validation data set to ccA or ccB, without assessing similarity to the rest of the tumors. Assignment was predicted for each sample 100 times with 80% pattern bootstrapping. A tumor was classified only if the assignment occurred in >75% of the prediction runs. Of the 177 ccRCC tumors, 83 were predicted to be ccA, 60 as ccB, and 34 remained unclassified with these stringent classification rules (Supplementary Table S8). When compared with the cluster assignment predicted by ConsensusCluster,
Figure 3. Pathway analysis of subtypes shows that ccA and ccB are highly dissimilar. (A) Heat map of the 6,213 probes differentially expressed between ccA and ccB as determined by SAM analysis; false discovery rate (FDR) < 0.000001. (B-G) Magnified heat maps of the genes from (A) that populate the ccA (B-D) or ccB (E-G) overexpressed Molecular Signatures Database curated gene sets of Brentani angiogenesis (B), beta-oxidation (C), HSA00071 fatty acid metabolism (D), epithelial to mesenchymal transition (EMT) up (E), transforming growth factor beta (TGFβ) C4 up (F), and Wnt targets (G).
we found a concordance of over 86%, thus validating LAD-predicted assignment as a sensitive measure of tumor assignment.

**VHL pathway analysis.** With the ability to assign individual tumors to ccA or ccB, we were able to further investigate an intriguing aspect of our pathway analysis. We had found that several of the pathways overexpressed in ccA tumors are typically considered as being perturbed in ccRCC (i.e., angiogenesis is considered a defining feature of ccRCC). A number of genes (e.g., EPAS1, EGLN3, PDGFC, HIG2, and CA9) tightly correlated with aspects of VHL inactivation and hypoxia inducible factor (HIF) signaling were found to be overexpressed in ccA relative to ccB.

We applied LAD analysis to our previously published data set that was well annotated for VHL inactivation. Of the 21 tumors, 10 were predicted to be ccA, 6 as ccB, and 5 as unclassified (Supplementary Table S9). In each category, there were VHL wild-type tumors, HIF1 and HIF2 overexpressing tumors, and HIF2-only overexpressing tumors. Our own analysis of VHL status also demonstrated the presence of VHL mutations and/or methylation in both the ccA and ccB clusters (Supplementary Table S1). These data suggest that ccA and ccB, despite having a similar frequency of VHL inactivation, have activation of different dominant biologic pathways, resulting in distinct patterns of gene expression.

**ccA and ccB have different survival outcomes.** Given that VHL is inactivated in tumors of both subtypes, we wanted to know whether the underlying differences in tumor biology would show survival differences. Cancer-specific survival and overall survival for the ccA and ccB classes from the 177 tumor validation set were plotted using Kaplan-Meier curves (Fig. 6 A and B), calculating 95% confidence intervals (Supplementary Table S10). For cancer-specific survival (Fig. 6A), the ccA subtype was associated with a highly significant survival advantage over ccB patients (P = 0.0002, median survival of 8.6 vs 2 years). At 5 years, cancer-specific survival was 56% in ccA patients and only 29% in ccB patients. Figure 6B shows the same trend for overall survival, with a significantly greater survival for ccA patients over ccB patients (P = 0.004, median survival of 4.9 vs 1.8 years). At 5 years, survival for ccA patients is 48% but only 23% for ccB patients.

**ccA/ccB subtype associates with clinical variables.** Fuhrman grade, tumor size (T stage), and performance status, the covariates in the UISS for predicting outcome in newly diagnosed patients, were evaluated and compared with our
Table 1. Logical Analysis of Data (LAD) Gene Set

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</table>

Probes identified through LAD to discriminate between ccA and ccB subtypes. All probes were significant at t test, P < 0.000001. Fold change was calculated as ccA/ccB. Full names, Unigene cluster IDs, and GenBank accession numbers are available in Supplementary Table S5.
molecular classification with regard to survival outcomes. As expected, molecular classification strongly associated with tumor stage \( (P = 0.009) \) and grade \( (P = 0.0007) \) but not performance status \( (P = 0.5684) \). Seventy-eight percent of grade 1 and 69% of stage 1 tumors clustered as ccA, while 65% of grade 4 and 58% of stage 4 tumors clustered as ccB tumors. As low-grade ccRCC tumors tend to have better prognosis and high-grade tumors poor prognosis,\(^4\) this result was expected. This observation also suggests that the biological characteristics responsible for grade and stage-specific prognosis in ccRCC are encompassed in the classification schema. Figure 6C demonstrates that the ccA/ccB subtype still significantly correlates with survival when limiting analysis to intermediate-grade (grades 2-3) tumors. As expected, a Kaplan-Meier curve limited to the highly aggressive grade 4 tumors shows a convergence of subtype-specific survival (Fig. 6D).

Molecular classification is independently associated with survival. To determine how our classification schema compares with current standard clinical parameters as a prognostic factor, univariate Cox regression analyses were performed (Table 2). Molecular subtype is strongly associated with survival, with a hazard ratio (HR) of 2.2 \( (P = 0.0003) \). Even in the absence of stage 4 (metastatic) tumors, subtype has a strong association with survival \( (HR = 2.143, P = 0.0233) \). In addition, the use of the Schwartz Bayesian criterion (SBC) suggests\(^24\) that whether the tumor is classified by ccA/ccB/unclassified, ccA/ccB, or LAD score, the measures are strongly associated with survival, with difference in adjusted SBC values of 8, 8.3, and 9, respectively. These results suggest that defining a tumor as ccA or ccB may be an important prognostic indicator for predicting outcome from patients with ccRCC.

Multivariate analyses were then performed to determine whether our classification schema was still independently associated with survival outcomes in the context of stage, grade, and performance status. The dichotomous classification of ccA/ccB provides a significant association with survival at the 0.1 level \( (P = 0.089) \), likely influenced by the smaller sample size of the 143 classified tumors. Increasing sample size to 177 by including unclassified tumors, the trichotomous classification increased significance to \( P = 0.0736 \). Statistical analyses
often show that continuous variables provide more statistical discrimination. In fact, LAD score is an independent predictor of survival \( (P = 0.0027) \) and is more predictive of outcome than Fuhrman grade \( (P = 0.0308) \). These data intimate that the classification schema presented in this article may provide independent prognostic information over and above that provided by standard clinical parameters.

**Discussion**

Unsupervised consensus clustering algorithms can identify distinct classifications of histologically similar tumors based on machine learning algorithms. In this analysis, a small gene set distinguishes two inherent molecular subtypes of ccRCC (ccA and ccB), characterized by divergent biological pathways and a highly significant association...
with survival outcomes. This unique analysis provides a powerful method to discriminate molecular subgroups of tumors that may be informative of tumor biology or influence tumor behavior.

A fundamental problem in gene expression analysis of human tumors is the measurement of genetic noise in pairwise comparisons across thousands of independent and dependent variables. Our combined use of principal component analysis (PCA), consensus clustering, and LAD is robust and, more important, identifies stable clusters within patterns of gene expression. This method is highly reproducible and able to classify samples into molecular and clinically meaningful categories. Within these categories, “core clusters” are sets of nonoverlapping samples that are distinguishable from each other with high accuracy. This method of tumor analysis permits a refined assignment into gene expression-defined classifications and yields predictive gene signatures based on a manageable sized number of gene features. These properties permit the identification of limited sets of highly predictive molecular features (i.e., genes) useful for the classification of individual samples outside of the primary analysis. The extension of biomarker molecular profiles to small groups of genes, which can assign classification to individual tumors, is a major step forward toward the development of a clinically relevant biomarker. Ultimately, such a classification scheme will be applied with such measures as quantitative RT-PCR.

The clinical heterogeneity of ccRCC, coupled with previous gene expression studies, suggests that at least two molecular subtypes of ccRCC exist. We demonstrated that there are likely only two primary subtypes of ccRCC stable under bootstrap analysis, although further subclassifications within these subtypes may be identified in much larger data sets, and rare tumors may represent unusual variants. Using the LAD predictions in the validation set, a third group of tumors shared pattern features with both ccA and ccB tumors. Such a third group, or other suggested classifications, may represent an intermediate manifestation of tumors undergoing progression from ccA to the ccB subtype or simply share common characteristics of both groups.

The subtypes ccA and ccB were associated with a significant difference in survival outcome, with ccA patients having a markedly better prognosis. While the continuous variable of LAD score proved to be an independent predictor of survival, the more immediately clinically useful dichotomous classification of ccA or ccB had a similar effect size and was statistically significant at the \( P = 0.1 \) level in the multivariable analysis. Future studies on larger numbers of patients are needed to validate the results of the preliminary multivariate analysis reported herein.

Pathway analysis showed that the better prognosis ccA group relatively overexpressed genes associated with hypoxia, angiogenesis, fatty acid metabolism, and organic acid metabolism, whereas ccB tumors overexpressed a more aggressive panel of genes that regulate VHL, the cell cycle, and wound healing. Intriguingly, ccA overexpresses genes associated with components of hypoxia and angiogenesis pathways, processes known to be broadly dysregulated in ccRCC. \( VHL \) inactivation and subsequent activation of the hypoxia response pathway is so highly correlated with ccRCC that many of these pathways are expected to be upregulated in virtually all ccRCC tumors. As expected, using both training set tumors and LAD assigned gene expression arrays from Gordan et al., we identified \( VHL \) inactivation in both clusters. Thus, ccB may have acquired additional genetic events that supplement \( VHL \) pathway events, contributing to a more biologically immature and aggressive phenotype that overwhelms the signature associated with \( VHL \) inactivation, which should be evaluated in future studies. In addition, it will be interesting in the future to determine if the key features that make up this classification are unique to ccRCC or if other histologic subtypes share the features of either the ccA or ccB classifications.

Finally, our small, robust panel of genes, whose expression levels can classify individual tumor samples into ccA and ccB subtypes with high accuracy, may provide a valuable resource for clinical decisions for patients following nephrectomy regarding frequency of surveillance or choices for adjuvant therapy in the future. This panel provides the basis for the development and validation by a prospective clinical trial to assign subtypes of ccRCC to individual tumor specimens for implementation in a prognostic algorithm.
Materials and Methods

Complete materials and methods can be found in the online supplementary material (Supplementary Data S11).

Samples. Fifty-one specimens from 48 ccRCC patients were collected from consenting patients undergoing nephrectomy for RCC from 1994 to 2008 (Supplementary Table S1), analyzed for quality, flash frozen, and accessed with appropriate institutional review board (IRB) approvals. The validation set of 177 cases was described previously. Survival data were updated with a median follow-up of 120 months (range, 66-271). The pVHL and HIF annotated data set was previously described.

Gene expression analysis. RNA was extracted using the Qiagen RNeasy kit (Valencia, CA), amplified, labeled, and hybridized against a reference on Agilent Whole Human Genome (4 × 44k) Oligo Microarrays. Expression data were tabulated, and missing data were imputed. Batches were combined using Distance Weighted Discrimination (DWD; https://cabig.nci.nih.gov/tools/DWD) and normalized. Data are posted on GEO (GSE16449). Gene expression data from the validation set were collected, GEO (GSE3538). Print runs were DWD combined and normalized. Gene expression data from the pVHL/HIF data set were posted on GEO (GSE11904).

Pathway analysis. Heat maps were generated using Cluster 3.0 (http://bonsai.ims.u-tokyo.ac.jp/~mdehoon/software/cluster/) and Java Treeview (http://jtreeview.sourceforge.net/). Genes were functionally annotated in DAVID (http://david.abcc.ncifcrf.gov/). SAM-GSA (http://sourceforge.net/) was performed using MSigDB curated gene sets (http://www.broad.mit.edu/gsea/mSigDB/).

PCA. ConsensusCluster was used for PCA (http://code.google.com/p/sensus-cluster/) and consensus clustering. Features whose coefficients were in the top 25% were selected from PCA eigenvectors representing 85% variation in the data, retaining 20 eigenvectors and 281 features.

Unsupervised consensus ensemble clustering. Consensus clustering was applied to PCA features to divide the data successively into $k = 2$, 3, 4 . . . clusters, with 80% bootstrapping of 300 subsamples of genes and/or samples. We applied two clustering techniques, K-Means and Self-Organizing Map.

LAD. Features mapped to genes that discriminate between the two subtypes $(t$ test, $P < 0.000001$) were retained. We then applied LAD (http://pit.kamick.free.fr/lemaire/software-lad.html). LAD patterns requiring only one gene for perfect discrimination were generated. LAD was reapplied to identify patterns of degree 1 and degree 2 (homogeneity and prevalence = 0.9). A classifier $CS = f_P - f_N$ assigned an unknown sample to a class, where $f_N/f_P$ is the fraction of negative/positive patterns satisfied. If the LAD score (CS) was negative/positive, the sample was predicted to class ccA/ccB, respectively.

Semi-quantitative RT-PCR. RNA from patient tumors (chosen by RNA or tumor availability) was reverse transcribed primarily using RNA extracted from a second sample of tumor. cDNA was amplified by 25 cycles of semi-quantitative PCR with primer sets for FLT1, FZD1, GIPC2, MAP7, NPR3 (http://www.idtdna.com/), or control 18S rRNA primers (Applied Biosystems, Foster City, CA). Full-sized gels are shown in Supplementary Figure S12.

VHL sequence and methylation analysis. DNA was extracted from tumor samples using proteinase K (Roche, Basel, Switzerland) and standard phenol/chloroform extraction. VHL exons were PCR-amplified and directly sequenced for mutations with a BigDye Terminator Cycle kit on a 3130xl sequencer (Applied Biosystems). Primers and protocols used were described previously. A CpG Wiz kit (Chemicon, Temecula, CA) and/or NotI digestion was used for methylation studies.

Statistical methods. Statistical analyses were performed using R v2.4.1 (http://www.r-project.org), SAS (SAS Institute, Cary, NC), or STATA (StataCorp, College Station, TX). Kaplan-Meier estimated the time-to-event functions of disease-specific and overall survival. Disease-specific or overall survival was time between nephrectomy to date of death due to disease or date of death, respectively. Log-rank test was used to test for differences between survival curves. Univariable logistic regression evaluated the association of covariates on the outcome probability of subtype ccA versus ccB. Univariable and multivariable Cox regression evaluated the association of individual and multiple covariates on disease-specific and overall survival. SBC assessed model fit.

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Declaration of Conflicting Interests

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