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Comprehensive Pan-Genomic Characterization of Adrenocortical Carcinoma

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SUMMARY

We describe a comprehensive genomic characterization of adrenocortical carcinoma (ACC). Using this dataset, we expand the catalogue of known ACC driver genes to include *PRKAR1A*, *RPL22*, *TERF2*, *CCNE1*, and *NF1*. Genome wide DNA copy number analysis revealed frequent occurrence of massive DNA loss followed by whole genome doubling (WGD) which was associated with aggressive clinical course, suggesting WGD is a hallmark of disease progression. Corroborating this hypothesis were increased *TERT* expression, decreased telomere length, and activation of cell cycle programs. Integrated subtype analysis identified three ACC subtypes with distinct clinical outcome and molecular alterations which could be captured by a 68 CpG probe DNA methylation signature, proposing a strategy for clinical stratification of patients based on molecular markers.

Graphical Abstract



INTRODUCTION

Adrenocortical carcinoma (ACC) is a rare endocrine malignancy with an annual incidence of 0.7–2 per million (Bilimoria et al., 2008; Else et al., 2014a). Current staging schemes such as by the European Network for the Study of Adrenal Tumors (ENSAT) broadly divides ACCs into 4 stages (Fassnacht et al., 2009). While stage I and II tumors are organ-confined and potentially curable by complete resection, advanced stage tumors are invasive (stage III) and/or metastatic (stage IV), with a dismal 5 year survival of 6–13% for stage IV patients (Else et al., 2014b; Fassnacht et al., 2009; Fassnacht et al., 2013). Histologic grading of ACC based on proliferation (Giordano, 2011; Weiss et al., 1989) refines treatment decisions for specific patient subgroups, but is not universally accepted (Miller et al., 2010). Chemotherapy, radiotherapy and the adrenolytic agent mitotane are the current standard therapeutic modalities for unresectable or metastatic ACC, but all are palliative (Else et al., 2014a). Our understanding of ACC pathogenesis is incomplete and additional therapeutic avenues are needed.

Molecular studies have nominated several genes as potential drivers involved in sporadic adrenocortical tumorigenesis, including insulin-like growth factor 2 (*IGF2*), β -catenin (*CTNNB1*), and *TP53* (Giordano et al., 2003; Tissier et al., 2005). β -catenin gain-of-function mutations are evident in approximately 25% of both benign and malignant sporadic adrenocortical neoplasms (Tissier et al., 2005). Recent genomic profiling efforts of ACC have identified candidate driver genes such as *ZNRF3* and *TERT*, and identified molecular subgroups with variable clinical outcomes (Assie et al., 2014; Pinto et al., 2015). Germline variants of these genes are also associated with Beckwith-Wiedemann, Familial Adenomatous Polyposis Coli and Li-Fraumeni syndromes, in which adrenocortical neoplasia occur.

Comprehensive and integrated genomic characterization of ACC may identify additional oncogenic alterations, provide a framework for further research, and guide development of therapies. As one of the rare cancer projects of The Cancer Genome Atlas (TCGA), we collected the clinical and pathological features, genomic alterations, DNA methylation profiles, and RNA and proteomic signatures of 91 cases of ACC. The comprehensive nature of the dataset combined with the many easy access routes to the individual and aggregated profiles will serve as a point of reference for many future ACC and comparative cancer studies. Here, we report this resource and an integrated analysis of the data.

RESULTS

Patient cohort, clinical annotation and analytical approach

We analyzed 91 histologically confirmed tumors and matched blood or normal tissue from a global cohort collected from 6 countries including 84 usual type, 4 oncocytic, 2 sarcomatoid and 1 myxoid variant. Median age at diagnosis was 49 years, and female:male ratio was 1.8. While 51 (56%) involved the left gland and 40 (44%) the right gland, more tumors in the left gland were diagnosed in males than females (72% vs 47%; p value=0.03, Fisher's exact test). The majority of tumors (57%) were functional; hypercortisolism was the most common endocrinopathy. Resected tumors spanned all stages (stage I, n=9; II, n=43; III, n=19; IV, n=17; 3 unavailable). Median overall survival was 78 months with 5-year survival of 59%. Locally invasive and metastatic tumors (grade III plus IV) had dramatically reduced median overall survival (18 months) with 5-year survival of 22%. Clinicopathologic data are summarized in Table S1. Patient stage at diagnosis and cortisol hypersecretion were predictive for both overall survival and disease free survival (Else et al., 2014b), but age was only associated with overall survival. Gender had no clear association with disease stage or clinical outcome (Table S1).

We generated a comprehensive molecular dataset of the 91 tumors, as follows: whole exome sequence (n=90), mRNA sequence (n=78), miRNA sequence (n=79), DNA copy number via SNP arrays (n=89), DNA methylation via DNA methylation arrays (n=79) and targeted proteome from reverse phase protein array (RPPA; n=45) (Table S1).

Our analytical approach consisted of four main parts. We began by identifying somatic single nucleotide variants, gene fusions, and copy number alterations. Given the significant copy number alterations observed, we performed an extended bioinformatic analysis that revealed the role of genome doubling and telomere maintenance in ACC. Using the various molecular datasets, we derived molecular classifications of ACC and integrated a three class solution with the somatic genomic landscape. We placed ACC in the broader context of cancer by performing several pan-cancer analyses. Finally, we quantified adrenal differentiation and tumoral hormone production across the entire cohort to correlate the genomic results with clinical parameters. All unprocessed data used in the analysis can be accessed through the TCGA portals.

Exome and RNA-sequencing nominate ACC driver events

Somatic single nucleotide variants (SSNVs) and small indels were detected using five independent mutation callers and variants called by at least three callers or independently validated by RNA sequencing were included (Figure S1A and S1B). This approach yielded a total of 8,814 high-confidence mutations (6,664 nonsynonymous and 2,150 synonymous; 3,427 of these were found in two tumors with ultramutator phenotype. Deep coverage resequencing (1,500×) achieved a validation rate of 95% (Supplemental Experimental Procedures and Table S1). The median somatic mutation density was 0.9 per Mb, similar to pancreatic adenocarcinoma but twice more than that of another endocrine tumor, papillary thyroid cancer (Figure S1C). Similar to other cancers, ACC demonstrated a marked heterogeneity in mutation density (range: 0.2 to 14.0 mutations/Mb, excluding the two ultramutators). Mutation density was correlated with numerous clinicopathological parameters including overall survival, time to recurrence, necrosis, stage, Weiss score, and mitotic count (Figure S1D). Interestingly, these associations remained when restricting to organ-confined stage I and II diseases.

We used MutSigCV (Lawrence et al., 2013) to identify five significantly mutated genes (SMGs): *TP53*, *CTNNB1*, *MEN1*, *PRKAR1A* and *RPL22*. The mutation frequencies ranged from 3.3% to 17.8% of the cohort (Figure 1A and Figure S1E). Mutations of *TP53* and *CTNNB1* in ACC are well recognized (Tissier et al., 2005). As expected, missense mutations in *CTNNB1* were confined to exon 3 (Figure 1A). Six (7%) tumors harbored inactivating mutations in *MEN1*, consistent with prior studies implicating *MEN1* in ACC (Assie et al., 2014). While our cohort is the largest to be sequenced to date, a much larger number of samples is needed to identify all candidate cancer genes (Lawrence et al., 2014). To overcome the limitation of sample size, we compared the mutated genes with the Cancer Gene Census (Futreal et al., 2004). This approach identified two cancer genes mutated in more than 5% of the cohort, *NF1* and *MLL4*, in addition to those nominated by MutSigCV.

In our cohort, 7 (8%) cases harbored inactivating mutations in the protein kinase cAMPdependent regulatory type I alpha gene (*PRKAR1A*) (Figure 1A). A homozygous deletion was found in three additional cases. While inactivating germline PRKAR1A mutations cause Carney complex and benign primary pigmented nodular adrenocortical disease (PPNAD) (Kirschner et al., 2000), malignant transformation has been reported in the adrenals of patients with this rare condition (Anselmo et al., 2012), and sporadic loss-of-function mutations in PRKAR1A have been found in adrenocortical adenomas and rare carcinomas (Bertherat et al., 2003). Interestingly, DNA sequencing of sporadic adrenocortical adenomas recently revealed a recurrent activating L206R mutation in the catalytic subunit of the cAMP-dependent protein kinase A (PKA) (PRKACA) (Beuschlein et al., 2014; Goh et al., 2014). This mutation results in constitutive PKA activity by disrupting the interaction between PRKACA and the regulatory subunits of PKA including PRKAR1A (Calebiro et al., 2014; Goh et al., 2014). While we found no *PRKACA* mutations in our cohort, we observed decreased PRKAR1A expression and increased MEK and BRAF protein expression (Figure S1F and S1G) in mutant cases, suggesting a potential role for inhibition of the RAF-MEK-ERK cascade in treatment of some ACCs.

Using two complementary analytical methods to detect fusion transcripts in mRNA-seq data (McPherson et al., 2011; Torres-Garcia et al., 2014), we identified 156 singleton but no recurrent gene fusion events in 48 of 78 (62%) tumors (range: 1–16) (Figure S1H and Table S1). Gene fusions occur at much lower frequencies than mutations and copy number variants (Yoshihara et al., 2015), and a larger cohort is needed to determine the frequency of fusions reported here. However, we did identify private in–frame fusions involving known cancer genes (Figure 1B). A highly expressed *EXOSC10-MTOR* fusion retained the mTOR catalytic domain and resulted in elevated levels of total and phosphorylated mTOR protein in this tumor (Figure S1I). The fusion point of a *MLL-ATP5L* fusion fell within the *MLL* breakpoint cluster region associated with acute myeloid leukemia (Krivtsov and Armstrong, 2007). Both fusion cases lacked mutations in SMGs. A fusion involving the gene *BRE*, reported to promote tumor cell growth and adrenal neoplasia (Chan et al., 2005; Miao et al., 2001), was identified in one tumor. While more data are required to ascertain their roles in ACC, these private fusions may represent functional transcripts.

Whole genome doubling is a common event in ACC

Lu, 2009).

We assessed somatic copy number alterations (SCNAs) and loss of heterozygosity in 89 tumors. Using GISTIC2 (Mermel et al., 2011), we identified recurrent focal amplifications of *TERT* (5p15.33), *TERF2* (16q22.1), *CDK4* (12q14.1) and *CCNE1* (19q12) and deletions of *RB1* (13q14.2), *CDKN2A* (9p21.2) and *ZNRF3* (22q12.1) (q 0.01; Figure 1C and Table S1). A focal deletion peak around 4q34.3–4q35.1 centered on a long noncoding RNA *LINC00290*, which has been reported as a deletion target in pediatric ACCs and other cancers (Letouze et al., 2012; Zack et al., 2013). *ZNRF3* homozygous deletions appeared in 16% (n=14) of tumors assayed; by including non-silent mutations, 19.3% of ACCs harbored alterations in this gene. *TERT* and *TERF2*, two telomere maintenance related genes (Blasco, 2005), were focally amplified in 15% and 7% of cases, respectively. *TERT* promoter hotspot mutations have been recently discovered in human cancers (Huang et al., 2013). We resequenced the *TERT* promoter region of all 91 tumors. In agreement with a recent study (Liu et al., 2014), we identified four cases with the C228T mutation, but no C250T mutations.

Arm-level copy number changes were frequent in ACC (Figure 2A and S2A). Clustering of 89 tumors based on their arm-level alterations produced three groups with striking differences: chromosomal (n=54; 61%); noisy (n=27; 30%); and quiet (n=8; 9%) (Figure S2A). The chromosomal group showed the highest frequency of whole chromosome arm gains and losses. The noisy group was characterized by a significantly higher number of chromosomal breaks as well as frequent loss of 1p with 1q intact. Tumors in the quiet group had few large copy number alterations. Kaplan-Meier analysis demonstrated a significant

decrease in survival in the noisy group relative to the chromosomal and quiet subtypes, suggesting that this copy number phenotype is characteristic of aggressive disease (Figure S2A). To validate these subtypes, we analyzed an independent data set of ACC tumors (n=119) profiled on different versions of the Illumina BeadArray platform (Assie et al., 2014). Based on the chromosome 1p/1q pattern, the noisy group and its survival association were recovered in this independent cohort (Figure S2B), suggesting that the group distinctions are robust. We did not observe copy number quiet ACC in the independent cohort.

We next determined tumor purity, ploidy and whole genome doubling (WGD) by integrating allelic copy number profiles and DNA mutation data using the previously validated ABSOLUTE algorithm (Carter et al., 2012). ACC samples were pure relative to other tumor types (tumor purity 0.82±0.15; Figure 2B). Consistently fractions of tumor infiltrated stromal cells estimated from gene expression signatures (Yoshihara et al., 2013) were low relative to a panel of 14 other cancer types (Figure S2C). Immune scores were lower in cortisol-secreting ACCs (p value=0.015), consistent with the suppression of T cell activity by glucocorticoids (Palacios and Sugawara, 1982). We found that hypodiploid karyotypes (ploidy 1.6) occurred more frequently in ACC than in eleven other tumor types (31% vs. 1%; Figure 2B), a frequency that was only matched by chromophobe renal cell carcinoma. WGD occurred in 68% of the noisy subtype, 51% of the chromosomal subtype, and none of the diploid quiet subtype cases. We inferred the temporal order of somatic mutations for genes implicated in ACC based on variant allele fractions (VAF) and genotype (Figure 2C and S2D). Mutations in TP53 (n=7), MEN1 (n=3), RPL22 (n=2), and ZNRF3 (n=2) were predicted to have occurred prior to WGD. Only 4/9 CTNNB1 mutations were predicted to be pre-doubling, while three were post-doubling and the remaining two showed further reduced VAFs suggestive of subclonality. PRKAR1A mutations split evenly as having occurred before WGD, after WGD or being subclonal.

We observed that loss of heterozygosity (LOH) patterns were nearly identical between hypodiploid and hyperdiploid cases in the chromosomal group (rho=0.96, p value< 10^{-5}) (Figure S3A and S3B). We hypothesized that the undoubled chromosomal ACCs were precursors to the chromosomal ACC that underwent whole genome doubling. This model is corroborated by the difference in outcome and absolute copy number (Figure 3A, S3C and S3D). We did not find a survival difference between non-WGD and WGD noisy samples suggesting that additional factors may contribute to disease course or that we were underpowered to detect a statistically significant signal. Mutation density further corroborated the WGD hypothesis as the higher mutation frequency in WGD cases was eliminated after normalization by ploidy (Figure S3E).

We employed gene set enrichment analysis using gene expression data to uncover differences between WGD and non-WGD tumors. We identified significantly enhanced pathways in WGD tumors including telomere regulation, cell cycle regulation and DNA replication repair (Figure S3F and S3G). The identification of cell cycle regulation was verified by an independent algorithm, Evaluation of Dependency DifferentialitY (Jung and Kim, 2014), which detected enrichment of the PARKIN pathway (Figure S3F) including *PARK2*, a recently reported master regulator of G1/S cyclins (Gong et al., 2014).

Amongst the telomere regulation pathway, TERT expression was significantly higher in the WGD group (FDR=0.05) (Figure 3B). Given the role of telomerase in maintaining telomere length (Blasco, 2005), we used spill over sequence generated by exome sequencing to infer telomere length of tumors and normal samples (Ding et al., 2014). Most tumors (73%) exhibited shorter telomeres than their matched normal samples (Figure 3C). WGD cases harbored shorter telomeres than non-WGD cases (Figure S3H). The association between WGD and *TERT* expression may suggest that increased *TERT* was required as a compensatory mechanism for telomere maintenance. Alternatively, TERT may have been non-functional with eroding telomeres as a consequence. This confirms previous observations that the majority of ACCs show telomerase activity, particularly in those with relatively short telomeres, while only a minority uses alternative telomere lengthening (ALT) as a telomere maintenance mechanism (Else et al., 2008). Telomere crisis is thought to drive tetraploidization (Davoli and de Lange, 2012) and may be directly associated with WGD events in these tumors. Similar to pediatric ACC and other cancers (Heaphy et al., 2011; Pinto et al., 2015), mutations in ATRX and DAXX(n=7) were associated with longer telomeres (p value= 5.4×10^{-5} , Fisher's exact test; Figure 3C). We did not find significant associations between telomere length and TERT promoter mutations or TERT amplification. Most cancers express *TERT* in the context of decreased telomere length (Maser and DePinho, 2002), suggesting that increased *TERT* maintains telomere homeostasis to a level that is sufficient for cancer cells to survive crisis. TERF2 was amplified in 7% of the cohort but similarly did not correlate with telomere length. TERF2 serves as an anchor protein of the shelterin complex which binds to telomeric DNA, but its direct involvement in telomere elongation has not been determined (Blasco, 2005). It has been documented that TERF2 might have telomere-independent functions (Stewart and Weinberg, 2006), raising interesting hypotheses about its role in adrenal tumorigenesis.

Molecular classes of adrenocortical carcinoma are captured by DNA Methylation signatures

To derive a robust molecular classification, we used unsupervised clustering to analyze the genomic and transcriptomic datasets (Figure S4 and Figure S2A). This approach yielded four mRNA expression groups (Figure S4A-B), six microRNA expression groups (Figure S4C-K), three DNA methylation groups (Figure S4L), three copy number groups (Figure S2A) and three protein expression groups (Figure S4M–N). Except for microRNA based clustering, the individual cluster analyses all resulted in classifications with significant differences in outcome. These subtypes are summarized in Table S2 and characteristics of each molecular classification are described in detail in the Supplemental Information. We integrated the ACC subsets identified across the DNA copy number, DNA methylation, mRNA expression and miRNA expression platforms through a Cluster of Cluster (CoC) analysis (Figure 4). Combined, the molecular classifications converged into three CoC subtypes (Figure 4A and S4O). Transcriptome clustering in a non-overlapping ACC sample set has previously identified an aggressive C1A subtype and an indolent C1B subtype (de Reynies et al., 2009; Giordano et al., 2009). A comparison between CoC and C1A/C1B showed that the majority of CoC I were classified as C1B, while most CoC II and CoC III were predicted as C1A (Figure 4A). Pan-cancer pathway enrichment analyses showed significant up-regulation of genes in immune-mediated pathways in CoC I tumors and

mitotic pathways in CoC III tumors (Figure S4P). We compared the clinical outcome of the three CoC clusters. Disease progression rates of the three CoCs were 7%, 56%, and 96%, respectively. Survival analysis showed a dismal median event-free survival of eight months for CoC III (Figure 4B) while median event-free survival time was not reached in CoC I. CoC II was more heterogeneous in outcome with an event-free survival of 38 months. Stage III/IV tumors represented 25%, 47%, and 52% of CoC I, II, III, respectively, and stage I/II cases in CoC III showed MKI67 values in accordance with their grade classification (Figure 5A).

While the CoC analysis showed that molecular data can determine outcome with high significance, implementing four parallel profiling platforms poses a clinical challenge. The four expression subtypes and three methylation subtypes rendered discriminative representations of each CoC group (Figure 4A), offering plausible routes for clinical implementation. Given the potential of the methylation platform to provide accurate data on formalin fixed, paraffin embedded tumor samples (de Ruijter et al., 2015; Thirlwell et al., 2010), we derived a methylation signature consisting of 68 probes that were included in both Illumina Human Methylation 450K and 27K arrays and tested its performance. This signature robustly classified our cohort into three ACC survival groups with 92.4% accuracy (Table S2). Classification of 51 methylation profiles from an independent cohort (Assie et al., 2014) accurately validated the prognostic power of the methylation signature (Figure S4Q).

Integrated genomic landscape of ACC

We generated a genomic landscape of ACC by further integrating multimodal mutational and epigenetic data with the CoC three class solution (Figure 5A). In addition to DNA copy number and mutations, we assessed epigenetic silencing and germline mutations of 177 manually selected genes (Table S3). We found 9 germline mutations, including two *TP53*^{R337H} in two Brazilian patients, two *MSH6*, one *MSH2* and one *NF1*^{R1534X} mutations. The *MSH6* and *MSH2* mutations support recent observations that ACC is a Lynch syndrome-associated cancer (Raymond et al., 2013). The two *TP53*^{R337H} mutated patients were younger than the rest of the cohort (23 and 30; median age of cohort: 49). We also found a single *TINF2*^{S245Y} variant that is associated with dyskeratosis congenita. The likely benign variant *APC*^{E1317Q} was observed in two cases (Rozek et al., 2006).

Collectively the genes altered most frequently by somatic mutations, DNA copy number alterations and epigenetic silencing were *TP53* (21%), *ZNRF3* (19%), *CDKN2A* (15%), *CTNNB1* (16%), *TERT* (14%) and *PRKAR1A* (11%). The majority of gene alterations were either mutation or copy number change, except *CDKN2A*, which was targeted by both deletion and epigenetic silencing through promoter DNA methylation. Alterations of *ZNRF3*, *CTNNB1*, *APC* and *MEN1* resulted in modification of the Wnt/β-catenin pathway in 41% of cases (Figure 5B). Wnt pathway activity, as measured by expression of canonical Wnt target genes, was increased in ACC with Wnt-pathway alterations relative to Wnt wild-type samples (Figure S5A). Somatic alterations in *TP53*, *CDKN2A*, *RB1*, *CDK4* and *CCNE1* emphasize the importance of the p53 apoptosis/Rb1 cell cycle pathway, and were altered in 44.9% of the cases (Figure 5B). Finally, histone modification genes (*MLL*, *MLL2*,

and *MLL4*) and chromatin remodeling genes (*ATRX* and *DAXX*) were collectively altered in 22% of cases, suggesting a role for epigenetic deregulation in ACC tumorigenesis (Figure 5C). Unsupervised HotNet2 analysis of protein-protein interaction network identified four significant subnetworks containing at least four genes (Figure S5B, p value<0.0001) (Leiserson et al., 2015). In addition to p53, Rb-cell cycle and Wnt-alteration subnetworks, HotNet2 showed a PKA network affecting 16 samples in total.

Combining somatic mutations, copy number alterations, and epigenetic modification, we found at least one alteration of potential driver genes in 69% of tumors (Figure 5A). The majority of CoC I tumors did not harbor a driver alteration that we could detect. We also examined the expression of *IGF2* and an established proliferation marker *MKI67*. *MKI67* expression was lower in CoC I tumors, consistent with the indolent phenotype of this subtype. *IGF2* expression was unanimously high independent of ACC classification. Expression of *IGF2* was relatively high in 67 of 78 (86%) ACCs (cutoff log₂(RSEM)=14 determined by expression distribution), with no association to either promoter DNA methylation or genome rearrangement.

Based on existing clinical trials and FDA-approved drugs for cancers, we found 51 potentially actionable alterations, including both mutations and copy number alterations, in 22 ACCs using precision heuristics for interpreting the actionable landscape (PHIAL) (Van Allen et al., 2014), from cyclin-dependent kinases to DNA repair protein poly ADP-ribose polymerase (PARP) (Figure S5C and Table S3).

Pan-Cancer Analyses provide context to ACC

Pan-cancer analysis of genomic data has provided insights on a wide range of cancer related questions. In an attempt to understand the driving mechanisms of ACC, we grouped our cohort on the basis of recurrent cancer-driving alterations using OncoSign (Ciriello et al., 2013). Our results broadly recapitulated the C class (OSC1–3, copy number driven) and M class (OSC 4–5, mutation driven) that have been described as a general trend across cancers (Ciriello et al., 2013) (Figure S6A).

The accumulation of somatic mutations in cancer is caused by mutational processes that can be deconvoluted as mutational signatures (Alexandrov et al., 2013). To examine mutational processes in ACC, we extracted 6 mutational signatures from 85 ACCs (mutation number 10) and about 2,900 other cancers using non-negative matrix factorization (Figure S6B). We compared the six signatures with the 22 signatures from an independent study (Alexandrov et al., 2013). Signature 1 resembled the age and DNA mismatch repair deficiency signatures, all featuring C>T substitution in the CG context. Signature 2 resembled the smoking signature featuring C>A substitution in the CG context. Signature 5 resembled UV and APOBEC signatures featuring C>G, C>T, and C>T substitutions in the contexts of TC, CC and TC, respectively. We did not find an association between signatures and sample country of origin (p value=0.9, Fisher's exact test). The majority of ACCs exhibited signatures 1, 2 and 4 (Figure 6A). Signature 1 captured the majority of gastrointestinal cancers (stomach, esophageal, colorectal) but also four ACC with a relatively high mutation frequency, all of which harbored mutations in the DNA mismatch repair pathway (Figure 6B). One case with a germline *MSH6* mutation but relatively modest mutation density (0.51 mutations/Mb) also

clustered with this signature. The smoking associated adenocarcinoma and squamous cell lung cancers in signature 2 also featured four high mutation frequency ACCs. Smoking has been listed as an ACC risk factor (Hsing et al., 1996). A set of Human papillomavirus (HPV)-driven cancers, including cervical, bladder and head and neck cancers, clustered together but did not capture any ACC. We analyzed RNA and exome sequencing data for microbial sequence reads (Chu et al., 2014) and identified human herpes virus sequences in nine exomes but not HPV sequence reads. De novo assembly of herpes virus sequences confirmed their presence but returned no evidence for viral genomic integration (Figure S6C).

Molecular correlates of ACC pathology and adrenocortical differentiation

We evaluated adrenocortical differentiation by using 25 genes with very high expression levels in the adult adrenal cortex and that are of importance for adrenal function, including steroidogenic enzymes, cholesterol transporters and their transcriptional regulator, SF1 (NR5A1) (Figure S7A and Table S4). We derived a single metric, termed Adrenocortical Differentiation Score (ADS), to measure adrenocortical differentiation (Figure 7). Sorting the tumors according to their ADS values we observed that functional tumors showed higher ADS values, which did not correlate with Weiss histopathology score (p value=0.41, ANOVA). TP53 mutations were present across the spectrum of ADS values (p value=0.56, Fisher's exact test), whereas Wnt-related mutations appeared to be enriched among tumors with higher ADS values (p value=0.0091, Fisher's exact test). Not surprisingly, the two sarcomatoid tumors had the lowest ADS values with very low expression of NR5A1 and steroidogenic enzymes (Figure S7B). One such patient had elevated serum cortisol levels, suggesting a mixed histology in which a differentiated component expressed steroidogenic enzymes and an undifferentiated component did not. The other tumor was histologically mixed with separate components of usual and sarcomatoid ACC. We suggest that these sarcomatoid, NR5A1-negative tumors do indeed represent dedifferentiated ACCs, rather than retroperitoneal sarcomas.

DISCUSSION

As part of TCGA, we present an integrated molecular characterization of a large cohort of ACC. The tumors were derived from four continents and thus represent a near-global sampling of this disease. The data presented here represents a resource for future investigations to facilitate ACC research across myriad avenues, including via pan-cancer analysis.

Our study builds upon prior efforts on adult ACC from European (Assie et al., 2014) and North American patients (Juhlin et al., 2015), as well as pediatric ACC patients from North America and Brazil (Pinto et al., 2015). Using high quality multidimensional genomic data, we confirm many alterations as essential for ACC development and progression but also expand the somatic genetic landscape of ACC to nearly double the known ACC driver genes. The high frequency of *PRKAR1A* mutations expands the role of PKA signaling in ACC and is consistent with *PRKACA* somatic mutations being the founder lesion of benign adrenal tumor associated with endocrinopathies such as Cushing syndrome (Beuschlein et al., 2014). By integrating data from multiple platforms we demonstrated the critical role of WGD. While WGD is common across cancer (Zack et al., 2013), an association between WGD and patient outcome has only been reported in ovarian cancer (Carter et al., 2012). The exceptional high level of tumor purity and the clear evidence that WGD is a marker of tumor progression make ACC a model disease for better understanding of the mechanisms that result in doubling and the molecular context needed to sustain it.

Weiss and colleagues (Weiss et al., 1989) first proposed that ACC consists of two pathologic classes that have different mitotic rates and distinct clinical outcomes. This proliferationbased two-grade (low and high) system has been confirmed by several transcriptome studies (de Reynies et al., 2009; Giordano et al., 2009) and has been clinically implemented by some centers (Giordano, 2011). We confirmed the two-grade classification with mRNA-seq data and importantly, used multidimensional data to extend the molecular classification to three classes that have markedly distinct biological properties and significantly different patient outcomes. Clinical implementation of this three-class grading system using DNA methylation profiles could facilitate improved patient care, although additional translational efforts are needed for its implementation. Despite these uncertainties, our results represent an extensive ensemble of molecular ACC subtypes and thus are likely to be useful in directing future development of clinically applicable classifiers. Moreover, our results illustrate how molecular data, combined with traditional clinicopathologic assessment, might inform therapeutic decisions and lead to advances in patient outcomes. The diversity of genomic alterations, especially the significant copy number changes seen in the majority of ACC, suggests that combined inhibition of disease pathways, however challenging, likely holds the key to successful targeted therapy for ACC.

EXPERIMENTAL PROCEDURES

Tumor and normal samples were obtained from patients after informed consent and with approval from local Institutional Review Boards (RIB). Details on all contributing Centers and their IRB approval can be found in the Supplemental Information. DNA, RNA and protein were purified and distributed throughout the TCGA network. In total, 91 primary tumors with associated clinicopathologic data were assayed on at least one molecular profiling platform. Platforms included exome sequencing, mRNA sequencing, miRNA sequencing, SNP arrays, DNA methylation arrays, and reverse phase protein arrays. Mutation calling was performed by 5 independent callers, and a voting mechanism was used to generate the final mutation set. MutSigCV (version 1.4) was used to determine significantly mutated genes (Lawrence et al., 2013). GISTIC2.0 was used to identify recurrent deletion and amplification peaks(Mermel et al., 2011). Consensus clustering was used to derive miRNA, mRNA, methylation and protein subtypes. Tumor purity, ploidy and whole genome doubling were determined by ABSOLUTE (Carter et al., 2012). The data and analysis results can be explored through TCGA data portal (https://tcga-data.nci.nih.gov/ tcga/tcgaCancerDetails.jsp?diseaseType=ACC), the Broad Institute GDAC FireBrowse portal (http://firebrowse.org/?cohort=ACC), Memorial Sloan Kettering Cancer Center cBioPortal (http://www.cbioportal.org/public-portal/study.do?cancer_study_id=acc_tcga), Regulome Explorer (http://explorer.cancerregulome.org/all pairs/?dataset=TCGA ACC).

See also the Supplemental Experimental Procedures and the ACC publication page (https://tcga-data.nci.nih.gov/docs/publications/acc_2016/).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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SIGNIFICANCE

Adrenocortical carcinoma is a rare endocrine cancer with limited therapeutic options and overall poor outcome. We comprehensively analyzed 91 ACC specimens from four continents using state of the art genomic technologies and computational methods. In addition to identification of ACC driver genes, pathways and refined subtypes, our analysis revealed whole genome doubling (WGD) as a milestone in disease progression. Our findings suggest that ACC can serve as a model for how WGD influences disease progression. Our dataset is standardized with other TCGA studies, fully available at the TCGA Data Portal, and thus will serve as a valuable research resource.

Highlights

• Standardized molecular data from 91 cases of adrenocortical carcinoma

- Driver genes including TP53,ZNFR3,CTNNB1, PRKAR1A,CCNE1 and TERF2
- Whole genome doubling event is a marker for ACC progression
- Three prognostic molecular subtypes captured by a DNA methylation signature

Zheng et al.



Figure 1. Mutational driver genes in ACC

(A) Protein domain structure of the five significantly mutated genes with somatic mutations aligned. Functional domains and mutation types are indicated in different colors and shapes as shown in the legend. Arm, Armadillo domain; TAD, transcription-activation domain; DBD, DNA binding domain; TMD, tetramerisation domain; RIIa, regulatory subunit of type II PKA R-subunit. (B) Sporadic gene fusions that involve cancer genes. All exons are represented, with red and blue indicating high and low expression respectively. Lines linking two exons indicate the fusion positions. Protein domains are related to the exons below the

gene diagram. HRDC, helicase and RNase D C-terminal; PI3Kc, phosphoinositide 3-kinase, catalytic domain; ATH, AT-hook motif; PHD, plant homology domain; SET, Su(var)3–9, enhancer-of-zeste, trithorax; PP2Ac, protein phosphatase 2A homologues, catalytic domain. (C) Focal recurrent amplifications and deletions in ACCs with the number of genes spanned by the peak in parentheses. Red and blue indicate amplification and deletion, respectively. The x-axis at the bottom and top of the figure represents significance of amplification/ deletion per q-value. See also Figure S1 and Table S1.

Zheng et al.



Figure 2. Landscape of DNA copy number alteration in ACC

(A) Three major copy number patterns in ACC. Unsupervised clustering divided the cohort (n=89) into quiet, chromosomal and noisy subtypes. (B) Pan-cancer purity and ploidy including ACC. Sample sizes are indicated on top. Average tumor purity is plotted as a grey line for each cancer type. The percentages of whole genome doubling and hypodiploidy (ploidy 1.6) are listed in red and blue, respectively. LUAD - lung adenocarcinoma; LUSC – lung squamous; HNSC – head and neck; KIRC – clear cell renal cell; BRCA – breast; BLCA – bladder; CRC – colorectal; THCA – thyroid papillary; UCEC – endometrial; GBM – glioblastoma; OV – ovarian; KICH – kidney chromophobe. (C) Purity adjusted variant allele fraction in genome doubled and undoubled tumors. Only tumors with high purity (0.8) and mutation density less than 5 were included. Cutoffs labeled in the figure are recognized as turning points in the density distributions of variant allele fractions. See also Figure S2.

Zheng et al.

Page 23



Figure 3. Comparison of WGD and non-WGD ACCs

(A) Event free survival of copy number subtypes and whole genome doubling groups. P value represents the statistical significance of event free survival differences between the five groups. (B) *TERT* expression in genome undoubled and doubled tumors. Boxplot shows median and interquartile range of *TERT* expressions, with whiskers extending to extreme values within 1.5 interquartile ranges from the upper and lower quartiles. Each dot corresponds to a tumor. (C) Telomere length was estimated using off-target exome sequencing data corrected for tumor purity and ploidy. Top panel shows *TERT* expression

with "x" representing missing values. *TERT* promoter C228T mutation (black), amplification (red) and *TERF2* amplification (red) are noted in the second panel. The bottom panel represents *ATRX* and *DAXX* mutations in light blue and dark red, respectively. See also Figure S3.



Figure 4. Cluster of clusters

(A) Cluster of clusters (CoCs) from four platforms (DNA copy number, black; mRNA expression, red; DNA methylation, blue; miRNA expression, purple) divided the cohort into 3 groups. Presence or absence of membership for each sample is represented by black or light blue ticks, respectively. Sample parameters are aligned on top of the heatmap. White tick indicates data not available. (B) Event free survival of the 3 CoC groups. Pairwise log-rank test p values are shown. See also Figure S4 and Table S2.

Zheng et al.



Figure 5. Genomic landscape of ACC

(A) The ensemble of mutations, copy number alterations, methylations, subtypes and clinicopathological parameters. (B) Aggregated alterations of the p53/Rb pathway and the Wnt pathways. Activating and deactivating alterations are indicated in red and blue, respectively. (C) Mutations in epigenetic regulatory genes. See also Figure S5 and Table S3.

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Figure 6. Pan-cancer mutational signature analysis

(A) Distribution of the mutational signatures extracted from pan-cancer analysis in the ACC cohort. Signature 2 is enriched in the cluster of clusters groups 1 and 2. (B) Circular plot of the mutational signatures in ACC and approximately 2,900 tumor samples from other cancer types. The distance to the center represents coding mutation density. Three small illustrations highlight ACC, lung squamous cell carcinoma and colorectal cancer to demonstrate their similarities in the featured directions. See also Figure S6.

Zheng et al.



Figure 7. Distribution of Adrenal Differentiation Score (ADS)

The 25-gene signature is shown in the expression heatmap. Adrenal cortex differentiation markers are listed on the left. Two sarcomatoid cases are indicated in red. See also Figure S7 and Table S4.