VHL deficiency drives enhancer activation of oncogenes in clear cell renal cell carcinoma

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Abstract
Protein-coding mutations in clear cell renal cell carcinoma (ccRCC) have been extensively characterized, frequently involving inactivation of the von Hippel Lindau (VHL) tumor suppressor. Roles for non-coding cis-regulatory aberrations in ccRCC tumorigenesis, however, remain unclear. Analyzing 10 primary tumor/normal pairs and 9 cell lines across 79 chromatin profiles, we observed pervasive enhancer malfunction in ccRCC, with cognate enhancer-target genes associated with tissue-specific aspects of malignancy. Super-enhancer profiling identified ZNF395 as a ccRCC-specific and VHL-regulated master regulator, whose depletion causes near-complete tumor elimination in vitro and in vivo. VHL loss predominantly drives enhancer/super-enhancer deregulation more so than promoters, with acquisition of active enhancer marks (H3K27ac, H3K4me1) near ccRCC hallmark genes. Mechanistically, VHL loss stabilizes HIF2α-HIF1β heterodimer binding at enhancers, subsequently recruiting histone acetyltransferase P300 without overtly affecting pre-existing promoter-enhancer interactions. Subtype-specific driver mutations such as VHL may thus propagate unique pathogenic dependencies in ccRCC by modulating epigenomic landscapes and cancer gene expression.

Significance
Comprehensive epigenomic profiling of ccRCC establishes a compendium of somatically altered cis-regulatory elements, uncovering new potential targets including ZNF395, a ccRCC master regulator. Loss of VHL, a ccRCC signature event, causes pervasive enhancer malfunction, with binding of enhancer-centric HIF2α and recruitment of histone acetyltransferase P300 at pre-existing lineage-specific promoter-enhancer complexes.
Introduction

Clear cell renal cell carcinoma (ccRCC) is the most common subtype of kidney cancer with 338,000 new cases in 2012 worldwide (1). With most ccRCCs being radio-chemo-resistant, metastatic ccRCC patients exhibit dismal 8% five-year overall survival (2). While targeted therapies inhibiting angiogenesis and mTOR pathways can lead to initial tumor control, most patients develop resistance in less than a year (3,4). A better understanding of ccRCC molecular dependencies and vulnerabilities is thus needed to develop new therapeutic strategies for patients who fail standard of care treatment.

Loss of the von Hippel-Lindau (VHL) tumor suppressor is a defining feature of ccRCC (5,6). When partnered with additional tumor suppressors (Pbrm1, Bap1, Trp53, Rb1, and/or Cdkn2a) and/or oncogenes (Myc), Vhl loss drives spontaneous ccRCC formation in mouse models (7-11). VHL encodes an E3 ubiquitin ligase (12,13) that targets hypoxia-inducible factor (HIF)-1α (HIF1A) and HIF2α (EPAS1), for degradation (14,15). VHL loss in ccRCC results in constitutive activation of HIF1/2α, and subsequent transactivation (16,17) of downstream genes regulating angiogenesis, glycolysis (18), lipogenesis (19,20), cell cycle (21) and anti-apoptosis (22).

Most reports studying VHL/HIF transcriptional activation have focused on HIF-bound promoters (23-28). However, recent evidence suggests an emerging role for distal enhancer elements in VHL/HIF transcriptional control (29,30). For example, HIF2α-bound distal enhancers activate the proto-oncogenes MYC (31) and CCND1 (21), and coincide with ccRCC genetic risk alleles. Nevertheless, such studies focused on individual enhancers, and the majority of distal elements in ccRCC remain largely unexplored.

Delineating the global ccRCC cis-regulatory landscape may also identify novel master regulators involved in tissue-specific disease processes. Compared to promoters that are largely cell-type invariant, distal enhancers integrate multiple lineage- and context-dependent
signals, catering to the specialized needs of diverse cell types and diseases (32,33). In cancer, such master regulators are frequently located near “super-enhancers” or “stretch-enhancers” marked by long stretches of H3K27ac signals (34,35). For example, subtype-specific genomic alterations such as EGFRvIII in glioblastoma (36) and EWS-FLI in Ewing’s sarcoma (37) induce de novo enhancers, causing reactivation of developmental master regulators required for self-renewal and lineage specification (36). While VHL inactivation has been shown to modulate protein levels of different histone modifiers (e.g. KDM5C/JARID1C (38), HDAC1 (39), JMJD1A (40), JMJD2B (40) and JMJD2C (41)), the impact of these protein alterations at specific epigenomic loci remains unclear. Moreover, previous studies profiling histone modifications in ccRCCs have also been limited by small sample sizes (2 cases, (42)), reliance on in vitro systems, and the lack of long-range interactome data and functional enhancer testing to accurately assign cognate enhancer targets.

In this study, we establish the most comprehensive collection of ccRCC histone profiles to date, annotating the precise genomic locations of altered promoters, enhancers and super-enhancers in ccRCC. Using isogenic cell lines with or without wild-type VHL, we further demonstrate that besides its well-defined role in oxygen sensing, VHL also safeguards the chromatin landscape; its loss induces tumor-specific enhancer gains around ccRCC hallmark genes such as angiogenic and metabolic targets through the stabilization of HIF2α/HIF1β (ARNT) heterodimers and recruitment of P300 histone acetyltransferase (EP300). One important target of epigenetic activation is ZNF395, a master regulator of ccRCC tumorigenesis. Taken collectively, our results reveal an epigenetic framework by which the major ccRCC-specific driver mutation, VHL, induces de novo enhancers, contributing to oncogenic transcription.
Results

Cis-regulatory landscapes in ccRCC tumors are aberrant

To explore whether ccRCC tumors display alterations in their cis-regulatory landscapes in vivo, we generated histone ChIP-seq profiles (3 marks: H3K27ac, H3K4me3, H3K4me1) in 10 primary tumor/normal pairs, 5 patient-matched tumor-derived cell lines, 2 commercially available ccRCC lines (786-O, A-498) and 2 normal kidney cell lines (HK-2, PCS-400) (Refer to Supplementary Table S1 for patient clinical information). Of the original 87 samples, 79 samples passed pre-sequencing quality control filters and were subjected to ChIP-seq processing and downstream analysis. In total, we generated 2,363,904,778 uniquely mapped reads (Refer to Supplementary Table S2 for sequencing statistics). On average, 89% of H3K27ac peaks, 98% of H3K4me3 peaks and 76% of H3K4me1 peaks obtained in our normal kidney tissues overlapped with peaks from adult kidney tissues in the Epigenomics Roadmap dataset (Supplementary Figure S1A). Among the 10 primary ccRCCs, 9 harbored VHL mutations, detected by targeted sequencing and confirmed by Sanger sequencing (Supplementary Table S3). Cell lines 786-O and A-498 also harbor VHL truncating mutations (Supplementary Table S3). The VHL mutations co-occurred with somatic mutations of other chromatin modifiers commonly found in ccRCC, including PBRM1 (7/10), SETD2 (1/10), KDM5A (1/10), KDM5C (1/10), ARID1A (1/10) and KMT2C (1/10).

Specific histone modifications can distinguish different categories of functional regulatory elements - H3K4me3 is generally associated with promoters, H3K4me1 with enhancers and H3K27ac with active elements (33,43). Integrating signals from 3 histone marks and GENCODE v19 annotated transcription start sites (TSS), we defined active promoters as H3K27ac⁺/H3K4me3⁺/±2.0 kb TSS regions, and distal enhancers as H3K27ac⁺/H3K4me1⁺ regions not overlapping with promoters. Focusing on epigenomic events specific to somatic
cancer cells, we derived cell lines from 5 primary tumors and, combined with the commercial lines, excluded peaks not found in any of the cell lines to reduce confounding effects from stromal cells. On average, we observed 80% overlap of ChIP-seq peaks between primary tumors and matched lines (Supplementary Figure S1B). Using these criteria, we identified 17,497 putative promoters and 66,448 putative enhancers (Figure 1A), numbers comparable to previous studies in other tumor types (43-45). The numbers of defined promoters and enhancers reached saturation after 4 and 16 samples respectively, suggesting that a sample size of 20 (10 tumor/normal pairs) is sufficiently powered to discover the majority of cis-regulatory elements in ccRCC (Supplementary Figure S1C, D). Principal components analysis (PCA) using the first 2 components of global H3K27ac intensities at promoters or enhancers (representing 83% and 64% of total variance respectively Supplementary Figure S1E, F) successfully separated normal and tumor samples, indicating that genome-wide pervasive alterations in cis-regulatory elements are a salient feature of ccRCC (Figure 1B).

We performed differential analysis to identify altered promoters and enhancers. To define gained or lost regions, we applied a fold difference of H3K27ac RPKM $\geq$ 2, an absolute difference $\geq$ 0.5, and for greater stringency no alterations in the reverse direction in the remaining tumor/normal pairs (see Methods and Supplementary Figure S1G for distribution of altered elements by number of patients). At the threshold of $\geq$5/10 patients, 80% of the altered regions achieved statistical significance ($q$-value $<$ 0.1, paired t-test, with Benjamini-Hochberg correction) (Supplementary Figure S1H) and at this same threshold, the increase in the fraction of samples meeting statistical significance reached a saddle point (Supplementary Figure S1I). Applying these criteria, we obtained a high-confidence and comprehensive set of 4,719 gained promoters, 592 lost promoters, 4,906 gained enhancers, and 5,654 lost enhancers (Figure 1A, C, Supplementary Table S4). Representative regions are presented in Supplementary Figure S2.
Supporting the reliability of this data, gained promoters and enhancers exhibited increased chromatin accessibility measured by higher FAIRE-Seq signals (46) in tumor tissues than normal tissues respectively ($p$-value < 0.0001), and also decreased DNA methylation based on TCGA data (47), consistent with reciprocal relationships between active regulatory regions and DNA methylation (Figure 1D). Interestingly, we also noted elevated expression of long non-coding RNAs (48) adjacent to gained promoters and enhancers in tumor tissues compared to normal tissues ($p$-value < 0.0001 respectively). Lastly, we confirmed that many of our cis-regulatory elements involved regions previously implicated in ccRCC – for example, we observed gains of H3K27ac signals and enrichment of H3K4me1 at a distal enhancer of $CCND1$ overlapping with an RCC susceptibility locus (rs7105934 (49) (21)) (Figure 1E). Our ability to re-discover this important enhancer in our unbiased profiling supports our data reliability.

**Tumor-specific enhancers are associated with hallmarks of ccRCC**

To identify genes modulated by the tumor-specific regulatory elements, we assigned enhancers using three approaches. The first approach utilized pre-defined linear proximity rules involving a set of highly confident genes (GREAT algorithm) (50) (Supplementary Table S5). MSigDB pathway analysis using GREAT-assigned genes revealed that gained enhancers exhibit a highly significant RCC-specific signature compared to gained promoters (enhancer $q$-value = $3.2 \times 10^{-26}$, promoter $q$-value = $1.5 \times 10^{-1}$, binomial FDR) (Figure 2A). While gained promoters were involved in general cancer processes (e.g. cell cycle, transcription and RNA metabolism, see Supplementary Table S6 for a complete list of promoter pathways), gained enhancers were enriched in disease-specific features of ccRCC including HIF1$\alpha$ network activity, pro-angiogenic pathways (platelet activation, PDGFR$\beta$ signaling), and SLC-mediated transmembrane transport (Figure 2A, Supplementary Table S7 for a complete list of enhancer pathways). Notably,
HIF1α network activity consistently emerged as one of the top 5 pathways, even with perturbations in the patient thresholds used to define gained enhancers (≥ 3-8 patients) (Supplementary Table S8).

Individual genes associated with gained enhancers included well-known hypoxic targets (VEGFA (Figure 2B), CXCR4) and metabolic genes involved in glycolysis, glutamine intake and lipid storage (GLUT1/SLC2A1 (Figure 2C), HK2, PFKFB3, PLIN2 (Supplementary Figure S3A) and SLC38A1 (Supplementary Figure S3B) (51)). The presence of enhancers around metabolic enzymes and transporters is largely consistent with the metabolic contexture of ccRCC, which involves increased glycolysis and glutaminolysis (19,52-55). Indeed, gene ontology (GO) analysis of gained enhancers strongly reflected hallmark metabolic changes associated with ccRCC, including monocarboxylic acid transmembrane transporter activity (binomial FDR q-value = 1.6 x10^{-10}) (Supplementary Figure S3C).

We also used a second method of enhancer-gene assignment based on correlations between H3K27ac signals and expression of genes within the same topological associated domain (TAD) (34). Using a q-value <0.05 based on Spearman’s correlation, we assigned 2311 gained enhancers to 2186 protein-coding targets (Supplementary Table S9). Reassuringly, H3K27ac signals of many gained enhancers were highly correlated with gene expression of their putative target genes. For example, H3K27ac levels of a VEGFA enhancer exhibited high correlation with VEGFA gene expression (r = 0.83, Spearman’s correlation), while H3K27ac signals of a SLC2A1 enhancer were highly correlated with SLC2A1 gene expression (r = 0.72, Spearman’s correlation) (Figure 2B, Supplementary Figure S3D). Similar to the GREAT approach, the TAD correlation approach also highlighted hypoxia (Krieg_Hypoxia_not_via_KDM3A, FDR q-value = 7 x 10^{-120}) and metabolism (Chen_Metabolic_Syndrome_Network, FDR q-value = 2 x 10^{-91}) as highly enriched pathways (Supplementary Table S10).
Thirdly, to independently validate the GREAT and TAD approaches in the specific context of ccRCC, we experimentally explored the interactome of ccRCC tumor-specific enhancers by performing Capture-C assays (56). Compared to other chromatin capture techniques, Capture-C offers both high-resolution (down to single Kb resolution) and high-throughput interrogation of user-defined regions (a usual working range of 10-500 regions). We designed probes against a subset of 56 gained enhancers and examined their interactions with protein-coding genes in 786-O cells. Each gene-enhancer pair revealed by Capture C was further filtered by correlations between gene expression and H3K27ac levels (q-value <0.05). The 56 gained enhancers were paired with 36 protein-coding genes (Supplementary Table S11) – of these, 58% were predicted by GREAT, and 80% by gene correlations within TADs. The median distance of interactions detected by Capture-C was 16 kb, and 83% of the interactions fell within a 100 kb window (Supplementary Figure S3E). As a visual example, Capture-C confirmed interactions between VEGFA enhancer with the VEGFA TSS, spanning a distance of ~100 kb (Figure 2B), and the interactions between the SLC2A1 enhancer and its promoter (Figure 2C). Taken collectively, these findings highlight the disease-specific nature of enhancer elements (33) and an important role for enhancer malfunction in modulating ccRCC pathology.

**Tumor super-enhancers identify ZNF395 as a master regulator of ccRCC tumorigenesis**

The importance of enhancers in ccRCC led us to examine the landscape of “super-enhancers” or “stretch-enhancers” – dense clusters of enhancers located near master regulators of cell identity and disease (35,57) 15. Using ROSE (35), we identified 1,451 super-enhancers in the ccRCC cohort, of which 1,157 were gained in tumor and 294 were lost in tumors (Supplementary Table S12).
Putative targets of top gained super-enhancers validated well-known oncogenes including MYC/PVT1, VEGFA and HIF2A (Figure 3A, Supplementary Figure S4A, B). In addition, we found several less known genes including ERGIC1, ZNF395, SLC28A1 and SMPDL3A (Figure 3B). These genes were highly overexpressed in tumors compared to their matched normal tissues (Figure 3B). Furthermore, they were unique to ccRCC and were not overexpressed in papillary and chromophobe RCCs, two other distinct ccRCC subtypes (Figure 3B). For instance, ZNF395 exhibited a tumor-normal ratio of ~7 in ccRCC (p-value = 1x10^{-22}, paired t-test) but experienced little over-expression in papillary and chromophobe RCC with tumor-normal ratios of 1.2 and 1.3 respectively (p-value = 0.02 in papillary and p-value = 0.06 in chromophobe, paired t-test).

Conversely, genes associated with lost super-enhancers were recurrently suppressed in ccRCC and included EFHD1, EHF, MAL, GCOM1 and HOXB9 (Figure 3B). In contrast to the lineage-specific nature of tumor super-enhancers, genes associated with lost super-enhancers were common between ccRCC and papillary RCC, implying a more universal function of tumor suppressor genes. For example, EHF/ESE-2, a tumor suppressor previously found in prostate cancer (58,59), exhibited reduced expression across all three RCC subtypes (ccRCC tumor/normal = 0.05, p-value = 3 x 10^{-15}; papillary tumor/normal = 0.1, p-value 2x10^{-6}; chromophobe tumor/normal = 0.1, p-value = 2x10^{-6}).

Since current therapeutic targets in kidney cancer are limited to angiogenesis and mTOR pathways (3), we sought to examine these less understood genes uncovered by super-enhancer profiling. We chose ZNF395 and SMPDL3A for their differential tumor expression (6-7 tumor-normal ratio) (Figure 3B) and high abundance (average RPKM of ZNF395 ~112, average RPKM of SMPDL3A ~58). Even though ZNF395 was previously identified as a potential ccRCC biomarker (60), its functional role in ccRCC malignancy remains unexplored. SMPDL3A shares
31% amino acid identity with the acid sphingomyelinase \textit{SMPD1}, and is a target of master
regulator of cholesterol metabolism, Liver X Receptors (LXR) (61).

Quantitative PCR (\textbf{Figure 3C}) and immunoblotting (\textbf{Supplementary Figure S4C}) confirmed
that A-498 and 786-O ccRCC cells exhibited high expression of \textit{ZNF395} and \textit{SMPDL3A}
whereas normal kidney proximal tubule cells, PCS-400 and HK-2, exhibited low expression of
both genes. SiRNA mediated knockdown of \textit{SMPDL3A} had a cell line dependent effect on
colony formation, inhibiting the growth of A-498 cells but having no observable effect on 786-O
cells (\textbf{Figure 3D}). On the other hand, \textit{ZNF395} consistently inhibited colony formation in both
786-O and A-498 cells but had minimal effect on normal kidney cells (\textbf{Figure 3D}, \textbf{Supplementary Figure S4D}). Consistent with this phenotypic observation, the \textit{ZNF395} super-
enhancer was only active in ccRCC cells (786-O, A-498) but silent in normal kidney cells (HK-2,
PCS400; \textbf{Figure 3E}). Furthermore, amongst 12 types of cancers profiled by TCGA, \textit{ZNF395}
was exclusively overexpressed in ccRCC tumors, consistent with the proposed lineage- and
disease-specific nature of super-enhancers (\textbf{Figure 3F}).

No study to date has functionally tested the tumorigenic requirement of \textit{ZNF395} in ccRCC or
any other cancer type. We validated \textit{ZNF395}’s tumor-promoting effect using individual shRNA
clones (\textbf{Supplementary Figure S4E, F}). Two independent \textit{ZNF395} shRNA clones drastically
decreased \textit{in vitro} colony formation (\textbf{Figure 3G}) and cell viability (\textbf{Figure 3H}) in both A-498 and
786-O cells. \textit{ZNF395} knockdown also resulted in increased apoptosis measured by cleavage of
Caspase3/7 substrates (\textbf{Figure 3I}) and Annexin V staining (\textbf{Supplementary Figure S4G}). \textit{In vivo}, tumor formation studies in mouse xenograft models revealed marked tumor suppression
by \textit{ZNF395} depletion (\textbf{Figure 3J}). Knockdown of \textit{ZNF395} led to elimination of A-498 tumors up
to day 74, when tumors in the control group began to exceed the size limits imposed by
institutional animal protocols. Similarly, \textit{ZNF395} depletion significantly slowed \textit{in vivo} tumor
growth of 786-O cells (Figure 3J). Taken together, we showed the indispensable role ZNF395 plays in ccRCC tumorigenesis.

**VHL deficiency remolds ccRCC enhancer landscapes**

To explore the extent to which epigenetic changes observed in primary ccRCCs (Figure 1) are directly driven by VHL loss, we examined chromatin changes in isogenic cell lines with and without VHL restoration. Consistent with earlier functional studies of VHL (62-64), VHL restoration in 786-O, A-498 and 12364284 cells had negligible effects on proliferation, colony formation and apoptosis *in vitro*, but profoundly delayed tumor growth *in vivo* (Supplementary Figure S5A-D), suggesting the importance of VHL in modulating processes required for *in vivo* tumorigenesis, including tumor-stroma crosstalk, angiogenesis, cell-matrix interactions, or tumor metabolism.

Focusing on the same regions defined in the primary tumors (4,719 gained promoters, 4,906 gained enhancers and 1,157 gained super-enhancers; Figure 1A), we examined VHL-driven H3K27ac changes in 4 different cell lines (two commercial cell lines – 786-O and A-498 and two patient-derived cell lines – 12364284 and 40911432). Consistently across all 4 cell lines, VHL restoration induced more pronounced changes on enhancers and super-enhancers than on promoters (Figure 4A, Supplementary Figure S6A-C). For example, in 786-O cells, after VHL restoration 12% of enhancers (549 enhancers) were significantly depleted, compared to 6.5% of promoters (321 promoters) (Figure 4A). We confirmed that a greater fraction of enhancers were significantly altered by VHL restoration than promoters (p < 2.2 x 10^-16, proportions test), and an even higher proportion involved gained super-enhancers (p < 2.2 x 10^-16, proportions test).

Even though gained enhancers were expected to show only depletion after VHL restoration, changes in H3K27ac levels were bi-directional (Figure 4A). However, only gained enhancers
with H3K27ac depletion were uniquely active in VHL-mutated ccRCC cell lines (786-O, A-498, 12364284) compared to VHL-wild-type ccRCC cells (86049102L), normal kidney cell lines (PCS-400, HK-2 and HKC-8), and 31 other cell lines of various cancer types (Figure 4B). The lack of H3K27ac signals in normal kidney cell lines argues against tissue lineage as the dominant contributor to the high H3K27ac ChIP-seq signals seen in ccRCC cell lines. On the other hand, gained enhancers with H3K27ac enrichment after VHL restoration showed high activity across multiple cancer types, suggesting that these enhancers are not unique to ccRCC (Figure 4B).

Furthermore, only gained enhancers showing H3K27ac depletion after VHL restoration were significantly associated with a concomitant downregulation of gene expression of their putative targets in both 786-O and 12364284 cells, whereas enhancers gained in primary ccRCCs and further H3K27ac-enriched after VHL restoration did not lead to significant gene upregulation on a global level (Figure 4C, Supplementary Figure S6D). These results suggest that the former enhancers (H3K27ac depletion) are likely to represent ccRCC- and VHL-specific epigenomic alterations, while the latter enhancers (H3K27ac enrichment) are likely to represent signify generic, compensatory mechanisms in response to VHL restoration.

Combining data from multiple lines, a total of 1564 enhancers were depleted by VHL restoration in ≥1 cell line, representing almost a third (32%) of all gained enhancers identified in primary ccRCC tumors (Supplementary Table S13). The proportion of VHL-responsive enhancers increased with the level of patient recurrence – only 7.8% of non-recurrent gained enhancers (1/10 patients) showed VHL-mediated H3K27ac depletion while 18% of enhancers recurrently gained in 9/10 patients and 20% of enhancers gained in 10/10 patients showed H3K27ac depletion in 786-O cells (Supplementary Figure S7A, p-value = 0.0001, proportions test), consistent with the high prevalence of VHL mutations (9/10 patients) in our discovery set. Interestingly, unsupervised clustering using the 1564 VHL-responsive gained enhancers
segregated the single VHL wild-type tumor (ID 75416923) away from the remaining VHL mutant 9 tumors (Supplementary Figure S7B), with the VHL-wild-type tumor showing low H3K27ac signals at the ZNF395 super-enhancer comparable to its patient-matched normal (Supplementary Figure S7C). Collectively, pathway analysis of enhancers depleted in ≥2 cell lines highlighted direct p53 effectors, integrin-linked kinase signaling and HIF1α transcription factor networks as the top 5 pathways (Supplementary Table S14), covering genes such as EGFR (Figure 4D), CCND1 (Figure 4E), ITGB3 (Figure 4F), VEGFA (Figure 4G), SLC2A1 (Supplementary Figure S7D), and HK2 (Supplementary Figure S7E). These results support a major role for VHL loss in ccRCC enhancer malfunction, even in the presence of other driver mutations.

We also examined whether other histone marks were concomitantly altered with H3K27ac marks. We found a surprisingly high degree of correlation between H3K27ac and H3K4me1 in response to VHL restoration in both 786-O cells (r = 0.77, Pearson’s correlation, Supplementary Figure S7F) and 12364284 (r = 0.61, Pearson’s correlation, Supplementary Figure S7G). Globally, enhancers exhibiting H3K27ac depletion also experienced concomitant H3K4me1 depletion (Figure 4H). We next examined whether VHL-restoration led to acquisition of the H3K27me3 repressive mark. Despite a moderate anti-correlation of H3K27ac and H3K27me3 (786-O cells: r = -0.28, Pearson’s correlation, Supplementary Figure S7H; 12364284 cells: r = -0.22, Pearson’s correlation, Supplementary Figure S7I), H3K27me3 levels remained low at gained enhancers even after VHL restoration (Figure 4H). These findings suggest that VHL restoration may result in a loss of enhancer identity by co-depletion of H3K27ac and H3K4me1, but not a formal transition to a poised enhancer state which would have retained H3K4me1 but acquired H3K27me3.
HIF2α-HIF1β heterodimer is enriched at VHL-responsive enhancers

We sought to investigate which transcription factors might mediate VHL-dependent chromatin remodeling at gained enhancers. Beginning with the primary ccRCC dataset, we looked for enrichment of trans-regulators in gained enhancers over lost enhancers. Using HOMER (65), we found that the top enriched motifs were AP-1 family, ETS family, NFĸB-p65-Rel and HIF1α/2α motifs (Figure 5A, full list of motifs in Supplementary Table S15). For subsequent in vitro validation, we chose c-Jun as a representative AP-1 family member because of its activation in ccRCC (66) and ETS1 as an ETS family representative because of its known interaction with HIF2α (67), but acknowledge that other family AP-1 and ETS family members may play a role in ccRCC. Immunoblotting of c-Jun, ETS1 and NFĸB-p65 showed variable protein expression in both normal and tumor cell lines, but expression of HIF1α and HIF2α restricted to tumor cells only (Figure 5B). HIF2α was expressed in a higher proportion of ccRCC cell lines than HIF1α (Figure 5B). We further examined gene expression of these transcription factors in the TCGA cohort, and found that ETS1, RELA (subunit of NFĸB-p65) and HIF2α were significantly overexpressed in tumors compared to normal tissues, with a range of tumor-association expression patterns similar to variations in ccRCC lines (Supplementary Figure S8A).

To further investigate chromatin occupancy of these factors, we generated ChIP-seq binding profiles of c-Jun, ETS1, NFkB cells and re-examined HIF2α, HIF1α and HIF1β binding profiles from previous literature (21,30), all performed in 786-O cells. Of note, because 786-O cells have lost endogenous HIF1α expression through genomic deletion, the HIF1α ChIP-seq was performed on 786-O cells genetically manipulated to re-express HIF1α protein (30). ChIP-seq results showed that all 6 transcription factors exhibited increased occupancy at gained enhancers compared to lost enhancers, validating the HOMER predictions (Figure 5C).
To determine which of these transcription factors might be directly dependent on VHL, we then compared their protein expression in VHL-mutated isogenic cell lines with and without wild-type VHL restoration. As shown in Figure 5D, VHL restoration consistently downregulated HIF2α expression in both 786-O and 12364284 cells lines but protein levels of other factors displayed contrasting trends between the two cell lines, implying that amongst the 6 factors examined, HIF2α protein expression was the most VHL-dependent. Indeed, supporting an important role for HIF2α in VHL-dependent enhancer remodeling, only HIF2α and HIF1β were significantly enriched at enhancers showing VHL-dependent H3K27ac depletion (Figure 5E). Moreover, amongst all known motifs in the HOMER database, HIF2α was the most enriched motif at VHL-responsive enhancers exhibiting H3K27ac depletion (p-value = 1x10^{-11}, Supplementary Table S16).

In contrast, HIF1α was not enriched at enhancers showing H3K27ac depletion (Figure 5E). Despite sharing many binding sites with HIF2α, HIF1α predominantly localized to promoter-proximal regions whereas HIF2α frequently occupied introns and intergenic regions in 786-O cells (Supplementary Figure S8B), consistent with a promoter-centric occupancy of HIF1α and an enhancer-centric occupancy of HIF2α (Figure 5F). Gained enhancers displayed a HIF2α occupancy twice that of tumor-specific promoters (p-value < 1x 10^{-16}, proportions test) in 786-O cells, suggesting that HIF2α may play a greater role at regulating enhancers than promoters.

To extend these HIF1α and HIF2α occupancy patterns findings to a system that expresses endogenous levels of both factors, we then performed HIF1α and HIF2α ChIP-seq in 40911432 ccRCC cells which abundantly co-express both HIFα subunits (Figure 5B). Similar to 786-O, in 40911432 cells, HIF1α showed a preferential occupancy at promoter-proximal regions while a large proportion of HIF2α were found in distal regions (introns and distal intergenic regions) (Supplementary Figure S8C). A higher proportion of HIF1α binding sites overlapped with gained promoters than HIF2α (68% of HIF1α vs. 41% of HIF2α, p-value = 0.002, proportions
test) (Figure 5G). Conversely, a higher proportion of HIF2α binding sites overlapped with gained enhancers than HIF1α (29% of HIF1α vs. 51% of HIF2α, p-value < 2.2 x 10^{-16}, proportions test). HIF2α’s preferential occupancy at enhancers was further substantiated by its higher enrichment at enhancers showing H3K27ac depletion after VHL restoration than HIF1α (Figure 5H). Specific examples of VHL-responsive enhancers bound exclusively by HIF2α but not HIF1α included an enhancer near Ubiquitin Protein Ligase E3 Component N-Recognin 4 (UBR4) (Figure 5I) and a super-enhancer near C-Maf Inducing Protein (CMIP) (Figure 5J).

Therefore, even in HIF1α/ HIF2α co-expressing ccRCC cells, these results suggest that HIF2α plays a greater role in VHL-mediated enhancer remodeling than HIF1α.

**HIF2α-HIF1β bound enhancers modulate gene expression**

To investigate the extent to which HIF2α silencing is sufficient to recapitulate the effects of VHL restoration, we performed H3K27ac ChIP-Seq and RNA-Seq in 786-O cells with HIF2α siRNA-mediated knockdown, and analyzed correlations between HIF2α siRNA knockdown and VHL restoration. When assessed against all genes, there was a low correlation (r = 0.1, p-value = 5.2 x 10^{-31}) between HIF2α knockdown and VHL restoration. Importantly however, this correlation increased to 0.23 (p-value = 5.8 x 10^{-14}) for genes nearby HIF2α binding sites (Figure 6A).

Similar results were obtained at the epigenomic level, where for gained enhancers the correlation was low at 0.06 across all gained enhancers (p-value = 1.9 x 10^{-5}) but increased substantially to 0.37 (p-value = 9.5 x 10^{-8}) at HIF2α-bound enhancers (Figure 6B) and at super-enhancers increased from 0.089 (p-value = 0.0025) to 0.25 (p-value = 0.00054) at HIF2α-bound super-enhancers (Figure 6C). As a visual example, H3K27ac signals at the ZNF395 super-enhancer were diminished after VHL restoration or HIF2α knockdown, concomitant with decreased ZNF395 gene expression (Figure 6D). Validation by RT-qPCR showed that HIF2α
siRNA knockdown downregulated VEGFA, SLC2A1 and ZNF395 expression to a comparable degree as VHL restoration (Figure 6E). Decreases in luciferase reporter activity of enhancer elements were also consistent between HIF2α siRNA knockdown and VHL restoration (Figure 6F).

We sought to establish a causal link between HIF2α-bound enhancers and control of gene expression. We performed Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) mediated genomic depletion of a ZNF395 enhancer region with the highest HIF2α peak (Figure 6G). All four clones with homozygous deleted ZNF395 enhancer consistently downregulated their ZNF395 expression compared to clones with the intact enhancer (p<0.05), providing evidence that ZNF395 expression is epigenetically controlled by HIF2α-HIF1β-bound enhancer (Figure 6G). Taken together, these results indicate that HIF2α is likely an important mediator of VHL-driven enhancer remodeling.

**VHL restoration reduced P300 recruitment but preserved promoter-enhancer interactions**

Finally, we sought to investigate why VHL restoration caused a decrease in H3K27ac levels. Previous pull-down assays have reported that both HIF2α and HIF1β can interact with histone acetyltransferase P300 (68-71). Indeed, P300 frequently marks enhancers (43) and is thought to be recruited by tissue-specific transcription factors (72). However, chromatin profiles of P300 have not been previously established in kidney cancer cell lines, so the contribution of P300 in shaping enhancers in ccRCC remains unclear. Therefore, we performed P300 ChIP-seq in 786-O cells, and confirmed its enrichment at gained enhancers over lost enhancers (Figure 7A). Comparing P300 ChIP-seq with HIF2α CHIP-seq yielded a surprisingly high degree of overlap between HIF2α and P300 (96%), even more than that of HIF2α and HIF1β (89%) (Figure 7B,
7C). In contrast, other transcription factors such as c-Jun, ETS1 and NFκB did not exhibit such high degree of overlap (≤60%) (Figure 7B).

We compared P300 binding at tumor enhancers with and without VHL. Despite increased P300 protein levels in 786-O cells after VHL restoration (Figure 7D), binding of P300 decreased across all 4 enhancers examined (Figure 7E). HIF2α depletion by siRNA knockdown also decreased P300 recruitment (Figure 7F), suggesting that loss of HIF2α may interfere with P300 recruitment.

We investigated whether VHL restoration and the subsequent loss of P300 binding disrupted promoter-enhancer interactions. We performed Capture-C of enhancer regions in paired 786-O cell lines with and without VHL restoration. Surprisingly, Capture-C interactions showed a relatively high correlation between VHL-deficient and VHL-restored 786-O cells at VHL-responsive regions (r = 0.74, Pearson’s correlation), even higher than correlations observed at non-VHL-responsive regions (r = 0.57, Pearson’s correlation) (Figure 7G). As a visual example, interactions between the VEGFA promoter and enhancer were intact even after VHL restoration (Figure 7H), indicating that loss of enhancer activity was not sufficient to dissociate promoter-enhancer interactions. Furthermore, many of these promoter-enhancers were lineage-specific, for example the interaction between SLC2A1 enhancer with its promoter was not detected in KATOIII, a gastric cancer cell line (Supplementary Figure S9). Therefore, promoter-enhancer interactions often pre-exist in kidney cells, frequently in a tissue-specific manner.
Discussion

Understanding epigenomic alterations and their genetic origin can identify new disease mechanisms (34), vulnerabilities (73,74) and therapeutic strategies (75-77). Through comprehensive profiling of histone modifications in primary normal-tumor pairs and cell lines, we generated a compendium of ccRCC-associated promoters and enhancers. Our study demonstrates that the most frequent ccRCC mutational event – VHL inactivation – leads to genome-wide enhancer and super-enhancer remodeling, which directly imparts ccRCC hallmarks including angiogenesis and metabolic reprogramming. ZNF395, epigenetically controlled by a VHL-responsive super-enhancer, emerged as a crucial regulator of ccRCC tumorigenesis.

Our work has three main advances. Firstly, to our knowledge, this is the most comprehensive atlas of histone profiles in ccRCC and will likely provide an invaluable resource to the ccRCC field. Using high-resolution multiplexed interactome data (Capture-C (56)) and H3K27ac-expression correlation, we minimized ambiguity in enhancer assignment, and further confirmed the dependency of enhancers on VHL/HIF status by reporter assays. Secondly, using isogenic cell lines, we show that VHL loss contributes significantly to enhancer remodeling. Even though another mutation in ccRCC, SETD2, can mediate widespread increases in chromatin accessibility (46) and DNA hypomethylation (78), its relatively low mutation frequency at ~10% in all ccRCC tumors (78) cannot explain epigenetic abnormalities in the vast majority of SETD2-widetype tumors. Lastly, an examination of somatically altered super-enhancers enabled us to identify a novel master regulator crucial to the pathogenesis of ccRCC, ZNF395. Even though ZNF395 overexpression in ccRCC has been previously reported (79-81) and its proximity to a super-enhancer was independently noted (42), our study is the first to pinpoint the specific VHL-dependent enhancer required for ZNF395 expression, and to show ZNF395's indispensable functional role for ccRCC tumorigenesis in vitro and in vivo.
Our data suggests that mechanistically, loss of VHL stabilizes HIF2α occupancy at tumor-specific gained enhancers, which in turn recruits histone acetyltransferase P300 (28,82) to maintain H3K27 acetylation, upregulating expression of ccRCC-specific genes such as ZNF395 (Figure 7I). Restoration of wild-type VHL resulted in co-depletion of H3K27ac and H3K4me1 marks and thus abrogation of active enhancer identity at tumor-associated enhancers. Compared to the promoter-centric occupancy of HIF1α, HIF2α is predominantly found at enhancers, pointing towards a key difference between HIF1α and HIF2α. We also found that HIF2α siRNA knockdown specifically attenuates the activity of HIF2α-bound enhancers/super-enhancers. Interestingly, the majority of promoter-enhancer interactions remained largely unaltered by VHL status, suggesting that these promoter-enhancer couplings are largely stable and pre-formed. This is consistent with a recent report demonstrating that promoter-enhancer interactions remain largely unchanged between normoxia and hypoxia (29). Our study demonstrating VHL’s impact on chromatin remodeling also suggests that other cancer genes with high tumor-type specific mutational penetrance, such as BRAF in melanoma (83) and APC in colon cancer (84) may also act to modify cellular epigenomes to effect broad yet disease-specific cellular changes, despite these genes not being classical chromatin modifiers.

Besides VHL, other mutations such as PBRM1, SETD2, ARID1A, SMARCA4, JARID1C, KDM6A/UTX have been reported in ccRCC (47,85), and these are likely to augment VHL’s core transcriptional effects (86), contributing to heterogeneity in disease phenotypes (87) and progression patterns (88). Using the example of 786-O cells, at least two other mutations in these cells may have a direct impact on chromatin - MLL3 (p.A3902G) and gain of function TP53 mutations (p.R248W). MLL3, a histone 3 lysine 4 methyltransferase, is directly responsible for formation of the H3K4me1 enhancer mark (89,90), and plays a critical role for enhancer regulation (91). Gain of function TP53 mutants also bind aberrantly to chromatin, especially near methyltransferases MLL1 and MLL2, potentially contributing to tumor growth via...
chromatin deregulation (92). Besides mutations, structural variants are also known to alter enhancers via enhancer hijacking (93) or copy number gains (94) in other cancers. Given the multitude of driver and bystander mutations in ccRCC, it is thus unlikely that VHL alone can account for all epigenomic changes observed in this tumor type. Nevertheless, by integrating data across multiple ccRCC cell lines, our data suggests that VHL inactivation is likely to account for almost a third (32%) of all gained enhancer regions, supporting its role as a dominant driver of epigenetic abnormalities in ccRCC despite the presence of other genetic changes.

Our epigenetic maps contain a wellspring of both well-validated and uncharacterized targets that may contribute to ccRCC tumorigenesis. We found extensive enhancer gains around well-characterized hypoxia-related targets (95) (VEGFA, CXCR4 (96), HK2), SLC-mediated membrane transporters (SLC2A1, SLC2A2, SLC38A1 (97)), SLC16A family (98)), and adipogenesis (PLIN2 (51,99)). New targets revealed in this study include SMPDL3A which could be another important ccRCC-specific oncogene given its role in lipid and cholesterol metabolism (61,100). Genes associated with lost super-enhancers, which could only be identified with normal-tumor pairs, implicated potential tumor suppressors (EHF, MAL, GCOM1 and HOXB9) that warrant further investigation.

One notable finding from this epigenomic study is the tumorigenic requirement of ZNF395 in ccRCC. ZNF395 is also known as Huntington’s disease gene regulatory region-binding protein-2 (HDBP2) (101) or papillomavirus binding factor (PBF) (102). ZNF395 is required for the differentiation of mesenchymal stem cells to adipocytes, by partnering with PPARγ2 to promote adipogenesis (103). ZNF395 has been shown to bind to the promoters of Huntington’s gene (101), interferon-induced genes, and to cause upregulation of cancer-related genes (MACC1, PEG10, CALCOCO1, MEF2C) (104) and pro-angiogenic chemokines including IL6 and IL8 under hypoxia (105). It remains to be elucidated in future studies the precise mechanism contributing to ZNF395’s tumorigenic role.
The enhancer landscapes profiled in this study have implications beyond ccRCC. The poorly perfused tumor core makes hypoxia a feature of virtually all solid tumors (106). MCF7 cells under hypoxia (but not normoxia) share similar H3K27ac profiles as 786-O (29). While ZNF395 is highly expressed in ccRCC, its low basal expression can be upregulated upon hypoxia in other cancer types including glioblastoma and skin cancer (104,105). Targeting ZNF395 or its downstream effectors in future studies may be therapeutically relevant to both ccRCC and other hypoxic solid malignancies. Direct targeting of ZNF395 using a peptide-based cancer vaccine is undergoing phase I trials in sarcoma patients (107-109), opening up the possibility of using immunotherapy to target the extracellular fragments of nuclear master regulators. Our study suggests that initiating a similar trial in ccRCC may be worthwhile. Moreover, given the recent progress in targeting transcription factors using various modalities including small molecules and stapled peptides (110-112), inhibitors of ZNF395 may provide an important therapeutic inroad for ccRCC treatment.
Methods

Patient information

Fresh frozen normal-tumor tissues were obtained from nephrectomy cases under approvals from institutional research ethics review committees and patient consent under IRB protocol 2010/735/B. Normal tissues were harvested from sites distant from the tumor. Refer to Supplementary Table S1 for detailed patient information.

Cell lines

Commercial cell lines (786-O, A-498, HK-2, PCS-400) were purchased from ATCC in 2000. Cell lines were maintained in RPMI (Invitrogen) with 10% FBS with the exception of primary renal proximal tubule epithelial cells, PCS-400, which were maintained in Renal Epithelial Cell Basal Medium (ATCC). Cell line authentication was performed by short tandem repeat (STR) analysis (Cancer Science Institute of Singapore) in 2015 against publicly available STR profiles. Mycoplasma testing was performed using the MycoSensor PCR assay kit (Stratagene).

Establishment of tumor-derived cell lines from primary tumors

Tumor cells were disassociated from primary tumors by collagenase, seeded and maintained in RPMI with 10% FBS. At 80-90% confluency, the cells were passaged at a 1:3 ratio. Cultured cells were considered to be successfully immortalized after 60 passages. Correct pairing of tumor tissues and cell lines was achieved by comparing the percentage identity of single nucleotide polymorphisms (SNPs) based on targeted sequencing. All tumor-cell line pairs showed identities of > 90% whereas shuffling of pairing showed identities < 80%. Tumors and cell lines from 12364284 and 40911432 showed the same VHL mutations but 86049102 tissue (named 86049102T) is VHL mutant while the cognate 86049102 cell line (named 86049102L) is VHL wild-type.
Stable VHL restoration in ccRCC lines

786-O cells (WT2, VHL+) and 786-O cells (RC3, VHL−) were kindly provided by Dr. Michael Ohh (University of Toronto). Stable transduction of VHL was performed in A-498, 12364284 and 40911432 cells as follows: HA-VHL wt-pBabe-puro plasmid (a gift from Dr. William Kaelin, Dana Farber Cancer Institute, Addgene plasmid # 19234) was transfected into PlatA cells (RV-102, Cell Biolabs) at 2 µg DNA/well of a 6-well plate using Lipofectamine 3000 (LifeTechnologies). A medium change was performed 10-16 hrs after transfection. The supernatant from PlatA cells containing retroviruses was harvested 48 hours later, and added to ccRCC cells, which were then selected with puromycin for 3 days post transduction.

Histone Nano-ChIP-Seq

Nano-ChIP-Seq was performed as previously described (113) with slight modifications. Fresh-frozen cancer and normal tissues were dissected using a razor blade to obtain ~5 mg of tissue. The tissues were fixed in 1% formaldehyde for 10 min at room temperature. Fixation was stopped by addition of glycine to a final concentration of 125 mM. Tissue pieces were washed 3 times with TBSE buffer. Pulverized tissues were lysed in 100 µl lysis buffer and sonicated for 16 cycles (30s on, 30s off) using a Bioruptor (Diagenode). The following antibodies were used: H3K27ac (ab4729, Abcam), H3K4me3 (07-473, Millipore), H3K4me1 (ab8895, Abcam) and H3K27me3 (07-449, Millipore). The total volume of immunoprecipitation was 1 ml and the amount of antibody used was 2 µg. The input DNA was precleared with protein G Dynabeads (Life Technologies) for 1 hr at 4°C and then incubated with antibodies conjugated protein G beads overnight at 4°C. The beads were washed 3 times with cold wash buffer. After recovery of ChIP and input DNA, whole-genome-amplification was performed using the WGA4 kit (Sigma-Aldrich) and Bpml-WGA primters. Amplified DNA was digested with Bpml (New England Biolabs (NEB)). After that, 30 ng of the amplified DNA was used with the NEBNExt ChIP-Seq
library prep reagent set (NEB). ChIP-seq in cell lines were performed using the same Nano-ChIP-seq protocol described above but with $1 \times 10^6$ cells. Each library was sequenced to an average depth of 20-30 million raw reads on HiSeq2500 using 101bp single end reads.

**Histone ChIP-Seq analysis**

Sequencing tags were mapped against the human reference genome (hg19) using Burrows-Wheeler Aligner (BWA-mem) (114)(version 0.7.10). Reads were trimmed 10 bp from the front and the back to produce 81 bp. Only reads with mapQ >10 and with duplicates removed by rmdup were used for subsequent analysis. Significant peaks were called using CCAT ($p$-value < 0.05) (115). The strength and quality of immunoprecipitation was assessed using CHANCE (116).

**Transcription factor Chip-Seq**

For each transcription factor, $3 \times 10^7$ cells were cross-linked with 1% formaldehyde for 10 min at room temperature, and stopped by adding glycine to a final concentration of 125 nM. Chromatin was extracted and sonicated to ~500bp (Vibra cell, SONICS). The following antibodies were used for chromatin immunoprecipitation, c-Jun (sc-1694, Santa Cruz), NFκB p65 (sc-372, Santa Cruz), ETS1 (sc-350, Santa Cruz), HIF1α (610959, BD Biosciences), and HIF2α (NB100-122, Novus Bio) and P300 (sc-585, Santa Cruz). The total volume of immunoprecipitation was 1.5 ml and the amount of antibody used was 15 µg. Input DNAs were precleared with protein G Dynabeads (LifeTechnologies) for 2 hr at 4°C and then incubated with antibody-conjugated protein G beads overnight at 4°C. The beads were washed 6 times with wash buffer at room temperature. At least 10 ng of the DNA was used with the NEBNext ChIP-Seq library prep reagent set (NEB). Each library was sequenced to an average depth of 30-50 million reads on a HiSeq2500 using 101 bp single end reads.
Capture C

Capture-C was performed as previously described (56). Briefly, $1 \times 10^7$ cells were crosslinked by 2% formaldehyde, followed by lysis, homogenization, DpnII digestion, ligation, and de-crosslinking. DNA was sonicated using a Covaris to 150-200bp to produce DNA suitable for oligo capture. A total of 3 µg of sheared DNA was used for sequencing library preparation (New England Biolabs). Enhancer sequences were double captured by hybridisation to customized biotinylated oligos (IDT) and enriched with Dynabeads (Life Technologies). Captured DNA was sequenced to an average depth of 2 million reads per probe on the Hiseq Illumina platform using 150 bp paired-end reads.

Capture C analysis and Gene assignment

Preprocessing of raw reads was performed to remove adaptor sequences (trim_galore, http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) and overlapping reads were merged using FLASH (117). In order to achieve short read mapping to the hg19 reference genome, the resulting preprocessed reads were then in-silico digested with DpnII and aligned using Bowtie (using p1, m2, best and strata settings). Aligned reads were processed using Capture-C analyser (118) to (i) remove PCR duplicates, (ii) classify sub fragments as ‘capture’ if they were contained within the capture fragment; ‘proximity exclusion’ if they were within 1Kb on either side of the capture fragment; or ‘reporter’ if they were outside of the ‘capture’ and ‘proximity exclusion’ regions, and (iii) normalize read counts per 100,000 interactions in bigwig format. We additionally used the r3Cseq package (119) on the capture and reporter fragments to identify significant interactions of the viewpoint against a scaled background ($q$-value <0.05). Gene assignment is defined by the overlap of significant Capture C peaks with genes with start and end defined by GENCODE v19. Interactions were plotted using Epigenome Gateway v40.0.
Identification of differentially enriched regions

Significant H3K27ac peaks called by CCAT were merged across all normal-tumor samples. The same was performed with H3K4me1 and H3K4me3 ChIP-seq data. Transcription start sites were based on GENCODE v19. Promoters were defined as regions of overlap between H3K27ac and H3K4me3 and also overlapping with ±2.0 Kb around the transcription start site. Enhancers were defined as regions of overlap between H3K27ac and H3K4me1 but not overlapping with promoters. To minimize stromal contamination, we performed further filtering using cell line data, where enhancers and promoters not overlapping with H3K27ac peaks in any of the cell lines were discarded. Wiggle files of window size 50 bp were generated using MEDIPs (120) from bam files. The input-subtracted signal for each promoter or enhancer region was computed using bigWigAverageOverBed to yield reads per kilobase per million (RPKM). The RPKM of H3K27ac, H3K4me1 and H3K4me3 ChIP-Seq from promoters and enhancers were corrected for batch effects using Combat. Tumor specific regions were defined as regions that have a fold difference of ≥2, and a difference of 0.5 RPKM from patient-matched normal tissue. Normal regions were defined as regions that have a fold difference of ≤ 0.5, and a difference of -0.5 RPKM from the corresponding regions in patient-matched tumor. Recurrently gained regions were defined as gain in ≥ 5/10 patients and no loss in any patients. Recurrently lost regions were defined as loss in ≥ 5/10 patients and no gain in any patients. Statistical testing for each cis-regulatory region was performed using paired t-tests with Benjamini-Hochberg correction. The differential regions were visualized using NGSplot (121).

Identification of super-enhancer regions

Super-enhancer regions were identified using ROSE (35) (with promoter excluded), using H3K27ac peak regions merged from all patients (both normal and tumor tissue). Wiggle files of window size 50 bp were generated using MEDIPs (120) from bam files. The input-subtracted
signal for each super-enhancer was computed using bigWigAverageOverBed (sum of reads over covered bases). The super-enhancer regions were ranked by the average difference of normal-tumor H3K27ac ChIP-seq signals. Gained super-enhancers were defined as regions that have average differential H3K27ac ChIP-seq signals >0. Lost super-enhancers were defined as regions that have average differential H3K27ac ChIP-seq signals <0.

Additional methods can be found in Supplementary Methods.

Date Accession

ChIP-seq and RNAseq data are available at Gene Expression Omnibus (GSE86095).


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Figure legends

Figure 1: *VHL* deficient ccRCC tumors exhibit an aberrant cis-regulatory landscape

(A) Putative active promoters are defined by co-occurrence of H3K4me3, H3K27ac and proximity to transcription start sites (TSSs) within 2 kb. Putative active enhancers are defined by the presence of H3K4me1, H3K27ac and exclusivity with promoters. (B) PCA analysis using all 17,497 promoters and 66,448 enhancers classify normal and tumors into distinct clusters. Patient IDs: (1-12364284; 2-17621953; 3-20431713; 4-40911432; 5-57398667; 6-70528835; 7-74575859; 8-77972083; 9-86049102; 10-75416923) (C) Heat maps show H3K27ac levels of altered promoters and enhancers in a paired patient tissue (patient 40911432, yellow high, black low). (D) The H3K27ac levels, chromatin accessibility (FAIRE-seq), DNA methylation of gained promoters and enhancers, and gene expression of the nearest ccRCC IncRNA are compared between normal and tumor tissues. ***p-value < 0.001, two-sided t-test. (E) Shown are tracks of histone ChIP-seq (H3K27ac, H3K4me1, H3K4me3) and RNA-Seq at the *CCND1* locus in a tumor-normal pair of patient 40911432. The histone ChIP-seq profiles of normal adult kidney tissue from the Epigenome Roadmap are displayed above the normal tissue generated by Nano-ChIP-seq for comparison. A cell line was derived from the tumor tissue and its histone profile is displayed below its matching primary tissue. This enhancer is known to interact with the *CCND1* promoter from a previous study (Schodel, 2012 (21)), and is situated close to a RCC susceptibility SNP rs7105934 (49).

Figure 2: Enhancer aberration is a signature of ccRCC

(A) Enriched pathways associated with gained promoters and enhancers revealed by GREAT (ranked by binomial FDR q-value). Red bars refer to ccRCC-specific pathways. (B) De novo enhancers are acquired in a ccRCC tumor tissue upstream of *VEGFA*. Capture-C confirmed interactions of this *VEGFA* enhancer (E) with its promoter (P) in 786-O cells. The arcs represent significant interactions detected by r3Cseq (q<0.05). The input-subtracted H3K27ac signals of this enhancer are highly correlated with *VEGFA* gene expression (Spearman’s correlation). (C) Similar to (B), a de novo tumor enhancer interacts with the *SLC2A1/GLUT1* promoter.

Figure 3: Tumor super-enhancers identify key oncogenic drivers

(A) A total of 1,451 super-enhancers are identified by ROSE and ranked by their differential H3K27ac intensity between normal and tumor tissues. Genes associated with the top gained and lost super-enhancers are listed. (B) TCGA RNA-seq data indicates that genes associated with top 10 gained enhancers are upregulated in tumors while genes associated with top 10 lost enhancers are downregulated. This tumor-specificity is restricted to ccRCC, but not the other two RCC subtypes, papillary and chromophobe. (C) Expression of ZNF395 and SMPDL3A are measured in a panel of normal kidney cell lines (black) and ccRCC cell lines (red) by RT-qPCR. (D) Pooled siRNA against ZNF395 inhibits colony formation of A-498 and 786-O but not HK-2 normal immortalized kidney cells. Pooled siRNA against SMPDL3A inhibits colony formation of A-498 but not 786-O. (E) H3K27ac ChIP-seq shows an active ZNF395 super-enhancer only in ccRCC cells (A-498 and 786-O) but not normal kidney cells (PCS-400, HK-2). (F) TCGA RNA-seq data shows exclusive overexpression of ZNF395 amongst 12 cancer types. (G-I) ZNF395 inhibition by two shRNA clones decreases colony formation (G), in vitro proliferation (H) and increases apoptosis measured by cleavage of Caspase3/7 substrate (I). *p-value < 0.05, two-sided t-test. (J) ZNF395 inhibition by shRNA leads to total elimination of A-498 tumors in vivo and delayed 786-O tumor growth. NC: n=7, shZNF395-1: n=7, shZNF395-2: n=6

Figure 4: *VHL* deficiency remolds ccRCC enhancers

(A) Log fold changes of H3K27ac ChIP-seq signals at gained promoters, enhancers and super-enhancers as defined in the primary ccRCC dataset after *VHL* restoration in 786-O cells. Red dots represent cis-
regulatory elements with significant changes (p-value <0.05, negative binomial) in H3K27ac levels after VHL restoration. The number and percentage of altered regions (p-value <0.05, negative binomial) are shown at the upper and lower right corners. (B) Read coverage of H3K27ac ChIP-seq at VHL-responsive enhancers in VHL-mutant ccRCC cell lines (red) compared to VHL-wild-type ccRCC (grey), and normal kidney cell lines (green) and 31 other cancer cell lines (black). (C) Changes in expression of genes linked to VHL-responsive tumor enhancers after VHL restoration in 786-O cells. *p-value < 0.05, two-sided t-test. (D-G) Examples of lost VHL-responsive enhancer/super-enhancers are associated with EGFR (D), CCND1 (E), ITGB3 (F) and VEGFA (G) in 786-O cells. (H) Log fold changes of H3K27ac (red), H3K4me1 (blue) and H3K27me3 (brown) signals at gained enhancers showing H3K27ac depletion after VHL restoration in 786-O cells.

Figure 5: HIF2α is enriched at enhancers of VHL-responsive tumor tumors

(A) Motif analysis of gained enhancers using HOMER reveals significant enrichment of AP-1 family, ETS family, NFκB and HIF1α/2α (hypergeometric test). Lost enhancers were used as background in the motif search to identify tumor-specific transcription factors. (B) Protein expressions of putative transcription factors enriched at gained enhancers in 9 tumor cell lines (4 commercial cell lines and 5 patient-derived cell lines*) and 2 normal cell lines. ACHN is a papillary RCC cell line. (C) ChIP-seq validated the enrichment of transcription factors in gained enhancers over lost enhancers. (D) Protein expression of transcription factors are shown in 786-O and 12364284 cells with and without wild-type VHL. (E) Transcription factor binding at VHL-responsive gained enhancers shows enrichment of HIF2α and HIF1β at enhancers with H3K27ac depletion (red) over regions with H3K27ac enrichment (black) after VHL restoration. (F) ChIP-Seq data shows distribution of exogenous HIF1α and endogenous HIF2α binding at altered promoters and enhancers in 786-O cells that have been genetically engineered to overexpress HIF1α. (G) ChIP-Seq shows distribution of endogenous HIF1α and HIF2α binding at altered promoters and enhancers in 40911432 cells. (H) Transcription factor binding at VHL-responsive enhancers shows higher enrichment of HIF2α than HIF1α at enhancers with H3K27ac depletion after VHL restoration (red) over regions with H3K27ac enrichment after VHL restoration (black). (I) Example of a VHL-responsive enhancer near UBR4 with only HIF2α binding but not HIF1α binding. (J) Example of a VHL-responsive super-enhancer near CMIP with only HIF2α binding but not HIF1α binding.

Figure 6: HIF2α-HIF1β bound enhancers modulate gene expression

(A) Pearson’s correlation of gene expression changes after either VHL restoration or HIF2α siRNA knockdown at all genes or genes adjacent to HIF2α binding sites. (B) Pearson’s correlation of H3K27ac changes after VHL restoration and HIF2α siRNA knockdown at either all gained enhancers or HIF2α-bound enhancers adjacent to binding sites. (C) Pearson’s correlation of H3K27ac changes after VHL restoration and HIF2α siRNA knockdown at either all gained super-enhancers or HIF2α-bound super-enhancers adjacent to binding sites. (D) Changes in RNAseq and H3K27ac ChIP-Seq signals after VHL restoration or HIF2α siRNA knockdown at ZNF395 super-enhancer (SE), together with binding profiles of transcription factors enriched at enhancers. (E) Both VHL restoration and HIF2α siRNA knockdown decreases expression of genes with HIF2α-bound enhancers in 786-O cells. *p-value < 0.05, two-sided t-test. (F) Both VHL restoration and HIF2α siRNA knockdown decrease enhancer activities measured by luciferase reporter assay in 786-O cells. *p-value < 0.05, two-sided t-test. (G) RT-qPCR measurement of ZNF395 expression in 4 wild-type clones (black) and 4 clones with ZNF395 enhancer depleted by CRISPR (red). Depleted region has the highest HIF2α binding at ZNF395 super-enhancer in 786-O cells (deleted region indicated in D). *p-value < 0.05, two-sided t-test.

Figure 7: VHL restoration reduces P300 recruitment but preserves promoter-enhancer interactions

(A) Enrichment of P300 binding at gained and lost enhancers based on ChIP-Seq. (B) Percentage of overlap between HIF2α and other transcription factors. (C) ChIP-Seq binding profiles of HIF2α and P300.
(D) Protein expression of P300 with and without VHL in 786-O and 12364284 cells measured by immunoblotting. (E) ChIP-qPCR of P300 binding at enhancers with and without VHL restoration in 786-O cells. NC – negative control regions. (F) ChIP-qPCR of P300 binding at enhancers with and without HIF2α siRNA knockdown in 786-O cells. NC – negative control regions. (G) Correlation of enhancer interactions measured by Capture-C (RPM – reads per million) between 786-O cells with and without VHL restoration at both VHL-responsive and non-VHL-responsive enhancers (H) Capture-C shows that VEGFA enhancer-promoter interactions are maintained even after VHL restoration. E – enhancer; P – promoter. (I) Schematics of VHL-driven enhancer aberration in ccRCC.
Figure 1

A

Promoter

Enhancer

-2kb TSS +2kb

H3K27ac

H3K4me3

H3K4me1

Regions | Definition | Total | Recurrently Gained in Tumor | Recurrently Lost in Tumor
--- | --- | --- | --- | ---
Promoter | H3K27ac+/H3K4me3+/-2kb TSS | 17,497 | 4,719 | 592
Enhancer | H3K27ac+/H3K4me1+/promoter | 66,448 | 4,906 | 5,654

B

PC1

PC2

Promoters

Enrichers

Normal Tumor

Fold enrichment of H3K27ac over input

Gain Promoters

Lost Promoters

Gain Enhancers

Loss Enhancers

Distance from TSS

-3kb TSS -3kb Center -3kb TSS -3kb Center -3kb TSS -3kb Center -3kb TSS -3kb Center

Distance from H3K27ac

summits

C

Fold enrichment of H3K27ac over input

Gain Promoters

Loss Promoters

Gain Enhancers

Loss Enhancers

Normal tissue Tumor tissue

Distance from TSS

-3kb TSS -3kb Center -3kb TSS -3kb Center -3kb TSS -3kb Center

D

Gain Promoters

Gain Enhancers

Normal tissue Tumor tissue

E

Chr11:69,196,493-69,539,519

CCND1

RNAseq

Normal tissue

Tumor tissue

Roadmap (Normal tissue)

Tumor derived cell line

Roadmap (Tumor tissue)

FAIREseq

H3K27ac

H3K4me3

H3K4me1

Normal tissue

Tumor tissue

Tumor derived cell line

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Figure 2

A

Gained Promoters

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- Cell Cycle
- Transcription
- RNA Metabolism
- mRNA Metabolism
- Deposition of CENPA-containing Nucleosomes
- RNA Polymerase I Promoter Opening
- RNA Polymerase I Transcription
- Chromosome Maintenance
- mRNA Splicing
- Meiotic Recombination

Gained Enhancers

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</table>

- HIF-1-alpha network
- Platelet activation
- Direct p53 effectors
- Renal cell carcinoma
- TCPTP Signaling
- AP-1 network
- PDGFR-beta signaling
- SLC-mediated transmembrane transport
- HGFR/c-MET
- ERK1/ERK2 MAPK Pathway

B

chr6:43,585,023-43,770,738

VEGFA

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Spearman’s correlation r: 0.83

C

chr1:43,365,540-43,566,851

SLC2A1

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<td>H3K27ac</td>
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Spearman’s correlation r: 0.72
**Figure 3**

**A** H3K27ac (T-N) raw counts

**B** ccRCC (n=72)

**C** Papillary (n=32)

**D** Chromophobe (n=25)

**E** chr8:28,192,829-28,284,799  ZNF395

**F** ZNF395 expression (RSEM)

**G** NC shZNF395-1 shZNF395-2

**H** Relative growth rate

**I** Cleaved Caspase3/7 substrates

**J** Tumor volume (mm3)
Figure 4

A. Gained promoters (786-O)

- Up p<0.05: 428 (8.7%)
- Down p<0.05: 321 (6.5%)

B. Enhancers with H3K27ac depletion after VHL restoration

C. Enhancers with H3K27ac enrichment after VHL restoration

D. EGFR

E. CCND1

F. ITGB3

G. VEGFA

H. H3K27ac, H3K4me1, and H3K27me3

Distance from H3K27ac summits

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Figure 5

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<th>Family</th>
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C

Gained enhancers —— Lost enhancers

D

786O 12364284

VHL

1.9 1.9 0.34

c-Jun

1.69 1.69 0.49

ETS1

1.21 1.21 0.64

NFKb-p65

0.12 0.12 0.43

HIF2α

Beta-Actin

E

Enhancers with H3K27ac depletion after VHL

Enhancers with H3K27ac enrichment after VHL

F

HIF1α (786-O)

Gained Enh | Lost Prom | Gained Prom | Lost Enh | Gained Enh | Lost Prom | Gained Prom | Lost Enh |

HIF2α (786-O)

Gained Enh | Lost Prom | Gained Prom | Lost Enh | Gained Enh | Lost Prom | Gained Prom | Lost Enh |

G

HIF1α (40911432)

Gained Enh | Lost Prom | Gained Prom | Lost Enh | Gained Enh | Lost Prom | Gained Prom | Lost Enh |

HIF2α (40911432)

Gained Enh | Lost Prom | Gained Prom | Lost Enh | Gained Enh | Lost Prom | Gained Prom | Lost Enh |

H

40911432

Enhancers with H3K27ac depletion after VHL

Enhancers with H3K27ac enrichment after VHL

I

chr1:19,352,768-19,383,967

H3K27ac (EV)

H3K27ac (VHL)

HIF1α

HIF2α

J

chr16:81,512,501-81,528,600

H3K27ac (EV)

H3K27ac (VHL)

HIF1α

HIF2α

K

Read count per million reads of TF ChIP-Seq

Distance from H3K27ac summits

L

CMIP

chr1:81,512,501-81,528,600

H3K27ac (EV)

H3K27ac (VHL)

HIF1α

HIF2α

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Gained enhancers (all regions)

Gained super-enhancer (all regions)

r = 0.25
r = 0.089

Gene expression (all genes)

Gene expression (HIF2α-bound)

log2 fold change in H3K27ac (HIF2a siRNA/NT)
log2 fold change in H3K27ac (VHL/EV)

log2 fold change in H3K27ac (HIF2a siRNA/NT)
log2 fold change in H3K27ac (VHL/EV)

Gene expression (EV)
Gene expression (VHL)
Gene expression (NT)
Gene expression (HIF2ai)

CRISPR deletion of ZNF395 enhancer
ZNF395 gene expression normalized to ACTB

Figure 6
VHL deficiency drives enhancer activation of oncogenes in clear cell renal cell carcinoma

Xiaosai Yao, Jing Tan, Kevin Junliang Lim, et al.

Cancer Discov  Published OnlineFirst September 11, 2017.

Updated version  Access the most recent version of this article at: doi:10.1158/2159-8290.CD-17-0375

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