Whole-Exome Sequencing Defines the Mutational Landscape of Pheochromocytoma and Identifies KMT2D as a Recurrently Mutated Gene

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As subsets of pheochromocytomas (PCCs) lack a defined molecular etiology, we sought to characterize the mutational landscape of PCCs to identify novel gene candidates involved in disease development. A discovery cohort of 15 PCCs wild type for mutations in PCC susceptibility genes underwent whole-exome sequencing, and an additional 83 PCCs served as a verification cohort for targeted sequencing of candidate mutations. A low rate of nonsilent single nucleotide variants (SNVs) was detected (6.1/sample). Somatic HRAS and EPAS1 mutations were observed in one case each, whereas the remaining 13 cases did not exhibit variants in established PCC genes. SNVs aggregated in apoptosis-related pathways, and mutations in COSMIC genes not previously reported in PCCs included ZAN, MITF, WDTC1, and CAMTA1. Two somatic mutations and one constitutional variant in the well-established cancer gene lysine (K)-specific methyltransferase 2D (KMT2D, MLL2) were discovered in one sample each, prompting KMT2D screening using focused exome-sequencing in the verification cohort. An additional 11 PCCs displayed KMT2D variants, of which two were recurrent. In total, missense KMT2D variants were found in 14 (11 somatic, two constitutional, one undetermined) of 99 PCCs (14%). Five cases displayed somatic mutations in the functional FYR/SET domains of KMT2D, constituting 36% of all KMT2D-mutated PCCs. KMT2D expression was upregulated in PCCs compared to normal adrenals, and KMT2D overexpression positively affected cell migration in a PCC cell line. We conclude that KMT2D represents a recurrently mutated gene with potential implication for PCC development.

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INTRODUCTION

Pheochromocytomas (PCCs) are rare, predominantly benign tumors arising from chromaffin cells of the adrenal medulla. Patients with PCC are diagnosed using catecholamine screening along with cross-sectional imaging and treated surgically (Welander et al., 2011; Brito et al., 2015). Approximately 40% of patients with PCCs have been reported to carry germline mutations in a growing list of genes (Dahia, 2014) currently including FH, EGLN1, EPAS1, KIF1Bβ, MAX, NF1, RET, SDHA, SDHB, SDHC, SDHD, SDHAF2, TMEM127, and VHL (Crossey et al., 1994; Ladroue et al., 2008; Schlüsis et al., 2008; Qin et al., 2010; Comino-Méndez et al., 2011; Welander et al., 2011; Zhuang et al., 2012; Castro-Vega et al., 2014; Dahia, 2014; Brito et al., 2015). Germline mutations in several of these susceptibility genes cause adrenomedullary tumor syndromes in which the patient presents with PCCs in addition to various syndromic manifestations (Favier et al., 2015). NF1 mutations cause neurofibromatosis type 1 in which 5% of patients can develop PCCs, RET mutations cause multiple endocrine neoplasia type 2, VHL mutations cause the von Hippel–Lindau syndrome, mutations in diverse SDHx genes have been linked to different hereditary paraganglioma and/or PCC syndromes named PGL1-4, mutations in EPAS1/HIF2A have been associated with the polycthemia-paraganglioma syndrome (Zhuang et al., 2012) and the Reed syndrome gene FH was recently found mutated in PCCs (Castro-Vega et al., 2014). Gene expression analyses have revealed that the PCCs can be clustered into mainly two different subgroups relating to genetic events and their aberrant pathways: VHL/SDHx/EPAS1-mutated tumors associate to stabilization of hypoxia inducible factors while KIF1Bβ/MAX/NF1/RET/TMEM127-mutated tumors correlate to the activation of kinase signalling pathways (Eisenhofer et al., 2004; Schlüsis et al., 2008; Dahia, 2014; Welander et al., 2014a; Favier et al., 2015). The genetics underlying sporadic PCCs are not yet clearly understood, and a majority of the tumors still lack a defined genetic driver event (Welander et al., 2011; Favier et al., 2015). Even so, somatic inactivating neurofibromin 1 (NF1) tumor suppressor gene mutations was recently discovered as a frequent event in PCCs (Burnichon et al., 2012; Welander et al., 2012) in addition to activating HRAS and EPAS1 mutations in subsets of cases (Comino-Méndez et al., 2013; Crona et al., 2013).

To further characterize the mutational landscape of adrenomedullary tumors and to identify novel gene candidates involved in disease development, 15 PCCs lacking established PCC susceptibility gene mutations as well as one RET-mutated PCC were collected and subjected to whole-exome sequencing (WES), and candidate genes were validated in an expanded cohort.

MATERIALS AND METHODS

Sample Acquisition, Preparations, and Ethical Statements

The study comprised a total of 99 PCCs; 89 Swedish cases and 10 US cases (Supporting Information Table 1). The PCCs are divided into a discovery cohort of 16 cases and a verification cohort of 83 cases. The discovery cohort subjected to WES consisted of fresh-frozen samples from 16 matched pairs of histologically confirmed PCC and normal tissues collected from surgery specimen at the Karolinska University Hospital, Stockholm, Sweden, and the verification cohort consisted of fresh-frozen tissues from a total of 83 histologically confirmed PCCs; 73 samples from the Karolinska University Hospital and 10 samples from the Yale-New Haven Hospital, New Haven, CT. All cases have been histologically confirmed as PCC using World Health Organization (WHO) criteria as part of the routine histopathology work-up (DeLellis et al., 2004). The distinction between malignant and benign PCCs was evaluated using the Armed Forces Institute of Pathology criteria (AFIP; Lack, 2007). All discovery cohort cases (n = 16) as well as approximately 70% of the cases in the verification cohort collected for extraction of genomic DNA were of sufficient sizes to allow for recutting and review by an experienced pathologist to confirm the representativity of PCC and eventual matched normal tissue for each case in parallel to the DNA extraction process (data not shown). Genomic DNA from somatic and constitutional tissues was extracted and validated using the DNeasy Blood and Tissue DNA isolation kit (Qiagen, Hilden, Germany) and Nano-Drop technology. DNA was obtained from both tumor and constitutional tissues for all cases in the discovery cohort and subsequently subjected to WES. As the exome sequencing was carried out in parallel for tumor and matched constitutional DNA for each case, recurrently mutated candidate genes on the somatic level were subsequently checked for germline variants using the exome sequencing data retrieved from constitutional tissues. The attaining of tissue and subsequent genomic analyses from both institutions were permitted by the local ethical
review board at Karolinska Institutet, Stockholm, Sweden and the Yale University Institutional Review Board, New Haven, CT, respectively.

Clinical and Genetic Characteristics

The overall clinical and genetic characteristics of the discovery and verification cohorts are detailed in Supporting Information Table 1. The discovery cohort of 16 PCCs used for WES includes 15 tumors (11 benign and 4 malignant PCCs) previously screened and found wild type for mutations in 13 known PCC susceptibility genes, namely EGLN1, KIF1Bβ, MAX, MEN1, NF1, RET, SDHA, SDHB, SDHC, SDHD, SDHAF2, TME1M27, and VHL. (Welander et al., 2014b). A PCC from a MEN2 patient exhibiting a constitutional RET gene mutation was also added to the discovery cohort, as this case displayed a highly equivocal pathology report suggestive of malignant features, but not fulfilling the AFIP or the WHO criteria for malignancy. As a very low PCC malignancy rate is reported for MEN2a patients, we analyzed this case as a part of the discovery cohort to pinpoint additional somatic driver mutations which could bear significance for malignant transformation in MEN2 PCCs. At the time of submission of the PCC discovery cohort for WES, the EPAS1 mutational status of the 16 cases was not known. Approximately 6 months after the submission of the discovery cohort material for WES, EPAS1 gene mutational status was included in the original study (Welander et al., 2014b).

Mutational data were available for 73 of the 83 PCCs included in the verification cohort, and the information was collected from three independent studies (genes investigated within parenthesis); (Welander et al., 2014a) (EPAS1, TME1M27), (Welander et al., 2012) (MAX, NF1, SDHB/D, RET, VHL) and (Welander et al., 2014b) (EGLN1, EPAS1, KIF1Bβ, MAX, MEN1, NF1, RET, SDHA/B/C/D, SDHAF2, TME1M27 and VHL). These data are detailed in Supporting Information Table 1.

Exome Capture, Massively Parallel Sequencing, Analysis, and Expression Studies

Genomic DNA samples generating adequate high-quality libraries were subjected to exome capture and sequencing, and the complete methodology regarding WES, sequence validation, ontology analyses, statistics, expression analyses, and functional experiments are detailed in the Supporting Information Materials and Methods.

RESULTS

WES of the Discovery Cohort

Using WES, a total of 130 somatic single nucleotide variants (SNVs) were detected across the 16 samples in the discovery cohort, with an individual tumor SNV count ranging from 0 to 18. Of the 130 SNVs detected, 97 (75%) were nonsynonymous, on average 6.1/sample (Fig. 1A, Supporting Information Table 2). Eighty-eight were missense alterations, five were positioned at exon–intron boundaries, and four were nonsense (truncating mutations). A mean coverage of 217 and 103 reads in tumor and normal samples, respectively, was obtained. Tumor samples were intentionally sequenced to a greater depth than normal tissue to maximize detection of heterozygous mutations in tumor cells intermingled with adjacent stromal tissue. Tumor purity ranged from 20.5 to 89.6% with a mean purity of 59%.

Somatic Mutations in PCC-Related Genes

Variants in PCC-related genes included the previously established constitutional RET variant in Case 34 as well as somatic mutations in HRAS (Q61R) and EPAS1 (P531S) in one case each (cases 88 and 94, respectively); however, the remaining 13 cases did not display any variant in PCC-associated genes. The HRAS Q61R mutation is a well-established activating variant that has been previously demonstrated in PCCs (Crona et al., 2013). The EPAS1 mutation P531S mutation has been similarly found in adrenomedullary tumors previously, and this variant affects the EPAS1 prolyl hydroxylation site (residues 530–539), which modulates the oncogenic function of EPAS1 (Comino-Méndez et al., 2013). While Case 88 lacks other somatic mutations in cancer-related genes, Case 94 also displays a heterozygous missense variant in the TSC2 tumor suppressor gene (Fig. 1A).

Recuously Mutated Catalogue of Somatic Mutations in Cancer (COSMIC) Database Genes

KMT2D, also known as mixed-lineage leukemia 2 (MILL2) encoding a histone methyltransferase that regulates DNA accessibility was identified as the most frequently mutated cancer-related gene in our discovery cohort, with two cases exhibiting somatic heterozygous KMT2D gene mutations and one additional case harboring a constitutional heterozygous KMT2D variant (3/16; 19%) (Fig. 1A, Table 1, Supporting Information Fig. 1). The two somatic mutations (p.N5223S

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and p.H5420Y) were both allocated to evolutionary-conserved amino acid positions within domains of the KMT2D protein with potential conserved roles related to chromatin regulation; namely the FYR-N and SET domains, respectively (Table 1, Fig. 2A). The constitutional variant (p.G2735S) is located in a region without known functional annotations, and is not reported as a polymorphism in the 1000 Genomes or Exome Variant Server databases. All three KMT2D variants as well as 13 additional variants from 12 genes detected by WES were confirmed using Sanger sequencing, and detailed information regarding these variants and primer sequences are available in Supporting Information Table 3.

Moreover, somatic missense mutations in the zonadhesin (ZAN) gene were found in two cases (Case 34; p.T823N and Case 88; p.A2396S). ZAN encodes a protein that is involved in sperm adhesion to the zona pellucida of the egg, and is represented in the COSMIC database as a recurrently mutated gene in lung adenocarcinoma and endometroid uterine carcinoma.

Somatic Mutations in Nonrecurrent COSMIC Genes

Several nonrecurrent somatic mutations in COSMIC annotated genes not previously associated to PCC development were observed in cases devoid of other known driver gene mutations (Fig. 1A, Supporting Information Table 2). For example, Case 19 (benign PCC) exhibited a missense p.R324S mutation in the microphthalmia-associated transcription factor (MITF) oncogene, a gene frequently amplified and mutated in melanoma (Cronin et al., 2009). Case 28 (malignant PCC) displayed a truncating p.L294X mutation in the WD and tetratricopeptide repeats 1 (WDTC1) gene, a candidate driver gene in microsatellite-unstable colorectal cancer (Alhopuro et al., 2012), and Case 37 (benign PCC) exhibited a
Figure 2. (A) Schematic representation of the *KMT2D* gene and mutational burden in PCCs. The 54 exon-spanning *KMT2D* gene is depicted with arrows indicating exon positions for each of the 14 *KMT2D* missense variants discovered in the discovery and verification cohorts. Recurrent variants (p.G2735S and p.N5223S) were found in two PCC cases each. Mutations within functionally important regions include N5222K/N5223S (FYR-N domain; amino acids 5175-5235), R5266H (FYR-C domain; amino acids 5236-5321), and H5420Y (SET domain; amino acids 5397-5513). (B) Search Tool for the Retrieval of Interacting Genes/Proteins (String) database interaction output illustrating well-characterized *KMT2D* (MLL2) interacting proteins. Blue lines denote confident binding partners as verified through experimental data and gray lines denote a predicted association in curated databases. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]
p.H154Y mutation in calmodulin binding transcription activator 1 (CAMTA1), a gene located on chromosome 1p36, a frequently deleted region in adrenergic medullary tumors (Edström Elder et al., 2002; Fig. 1A). Moreover, Case 34 displaying a constitutional RET gene mutation exhibits an equivocal pathology report suggestive of malignant features, but not fulfilling histological criteria for malignancy. As a very low malignancy rate is reported for PCCs in MEN2a patients, we included this case as a part of the discovery cohort to detect eventual additional somatic driver gene mutations of interest. Indeed, in Case 34, we observed a somatic p.D76G mutation in the cyclin-dependent kinase inhibitor 2C (CDKN2C, p18), a gene previously showed to be mutated in human RET-associated PCCs (van Veelen et al., 2009; Fig. 1A, Supporting Information Table 3).

**Mutational Gene Ontology Analysis**

Gene ontology analyses were performed using The Database for Annotation, Visualization and Integrated Discovery (DAVID) database. These analyses identified an enrichment of somatic mutations in apoptosis-related pathways, as nonsynonymous mutations in 10 different genes annotated as “apoptosis-related” were seen (ANXA1, BIRC6, CD5, CDKN2C, HMOX1, HRAS, MITF, NOX5, NTF3, and RYR2). This finding constituted the top enriched biological process, as these genes constitute approximately 10% of all mutated genes in the discovery cohort. Other significant gene ontology enrichments comprised chordate embryonic development (seven mutated genes; EPAS1, KMT2D/MLL2, PROX1, RIC8A, SFRP2, TSC2, WDTC1) and chemotaxis (four mutated genes; CXCL13, HRAS, PLD1, TSC2).

**Custom Amplicon KMT2D Sequencing**

A verification cohort of 83 additional PCCs was collected and analyzed for KMT2D gene mutations by focused sequencing of exomic regions and exon–intron boundaries using molecular inversion probes (Supporting Information Table 1). This lead to the discovery of 11 additional heterozygous missense variants in 83 PCCs (13%), all verified using Sanger sequencing. Ten of the KMT2D variants detected in PCCs were verified somatic based on the finding of wild-type sequences in DNA from constitutional tissues, whereas two variants were found also in

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**TABLE 1. Summary and Computational Functional Prediction of the 14 KMT2D Gene Variants in 99 Pheochromocytoma Samples**

| Sample number | Variant (amino acid change) | Nucleotide change | Somatic/constitutional origin | CHASM driver score | P value | CHASM functional score | P value | PolyPhen2 score | PolyPhen2 prediction | FYR/SET domain |
|---------------|-----------------------------|-------------------|-------------------------------|-------------------|--------|------------------------|--------|-----------------|----------------------|------------------|
| Discovery cohort (n=16) | | | | | | | | | | |
| 6 | G2735S | GCC>AGC | Constitutional | 0.562 | 0.254 | 0.291 | 0.318 | 0.008 | Benign | No |
| 36 | N5223S | AAT>AGT | Somatic | 0.420 | 0.063 | 0.174 | 0.481 | 0.887 | Possibly damaging | FYR-N |
| 92 | H5420Y | CAC>TAC | Somatic | 0.364 | 0.030 | 0.596 | 0.123 | 0.998 | |
| Verification cohort (n=83) | | | | | | | | | | |
| 3 | G2735S | GCC>AGC | N.d. | 0.562 | 0.254 | 0.291 | 0.318 | 0.008 | Benign | No |
| 12 | R5266H | CGC>CAC | Somatic | 0.504 | 0.155 | 0.471 | 0.184 | 0.98 | Probably damaging | FYR-C |
| 27 | Q1023K | CAG>AAG | Constitutional | 0.700 | 0.605 | 0.119 | 0.608 | 0.002 | Benign | No |
| 30 | N2965S | AAC>AGC | Somatic | 0.402 | 0.052 | 0.006 | 0.996 | 0.000 | Benign | No |
| 43 | N5223S | AAT>AGT | Somatic | 0.420 | 0.063 | 0.174 | 0.481 | 0.887 | Possibly damaging | FYR-N |
| 52 | L2610P | CTA>CCA | Somatic | 0.394 | 0.047 | 0.212 | 0.418 | 0.416 | Benign | No |
| 66 | P4048L | CCG>CTG | Somatic | 0.480 | 0.125 | 0.188 | 0.455 | 0.039 | Benign | No |
| 67 | L4222V | CTA>GTA | Somatic | 0.548 | 0.282 | 0.268 | 0.344 | 0.967 | Possibly damaging | FYR-N |
| 83 | R2922W | CGG>TGG | Somatic | 0.302 | 0.012 | 0.609 | 0.119 | 0.928 | Possibly damaging | |
| 87 | N5222K | AAC>AAA | Somatic | 0.704 | 0.616 | 0.235 | 0.386 | 0.446 | Benign | FYR-N |
| U100 | F2536S | TTC>TCC | Somatic | 0.530 | 0.196 | 0.76 | 0.059 | 0.535 | Benign | No |

*a* N.d. = not determined due to absent normal tissue.

*b* CHASM = Cancer-specific High-throughput Annotation of Somatic Mutations, driver score interpretation: true driver mutations close to 0, functional score: close to 1 means functional effect.

*c* PolyPhen2 = Polymorphism Phenotyping v2, non-synonymous missense variant prediction.
constitutional tissues and one was termed undetermined due to the lack of normal tissues (Table 1, Supporting Information Fig. 1, Supporting Information Table 3). In total, heterozygous missense KMT2D variants were found in 14 out of 99 PCCs sequenced (14%; 13 of 89 Swedish cases, 1 of 10 US cases), and two recurrent variants (p.G2735S and p.N5223S) were seen in two independent PCCs, respectively.

In Silico Mutation Prediction Analyses

To assess the pathogenic nature of the KMT2D mutations discovered in our cohorts, we assessed the individual variants using a bioinformatical screening process incorporating the established mutation prediction softwares cancer-specific high-throughput annotation of somatic mutations (CHASM) and PolyPhen 2 HumDiv (Carter et al., 2009; Adzhubei et al., 2010). The results indicate that subsets of the detected KMT2D mutations are predicted to be of functional significance as supported by these independent prediction softwares (Table 1). Moreover, a schematic overview of significant KMT2D binding partners was obtained through the Search Tool for the Retrieval of Interacting Genes/Proteins (String) database (Fig. 2B).

Constitutional KMT2D Variants

Two constitutional variants (p.G2735S in Case 6 and p.Q1023K in Case 27) were observed. Case 6 is a 44-year-old female with a benign PCC without signs of recurrent disease, also displaying history of an invasive ductal carcinoma of the breast. Case 27 is a 52-year-old female with a benign PCC, no recurrences and no additional tumors. Both patients lack positive family history indicative of tumor susceptibility syndromes, and no indications of Kabuki syndrome-related features were found. No parental samples were available for further genetic characterization. The same screening was carried out for a third patient (Case 3) with a KMT2D variant that could not be verified as somatic or constitutional due to lack of normal tissue.

Comparison to Established Genotypes and Clinical Parameters

Using the previously established genotypes in the Swedish PCC subset of the verification cohort (n = 73), a detailed mapping of case-by-case mutational burden regarding the 10 out of the 13 susceptibility genes that were found mutated in any of the PCCs (including KMT2D) was performed (Welander et al., 2012, 2014b; Fig. 1B). KMT2D was found mutated in 13 Swedish PCC cases, constituting 25% of the PCCs with any known susceptibility gene mutation (Fig. 1B). Seven of the 13 PCCs with KMT2D mutations did not display mutations in other PCC susceptibility genes (54%), whereas the remaining six cases also carried mutations in either NF1 (three cases), RET (two cases), or TMEM127 (one case) (Fig. 1B).

Tumors with KMT2D mutations were found to be significantly larger than tumors with other known PCC susceptibility gene mutations (Two-tailed Mann–Whitney U test, P = 0.039, Fig. 3A). Full biochemical data was available for 11 PCCs with KMT2D mutations, 10 of which displayed increased serum norepinephrine levels (Supporting Information Table 1). Although not statistically significant, this trend may indicate an underlying biochemical correlation between KMT2D mutations and serum norepinephrine. KMT2D mutations were not significantly associated with gender, age, or malignancy status (Supporting Information Table 1).

KMT2D Expressional and Copy Number Analyses

The copy number of KMT2D was determined in 86 PCCs using a TaqMan Copy Number Assay targeting the KMT2D locus. While the vast majority of samples (n = 78; 91%) were diploid for KMT2D, a small subset did exhibit one (n = 2; 2%), three (n = 5; 6%), or four (n = 1; 1%) copies (Supporting Information Table 4). All KMT2D-mutated cases available for copy number analyses were diploid for the KMT2D locus, and hence mutations and copy number alterations were mutually exclusive.

KMT2D expressional analyses using quantitative RT-PCR (qRT-PCR) were undertaken for PCCs for which RNA was available (n = 69) and for normal adrenal samples (n = 10) as well as normal adrenal medulla (n = 1). The KMT2D gene was found expressed in all PCCs tested, with a relative expression compared to normal adrenal mean ranging from 0.429 to 6.257 and the normal adrenomedullary biopsy exhibited KMT2D levels on par with the 10 normal whole-adrenals used as normalization controls (Fig. 3B, Supporting Information Table 4). Furthermore, KMT2D expression was significantly increased in PCCs compared to normal adrenals (Two-tailed Mann–Whitney U test, P = 0.017) but did not significantly differ between KMT2D-mutated and wild-type cases.

KMT2D immunohistochemistry was performed for all cases with available formalin-fixated and
paraffin-embedded (FFPE) tissues \( (n = 65) \). All PCCs either stained positive (>75% positive tumor nuclei, \( n = 28 \)) or partially positive (50–75% positive tumor nuclei, \( n = 37 \)) for nuclear KMT2D (Fig. 3C, Supporting Information Table 4). Statistically, KMT2D-mutated cases more frequently displayed positive KMT2D nuclear staining than wild-type cases (Fisher’s exact, \( P = 0.032 \)). Moreover, 19 tumors (10 KMT2D wild-type and 9 with somatic KMT2D mutations) were furthermore assessed for immunohistochemistry using an anti-H3K4me3 antibody, as KMT2D conveys trimethylation of histone-3 lysine-4 (Kim et al., 2014). Thirteen PCCs (of which 6 KMT2D mutated) displayed positive nuclear staining and the remaining six (including 3 KMT2D mutated) exhibited partially positive immunoreactivity (Fig. 3C, Supporting Information Table 4). A significant correlation between cases with positive KMT2D nuclear staining and cases with positive H3K4me3 nuclear staining was observed (Fisher’s exact, \( P = 0.038 \)).

A subset of PCCs (\( n = 18 \); eight KMT2D mutated and 10 KMT2D wild type) was assessed for H3K27me3, as activation of gene expression in the related tumor-type medulloblastoma is concerted through increased H3K4me3 and reduction of H3K27me3 levels, respectively (Dubuc et al., 2013). Sixteen cases either stained negative \( (n = 6) \) or mixed, displaying nuclear H3K27me3 immunoreactivity in 25–75% of tumor cells \( (n = 10) \). Interestingly, all six H3K27me3 negative cases exhibited strong H3K4me3 staining, and
four of them displayed somatic KMT2D mutations. 11 out of the 13 H3K4me3-positive PGCs (85%) exhibited either negative or mixed H3K27me3, only two were H3K27me3 positive. These results suggest that the majority of H3K4me3-positive cases seem to exhibit reduced levels of H3K27me3, and that H3K27me3-negative PGCs are associated to strong H3K4me3 levels as well as KMT2D mutations (Supporting Information Table 4).

**SDHB Expressional Analyses**

All PGCs (n = 16) in the discovery cohort and 73 of the 83 PGCs included in the verification cohort have been previously screened for PCC susceptibility gene mutations, (including SDHB), however, no SDHB mutations were found among the 89 PGCs tested (Supporting Information Table 1). To test whether KMT2D mutations could potentially affect the SDHB expressional status, we gathered available FFPE tissue from 14 PGCs (including five cases with KMT2D mutations) and 13 paragangliomas. The paragangliomas were included as negative and positive controls. SDHB immunohistochemistry was performed using a standardized protocol as detailed in the Supporting Information Materials and Methods section. All PGCs analyzed, except two cases endowed with SDHA mutations, displayed strong cytoplasmic SDHB immunoreactivity (Supporting Information Table 4). This is in line with the wild-type SDHB status for these cases. Also, the retained SDHB expression found in KMT2D-mutated PGCs suggest that SDHB expression is not disrupted through KMT2D mutations. Five of the 13 paragangliomas displayed SDHB mutations, and all SDHB-mutated cases displayed negative SDHB expression as expected, serving as negative controls for the experiments (data not shown).

**Functional KMT2D Experiments**

The functional outcome of KMT2D siRNA knockdown and KMT2D constitutive overexpression in the rat PGC cell line PC12-Adh was analyzed. While no significant effect on viability or loss of viability was seen upon KMT2D siRNA knockdown (Supporting Information Fig. 2), KMT2D silencing significantly reduced the motility of PC12-Adh cells (Fig. 4). A significant increase in cellular motility was similarly observed when constitutively overexpressing KMT2D in the same cell line, and the findings were furthermore supported by transient overexpression analyses (Fig. 4).

**Expressional Profiling Of KMT2D Overexpressed PC12 Cells**

The RNA expressional profile of PC12 cells with and without KMT2D overexpression was studied using a high-resolution Affymetrix array technique as described in detail in the Supporting Information Materials and Methods section. A significant difference in expressional patterns between mock-transfected and KMT2D-transfected PC12 cells was found for a total of 594 transcripts (P = 0.01; Supporting Information Table 5). Gene ontology analyses were undertaken using the KEGG pathway analysis at the DAVID database suggesting significant enrichment of genes within the Transforming growth factor beta (TGF-beta) signaling network and extracellular matrix–receptor interaction pathways (P = 6.7 × 10^-12 and 1.1 × 10^-10, respectively). Furthermore, in descending order of significance, the DAVID database suggests regulation of the five top five pathways when comparing PC12 cells with and without KMT2D overexpression (all with significant associations P < 5.3 × 10^-6): axon guidance, regulation of actin cytoskeleton, focal adhesion, and pathways in cancer. The finding of axon guidance and TGF-beta signaling pathways as significantly altered molecular networks in KMT2D-transfected PC12 cells compared to mock controls was furthermore supported by supplementary gene ontology analyses in which only transcripts with absolute fold changes of 1.5 or above were considered biologically relevant and included in the analysis (data not shown). Moreover, as detailed in Supporting Information Table 5, CDH2 and ITGBL1 (Cadherin-2 and Integrin beta-like 1 respectively) were two of the most down regulated genes.

**DISCUSSION**

In this study, WES was performed for a well-characterized PCC cohort devoid of established constitutional susceptibility gene mutations, identifying several interesting alterations, including the novel recognition of KMT2D and ZAN as recurrently mutated COSMIC genes. Furthermore, nonrecurrent mutations in the cancer-associated genes MITF, WDTC1, CAMTA1, and CDKN2C were detected in cases lacking other credible somatic driver events. Overall, somatic mutations

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aggregated in apoptotic-related pathways, which might indicate that aberrancies in signaling pathways controlling programmed cell death partly account for the development of PCCs.

Although adrenomedullary tumors have been previously investigated using next-generation sequencing, this is to our knowledge the first study that specifically targets a cohort enriched for genetic orphan PCCs to detect novel genes implicated in PCC development. We hypothesized that residual, nonestablished genetic events in PCCs potentially detectable through WES would probably occur at low frequencies. To ensure an adequate sensitivity for detecting these remaining causal variants, a large number of PCCs would be needed. Given the well-known rarity of adrenomedullary tumors, we selectively chose PCCs devoid of known PCC susceptibility gene mutations (somatic and constitutional) to counter this issue.

*KMT2D* was identified as the most recurrently mutated cancer gene in our discovery cohort, and extended investigations in a validation cohort revealed missense variants in 14% of the PCCs studied. Expressional *KMT2D* analyses indicate overexpression in PCCs compared to normal

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**Figure 4.** *KMT2D* affects the migratory potential of PC12-Adh cells. (A) PC12-Adh cells were treated with Lipofectamine 2000 (denoted Lipo), scrambled siRNA (denoted Neg), or specific siRNA (denoted siRNA) against *KMT2D* for 48 hr, trypsinized and allowed to migrate through a modified Boyden chamber for 2 hr; fixed, stained and the migrated cells were counted. Downregulation of *KMT2D* mRNA (B) and protein (C) were determined by qRT-PCR and Western blotting, respectively. (D) Constitutive *KMT2D* expression in PC12-Adh cells leads to an increase in migration through a modified Boyden chamber counted after 2 hr; PC12: untransfected control, PC-Neo: mock-transfected clone, and PC-MLL2: *KMT2D*-transfected clone. Upregulation of *KMT2D* mRNA was demonstrated by qRT-PCR (E).
tissues, and nuclear KMT2D expression was significantly associated to trimethylation of H3K4. The protein encoded by KMT2D is a histone methyltransferase that regulates DNA accessibility (Kim et al., 2014). Germline mutations in this gene have been shown to be a cause of Kabuki syndrome (OMIM #147920), a developmental disorder characterized by postnatal dwarfism, specific facial features as well as intellectual disability (Ng et al., 2010; Li et al., 2011; Paulussen et al., 2011). KMT2D is recurrently mutated in non-Hodgkin lymphoma (Morin et al., 2011; Okosun et al., 2014), and mutations have also been reported in epithelial tumors such as medulloblastoma, urinary bladder carcinoma, esophageal squamous cell carcinoma, and small cell lung cancer (Parsons et al., 2011; Balbás-Martínez et al., 2013; Dubuc et al., 2013; Lin et al., 2014; Ross et al., 2014; Song et al., 2014). Knockdown of KMT2D causes dysregulation of adhesion-related cytoskeletal events in vitro that in turn affect cell growth and survival, and KMT2D has been shown to exhibit tumor suppressor as well as oncogenic properties for various tumors (Issaeva et al., 2007; Natarajan et al., 2010; Guo et al., 2013; Kim et al., 2014). In this study, PCCs with KMT2D mutations were found to be significantly larger than tumors with other known PCC susceptibility gene mutations, suggesting that KMT2D mutations might positively influence tumor growth in PCCs. Overall, five PCCs displayed somatic mutations in the functional FYR or SET domains of KMT2D (Table 1), constituting 36% of all KMT2D-mutated PCCs in this study (Fig. 2A). Moreover, the finding of strong nuclear H3K4me3 levels in cases with KMT2D mutations vaguely suggests that the mutations affect the histone methyltransferase activity of KMT2D. However, as a significant proportion of the KMT2D mutations detected were not located within KMT2D domains with established methyltransferase-associated functions, it is possible that a subset of these mutations exert their function through disruption/increased affinity of important KMT2D binding partners, such as ESR1, KDM6A, and WDR5 to name a few (Fig. 2B). Notably, in silico analyses did not support a pathogenic role for a subset of the observed mutations, characterized by low PolyPhen2 or CHASM functional scores. Together with the fact that KMT2D expression did not significantly differ between KMT2D-mutated and wild-type cases, this could imply that subsets of the observed KMT2D mutations in fact are passenger events occurring randomly. As the size of KMT2D prevented a site-directed mutagenesis approach to study the functional consequences of the missense variants, the true pathogenic nature of the KMT2D somatic mutations discovered in this study remains to be established. Future studies will possibly elucidate whether KMT2D should be regarded as an oncogene or tumor suppressor in adrenomedullary tumors. Both patients (cases 3 and 27) exhibiting PCC and constitutional KMT2D variants lacked positive family history for either Kabuki syndrome as well as adrenomedullary disease. Also, the mutation prediction analyses did not point out the two
constitutional variants to be of pathogenic significance, and hence the importance of these findings remains unclear. Even so, cases 3 and 6 both exhibited the recurrent p.G273S variant, and both cases displayed ductal carcinoma in situ and invasive ductal carcinoma of the breast, respectively. Although Case 3 was not available for constitutional testing, the co-occurrence of PCC and malignant breast tumors in these two unrelated cases with an identical KMT2D variant is intriguing, and might suggest an underlying tumor phenotype.

To conclude, KMT2D is a recurrently mutated gene in PCCs. As more than half of the KMT2D-mutated tumors lack mutations in other known PCC susceptibility genes, KMT2D mutations could denote a novel genetic mechanism with possible implications for PCC tumorigenesis. The observation that KMT2D regulates adrenomedullary cell migration indicates that dysregulation of this intriguing methyltransferase might constitute a potential novel pathogenic mechanism for subsets of PCCs.

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