

RED CELLS, IRON, AND ERYTHROPOIESIS

Identification of a new *VHL* exon and complex splicing alterations in familial erythrocytosis or von Hippel-Lindau disease

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KEY POINTS

- Mutations in a *VHL* cryptic exon may be found in patients with familial erythrocytosis or VHL disease.
- Synonymous mutations in *VHL* exon 2 may induce exon skipping and cause familial erythrocytosis or VHL disease.

Chuvash polycythemia is an autosomal recessive form of erythrocytosis associated with a homozygous p.Arg200Trp mutation in the von Hippel-Lindau (*VHL*) gene. Since this discovery, additional *VHL* mutations have been identified in patients with congenital erythrocytosis, in a homozygous or compound-heterozygous state. *VHL* is a major tumor suppressor gene, mutations in which were first described in patients presenting with VHL disease, which is characterized by the development of highly vascularized tumors. Here, we identify a new *VHL* cryptic exon (termed E1') deep in intron 1 that is naturally expressed in many tissues. More importantly, we identify mutations in E1' in 7 families with erythrocytosis (1 homozygous case and 6 compound-heterozygous cases with a mutation in E1' in addition to a mutation in *VHL* coding sequences) and in 1 large family with typical VHL disease but without any alteration in the other *VHL* exons. In this study, we show that the mutations induced a dysregulation of *VHL* splicing with excessive retention of E1' and were associated with a downregulation of *VHL*

protein expression. In addition, we demonstrate a pathogenic role for synonymous mutations in *VHL* exon 2 that altered splicing through E2-skipping in 5 families with erythrocytosis or VHL disease. In all the studied cases, the mutations differentially affected splicing, correlating with phenotype severity. This study demonstrates that cryptic exon retention and

exon skipping are new *VHL* alterations and reveals a novel complex splicing regulation of the *VHL* gene. These findings open new avenues for diagnosis and research regarding the *VHL*-related hypoxia-signaling pathway. (*Blood*. 2018;132(5):469-483)

Introduction

Congenital erythrocytosis represents a heterogeneous group of rare disorders. Genetic changes affecting all parts of the regulatory pathway of erythropoiesis, including oxygen sensing, erythropoietin sensitivity, and hemoglobin oxygen affinity, have been described in patients with congenital erythrocytosis. The detection of underlying genetic changes in patients with presumed hematological pathology may have important implications for adequate clinical management. However, even with the use of next-generation sequencing panel diagnostics, the underlying genetic cause of presumed congenital erythrocytosis has been identified in fewer than one third of patients in most published cohorts.

The molecular basis of von Hippel-Lindau (*VHL*)-related congenital erythrocytosis was first described in the autonomous Russian Republic of Chuvashia, where this condition is an endemic disorder.¹ Chuvash polycythemia is frequently associated with rubor, vertebral hemangiomas, varicose veins, and low blood pressure. Chuvash patients have reduced survival rates associated with a higher prevalence of arterial and venous thromboses and pulmonary hypertension in addition to hemorrhagic events.²

Chuvash polycythemia arises from a homozygous c.598C>T p.Arg200Trp (R200W) mutation in the *VHL* gene. This specific *VHL* R200W mutation has also been identified in combination with other *VHL* mutations (compound heterozygosity), and other missense *VHL* mutations in both alleles have been described in patients with congenital erythrocytosis.^{3,4} Interestingly, it has been described in some unexplained cases of patients with erythrocytosis carrying only one heterozygous mutation.^{4,6}

VHL is located on 3p25-26 and has been reported to contain 3 exons (E1, E2, and E3). The commonly described *VHL* transcript contains the 3 spliced exons that encode a 213 amino acid (aa) protein (pVHL213; also termed pVHL30) and a smaller isoform (pVHL160 or pVHL19) initiated from an in-frame internal translation start site.⁷ A naturally occurring splice variant, expressed at low levels in some tissues, comprises E1 directly spliced to E3 and is translated into a protein product termed pVHL172 (pVHLΔE2), the functions of which are still under investigation.⁸⁻¹² pVHL213 and pVHL160 are involved in a variety of functions, the most studied being the regulation of the cellular oxygen-sensing pathway. The main player of this pathway is the hypoxia inducible factor (HIF). Under normal oxygen supply, the α subunits of HIF (HIF-1 α , -2 α , and -3 α) are hydroxylated by the prolyl-4 hydroxylase domain enzymes (PHD1, -2, and -3) and subsequently targeted by pVHL, a subunit of an E3 ubiquitin-ligase complex that promotes HIF- α ubiquitination and subsequent proteasomal degradation.^{13,14} Under hypoxic conditions or when *VHL* is mutated, HIF- α remains stable and heterodimerizes with HIF- β , constituting a functional HIF. HIF transcriptionally activates a variety of genes involved in adaptation to reduced oxygen supply (eg, erythropoiesis, angiogenesis, metabolism, and

cell survival). Dysregulation of the hypoxia pathway^{13,14} is central to the development of erythrocytosis^{1,4} (via upregulation of erythropoietin, a HIF-2 α target gene), but also in the development of tumors.¹⁵ Indeed, *VHL* is a tumor suppressor gene, heterozygous mutations of which are associated with *VHL* disease (Figure 1A).^{16,17} *VHL* disease, described in 1936, is an autosomal dominant disorder with high penetrance characterized by the development of highly vascularized tumors like central nervous system and retinal hemangioblastomas, pancreatic neuroendocrine tumors, pheochromocytomas, and clear cell renal cell carcinomas.¹⁸⁻²⁰

In patients carrying *VHL* mutations, the precise mechanistic aspects that underpin the different phenotypes remain obscure. Although most patients carry mutations in the *VHL* gene that induce a partial or complete loss of protein function, some cases remain unsolved. Indeed, some patients with erythrocytosis have been found to be heterozygous rather than homozygous for the expected alteration^{4,6} or carry homozygous synonymous mutations that leave the amino acid sequence intact. In addition, some patients present with *VHL* disease in the absence of identified mutations or deletions in *VHL* or carry heterozygous synonymous mutations. Here, we report an investigation of 12 families linked to unexplained disease, including 9 families with erythrocytosis and 3 with *VHL* disease (Figure 1A). This study led to the discovery of a novel cryptic exon in the *VHL* gene and complex regulation of *VHL* splicing.

Methods

Complete materials and methods are detailed in the supplemental Material and methods (available on the *Blood* Web site).

Study approval

Informed consent for medical diagnosis and research was obtained from the patients and their relatives. This study was agreed by the Comité Consultatif de Protection des Personnes dans la Recherche Biomédicale (French ethical committee) Paris-Sud at Bicêtre Hospital.

Sanger sequencing

Exons and exon-intron junctions of the *VHL* gene were sequenced from DNA extracted from whole blood, as previously described.²¹

WGS

Whole-genome sequencing (WGS; for families F2, F3, and F7) was performed at the clinically accredited Molecular Diagnostics Laboratory at the John Radcliffe Hospital using the Hi-Seq4000 platform (Illumina Inc., San Diego, CA) in high-throughput mode.^{22,23} Analysis of single-nucleotide variants, short insertions/deletions, and copy number variants was conducted and is explained in detail in the supplemental Methods.

Transcript detection and quantification

After reverse-transcription reactions (ThermoScientific), exon-specific polymerase chain reaction (PCR) was performed using

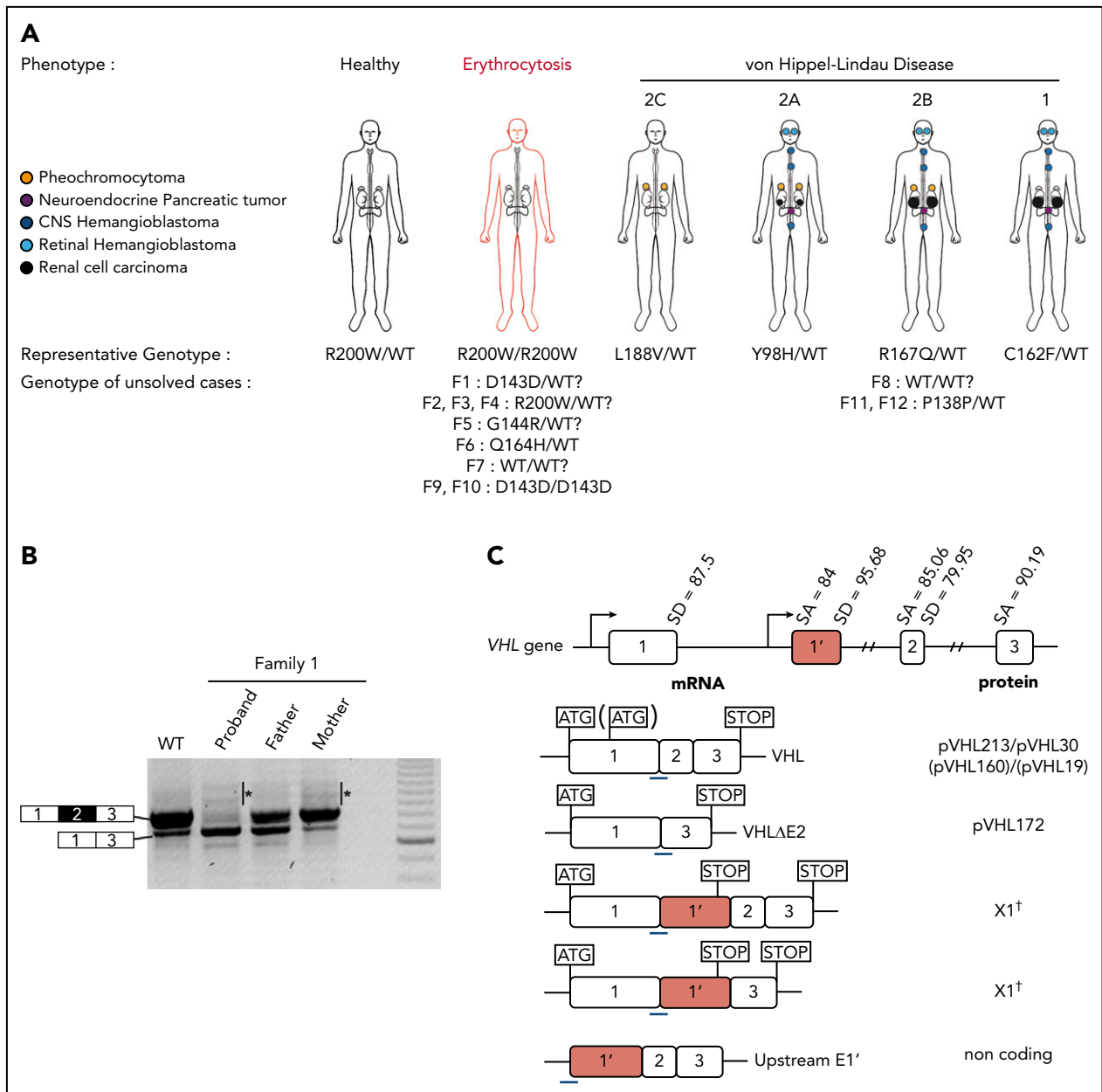


Figure 1. Clinical manifestations of patients carrying *VHL* mutations and identification of a new *VHL* spliced isoform containing a cryptic exon. (A) Mutations in the *VHL* gene predispose to different phenotypes. *VHL* disease is characterized by the development of central nervous system (CNS) and retinal hemangioblastomas, neuroendocrine pancreatic tumors, pheochromocytomas, and clear cell renal cell carcinomas. Chuvash polycythemia (erythrocytosis) is characterized by elevated red blood cell numbers. This study describes families with typical *VHL*-related phenotypes associated with an unexpected *VHL* status (ie, either synonymous mutations or no alterations in *VHL*). (B) Reverse transcription polymerase chain reaction (RT-PCR) using primers specific for E1 and E3 was performed on messenger RNA (mRNA) extracted from lymphoblastoid cell lines (LCLs) established from controls and patients of family 1 (F1). (C) Schematic representation of the *VHL* gene and its products. The different *VHL* exons are represented on a scale: E1, 340 bp from the ATG initiation codon; E1', 259 bp; E2, 123 bp; E3, 179 bp to the Stop termination codon. The full-length *VHL* mRNA isoform encodes pVHL213 (also named pVHL30). *VHL* E1 contains an internal translation initiation codon that initiates the production of pVHL160 protein (pVHL19). The isoform containing E1 spliced in phase with E3 encodes pVHL172 that lacks E2. The isoforms containing E1 spliced with E1' may theoretically encode a protein termed X1 of 193 aa (114 aa encoded by E1, and 79 aa encoded by E1'). Consensus values of donor (SD) and acceptor (SA) splice site sequences are indicated above the *VHL* gene, as calculated by the human splicing finder in silico tool. Horizontal blue lines indicate the location of probes used in TaqMan assays. *Denotes larger fragments that were subsequently cloned and sequenced. †The isoforms containing E1' spliced with *VHL* exons were identified by cloning and sequencing in the laboratory but were described later by the National Center for Biotechnology Information as transcripts able to produce a protein termed X1. WT, wild type.

primers localized to flanking E1 and E3 exons. TaqMan real-time PCR were performed on 20 ng of complementary DNA (cDNA) with the quantitative PCR Mastermix (Eurogentec). Quantification of *RPLPO* transcripts was used as internal control. The thresholds were determined using dilutions of plasmids containing coding sequences of each gene.

RNA sequencing

Library construction was performed with SureSelect Strand-Specific RNA Library Prep for the Illumina Multiplexed kit (Agilent Technologies). After purification (Macherey-Nagel), the fragment size of libraries was controlled using the 2200 TapeStation system (Agilent Technologies). Ten pM of each

Table 1. Description of variants identified in VHL E1' and VHL E2 with associated clinical manifestations

Family no., patient	Nucleotide variant on allele 1/allele 2	Impact on protein	Year of birth/age at diagnosis	Sex	Hb, g/dL	Ht, %	Red cells, $\times 10^9/L$	EPO, mU/mL	Phenotype	Other mutation
F1, II.1	c.429C>T/c.340+770T>C	VHL p.Asp143Asp/X1 p.Ser179Pro?	1991/15 y	M	16.6	57	6.9	163	Erythrocytosis, kidney ischemic infarct	—
F2, II.1	c.598C>T/c.340+770T>C	VHL p.Arg200Trp/X1 p.Ser179Pro?	1957/21 y	M	19.9	67	—	22.2	Erythrocytosis	—
F3, II.1	c.598C>T/c.340+770T>C	VHL p.Arg200Trp/X1 p.Ser179Pro?	2003/4 mo	M	17.9	54	7.5	60	Erythrocytosis	—
F3, II.2	c.598C>T/c.340+770T>C	VHL p.Arg200Trp/X1 p.Ser179Pro?	2005/6 mo	M	14.6	45	—	—	Erythrocytosis	—
F4, II.1	c.598C>T/c.340+694_711dup	VHL p.Arg200Trp/X1 p.Trp159X?	1990/7 y	M	20.6	64	8.25	49.9	Erythrocytosis, deep vein thrombosis, intracerebral hemorrhage	FV Leiden
F5, II.2	c.430G>A/c.340+694_711dup	VHL p.Gly144Arg/X1 p.Trp159X?	1975/13 y	M	16.2	60	5.9	—	Erythrocytosis	—
F6, II.1	c.492G>T/c.340+574A>T*	VHL p.Gln164His/SA	2014/6 mo	M	15.6	49	6.9	1167	Splenomegaly	—
F7	c.340+816A>C/c.340+816A>C	X1 p.*194Serext*24/X1 p.*194Serext*24	7 y	M	20	64	8.09	33	Erythrocytosis	—
F8	c.340+617C>G + c.340+648T>C†/WT	X1 p.Leu128Val + X1 p.Leu138Pro/WT	—	—	N	N	N	N	VHL disease	—
F9, II.1	c.429C>T/c.429C>T	VHL p.Asp143Asp/VHL p.Asp143Asp	1997/5 mo	M	16.2	49.3	5.2	186	Erythrocytosis	Hb sitia
F10, II.1	c.429C>T/c.429C>T	VHL p.Asp143Asp/VHL p.Asp143Asp	2002/4.5 y	F	22.5	64-77.4	12.6	264	Erythrocytosis, splenomegaly	β thalassemia
F11	c.414A>G/WT	VHL p.Pro138Pro/WT	—	—	N	N	N	N	VHL disease	—
F12	c.414A>G/WT	VHL p.Pro138Pro/WT	—	—	N	N	N	N	VHL disease	—

The second column indicates the position of the nucleotide variants identified in the VHL gene regarding the current nomenclature (sequence encoded by the VHL E1-E2-E3). The third column indicates the impact either on the VHL protein (for variants located in E1, E2, or E3) or on the potential X1 protein, if encoded by E1-E1' (for variants located in E1'). Normal values correspond to: Hb: M, 13 to 18 g/dL and F, 12 to 15 g/dL; Ht: M, 40% to 52% and F, 37% to 47%; red cells: M, 4.2 to 5.7 $\times 10^9/L$ and F, 4.2 to 5.2 $\times 10^9/L$; EPO: 5 to 25 mU/mL.

EPO, erythropoietin; Hb, hemoglobin; Ht, hematocrit; N, normal value.

*The nucleotide change was reported as rs982745672 with a global minor allele frequency that corresponds to T = 0.00007/2 (TOPMed).

†The nucleotide change was reported as rs73024533 with a global minor allele frequency that corresponds to C = 0.0026/13 (1000 Genomes) and C = 0.0050/147 (TOPMed).

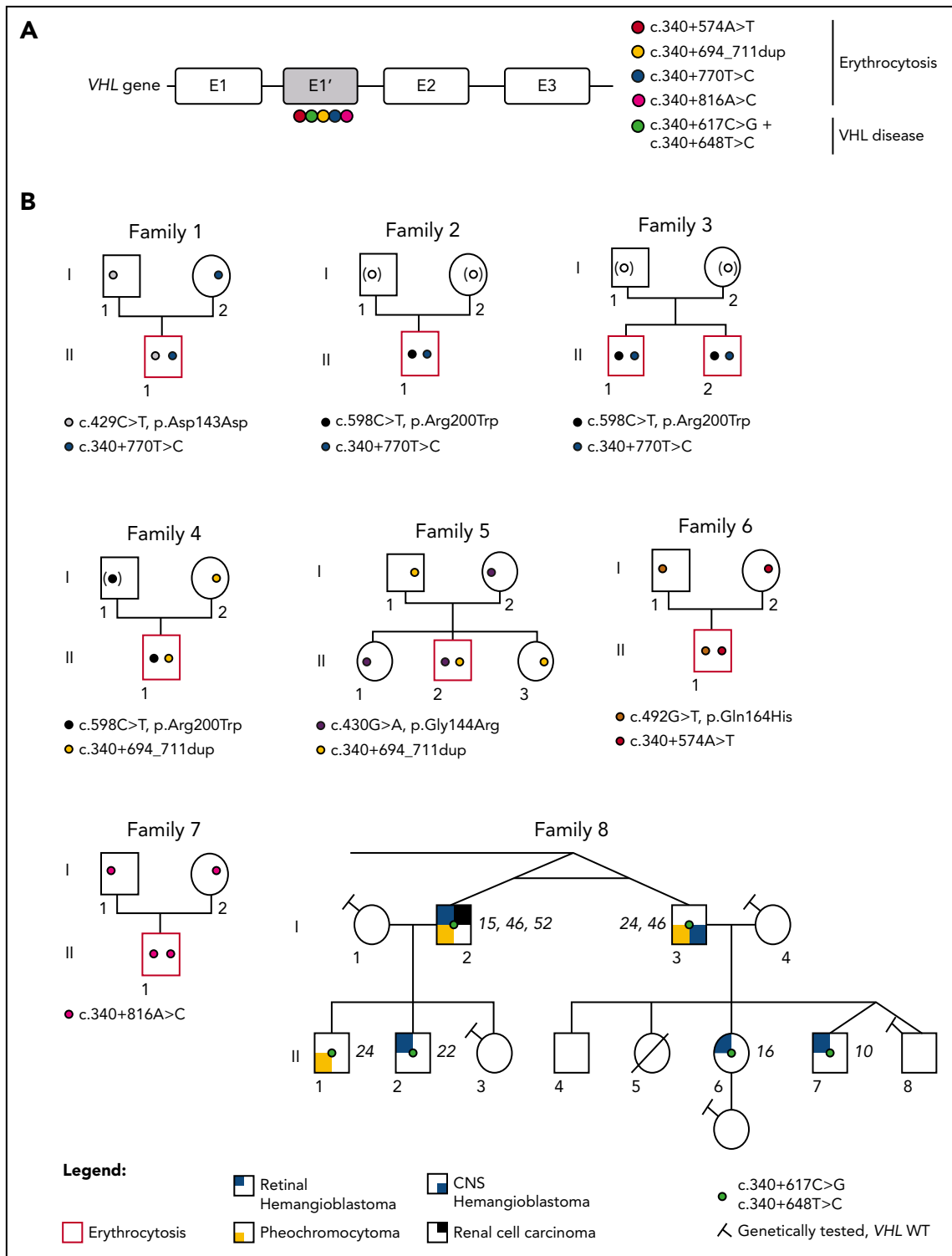


Figure 2. Identification of mutations in the new VHL cryptic exon in 7 families with erythrocytosis and a large family with VHL disease. (A) Schematic representation of the VHL gene and location of the identified mutations in the new VHL cryptic exon E1'. (B) Pedigree of families with erythrocytosis or VHL disease. The genotypes were elucidated by sequencing both parents and proband (F1, F5, and F6), deduced by sequencing of 1 parent and proband (F4, the mutation deduced in brackets), or deduced from allele cloning of proband carrying the conserved Chuvash mutation and core haplotype (F2 and F3), confirming the transmission of the mutations by 1 of each parent (for F2 and F3, the identity of the transmitting parent being unknown, the mutation is represented by a white circle in brackets). The genotype of parents from F7 was elucidated from WGS data. The numbers in italics (F8) indicate the age of the patient at tumor diagnosis. CNS, central nervous system.

library was pooled and prepared according to the denaturing and diluting library protocol for HiSeq and GAllx (Illumina) for cluster generation on the cBot system. Paired-end sequencing

was carried out in a single lane on the HiSeq 2500 system (Illumina). Processing of reads is detailed in the supplemental Methods.

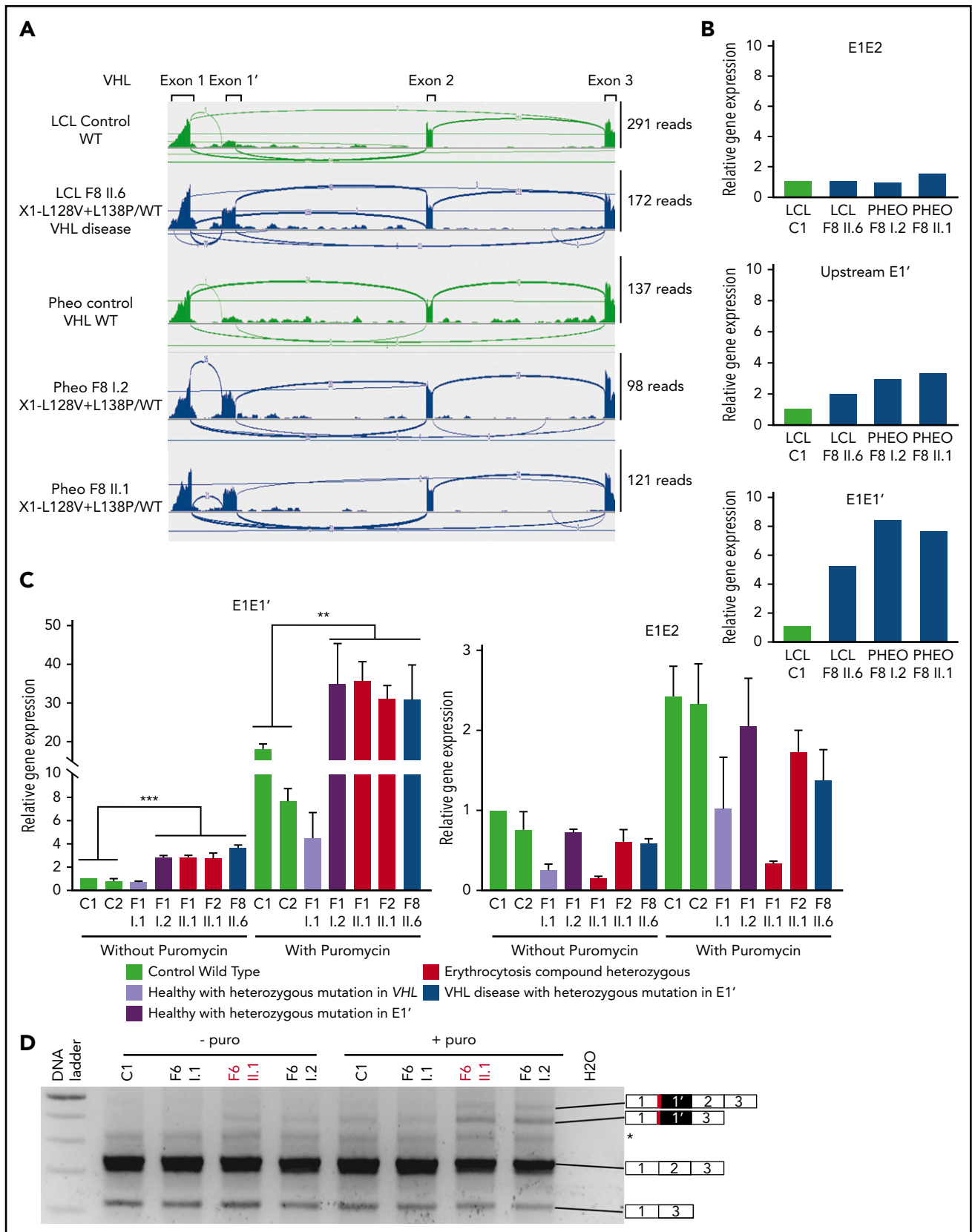


Figure 3. Expression study of the new VHL transcript isoforms in patient cells. (A) Sashimi plots from RNA-seq data obtained from samples (LCLs or pheochromocytomas [Pheo]) of 3 patients from F8. The positions of the different VHL exons are indicated, with the maximal number of reads indicated at the right. Splice junctions are denoted by the horizontal links, with details provided in supplemental Figure 8. (B) TaqMan quantification of the different VHL isoforms from samples of F8 performed using probes specific to VHL E1-E2, the translated sequence upstream E1', or E1-E1' junctions. Relative gene expression was normalized to LCL control (C1) fixed at 1 (mean results of technical duplicates). (C) TaqMan quantification of the different VHL isoforms in LCLs (established from 2 independent controls and from patients of F1, F2, and F8) cultured in the absence

Reporter assays

The plasmid encoding the WT hemagglutinin-tagged VHL was kindly provided by William G. Kaelin Jr. An expression plasmid for the hypothetical X1 protein was constructed by subcloning the coding sequence after its synthesis (Life Technologies). Mutations of interest were introduced by site-directed mutagenesis (New England Biolabs); 786-O cells were transfected (Ozyme) with constructs encoding WT or mutant proteins, firefly luciferase under the control of hypoxia response elements, and *Renilla* luciferase for normalization.²⁴ After 24 hours, luciferase assays were performed using the Dual-Luciferase Reporter Assay System (Promega).

Western blotting

Cell lysates from the luciferase reporter assay were loaded into a Bis-Tris Mini Gel (4% to 12%; Invitrogen). After transfer, the membrane (GE Healthcare) was subsequently incubated with a mouse anti-hemagglutinin antibody (BioLegend) and then a goat anti-mouse horseradish peroxidase-conjugated antibody (Jackson Immuno Research). Western blot using the mouse monoclonal antibody JD-1956 (patent #14305925.1-1402-2014; Centre National de la Recherche Scientifique Etablissement Français du Sang) raised against human VHL was performed as described.⁸

Minigene experiments

Minigene constructs were prepared in pCas2 plasmid, containing 2 artificial exons A and B (as described by Gaildrat et al²⁵), between which VHL E1' or VHL E2 with intronic flanking sequences were cloned. Cells were transfected (Ozyme) or nucleofected (Lonza). RNA was extracted 24 hours after transfection and reverse transcribed. PCR amplification was performed using the PCR GoTaqQ2 kit (Promega) with primers against artificial (A and B) exons. PCR products were resolved in a 2% agarose gel.

Results

Identification of new VHL spliced isoforms containing a cryptic exon

We first focused our study on a patient with erythrocytosis in whom a synonymous VHL c.429C>T p.Asp143Asp (D143D) mutation in the heterozygous state had previously been identified (F1; Table 1). No other mutations in the 3 VHL canonical exons had been identified in genomic DNA of this patient. RT-PCR using primers in E1 and E3 was performed using mRNA samples extracted from LCLs established from different family members. The results showed a strong decrease of the E1E2E3 isoform and an upregulation of the E1E3 isoform (Figure 1B) compared with WT LCLs. Some minor extra fragments of larger size were observed in this patient and his mother's sample. Subcloning and sequencing of these fragments allowed us to identify new VHL transcripts that contained intronic sequences. This intronic sequence, which we termed the E1' cryptic exon, was spliced to E1 at its 5' end and to either E2 or E3 at its 3' terminus (Figure 1C). We show that these isoforms were expressed in a variety of tissues and cell lines (supplemental Figure 1). Their translation may theoretically lead to

the production of a protein of 193 aa that contains the first 114 aa encoded by E1²⁶ and additional 79 aa of unknown function encoded by E1'. During the course of our study, data from an automated computational analysis for an isoform containing the E1' cryptic exon (E1E1'E2 isoform) were deposited in the National Center for Biotechnology Information. This isoform was predicted to encode a protein of 193 aa, named X1 (XP_011532380.1), with the same sequence (114 aa from E1 in addition to 79 aa encoded by E1') (Figure 1C). The analysis of this region revealed strong conservation in primates, but a moderate to low conservation in more distant species (supplemental Figure 2A-C); notably, the splice sites were identical to canonical sites (ttcag/TC, AG/gtaag) and were highly conserved. In silico analysis of the donor and acceptor splice sites of E1' showed similar consensus values compared with other VHL exons (Figure 1C, upper panel). The capacity to translate a potential X1 protein was only conserved in higher primates (supplemental Figure 2D). The deposited sequence was removed and replaced with a noncoding isoform containing E1' spliced with VHL E2 and E3 (ENST00000477538.1; Figure 1C, isoform on the bottom). This isoform may be initiated by an alternative promoter. Indeed, the sequence located at the 5' end of E1' represents a transcriptionally active region, as illustrated by epigenetic marks (supplemental Figure 3). We confirmed the expression of an additional transcript initiated from the upstream region of E1' (that we termed upstream E1') in different tissues and cell lines (supplemental Figure 1).

New E1' cryptic exon is mutated in patients with erythrocytosis or VHL disease

Sanger sequencing of this new cryptic exon in the proband (F1-II.1) identified a variant that had not been reported in databases: c.340+770T>C (Figure 2B; Table 1; supplemental Figure 4). Sequencing of the germline DNA of the mother revealed the same variant, indicating that the proband F1-II.1 was compound heterozygous. This result prompted us to sequence additional patients with erythrocytosis described as heterozygous for VHL mutations.

We investigated patients with Chuvash polycythemia for whom VHL R200W had been found in a heterozygous state. Sanger sequencing identified the identical intronic variant c.340+770T>C in a singleton (F2)⁵ and 2 affected brothers (F3; Figure 2B). In addition, we identified a duplication c.340+694_711dup in the proband of F4 and F5⁶ previously described as VHL R200W/WT and VHL G144R/WT, respectively (Figure 2B). In F6, previously diagnosed as VHL Q164H/WT, we identified a genetic variant c.340+574A>T that altered the consensus SA site ag/TC of E1'. This variant is described as a rare polymorphism (rs98274567) in the National Center for Biotechnology Information database (Table 1).

Because biological material from the parents of F2 and F3 was not available, we cloned the intronic region containing the E1' variant and the Chuvash core haplotype single-nucleotide polymorphism rs779808 associated with the VHL R200W mutation (described²⁷ to be located downstream of E1', in position

Figure 3 (continued) or presence of puromycin (puro), an inhibitor of nonsense-mediated mRNA decay. TaqMan probes were specific to the VHL E1-E1' or E1-E2 junction. The graph resumes experiments performed on LCLs cultured independently 3 times and quantified in duplicate. Data are represented as the mean \pm standard error of the mean. (D) RT-PCR using primers specific for E1 and E3 was performed on mRNA extracted from LCLs established from controls and patients of F6 (carrying the mutation c.340+574A>T that targets the SA site of E1'). Patient with erythrocytosis is indicated in red. On the right, the spliced isoforms are schematically represented. ** $P < .05$, *** $P < .01$ based on Student t test. *Denotes larger fragments that contained E1' spliced with E1, but with 15 nucleotides deleted (represented in red) by the use of an alternative SA site (sequences of the cloned bands are detailed in supplemental Figure 10).

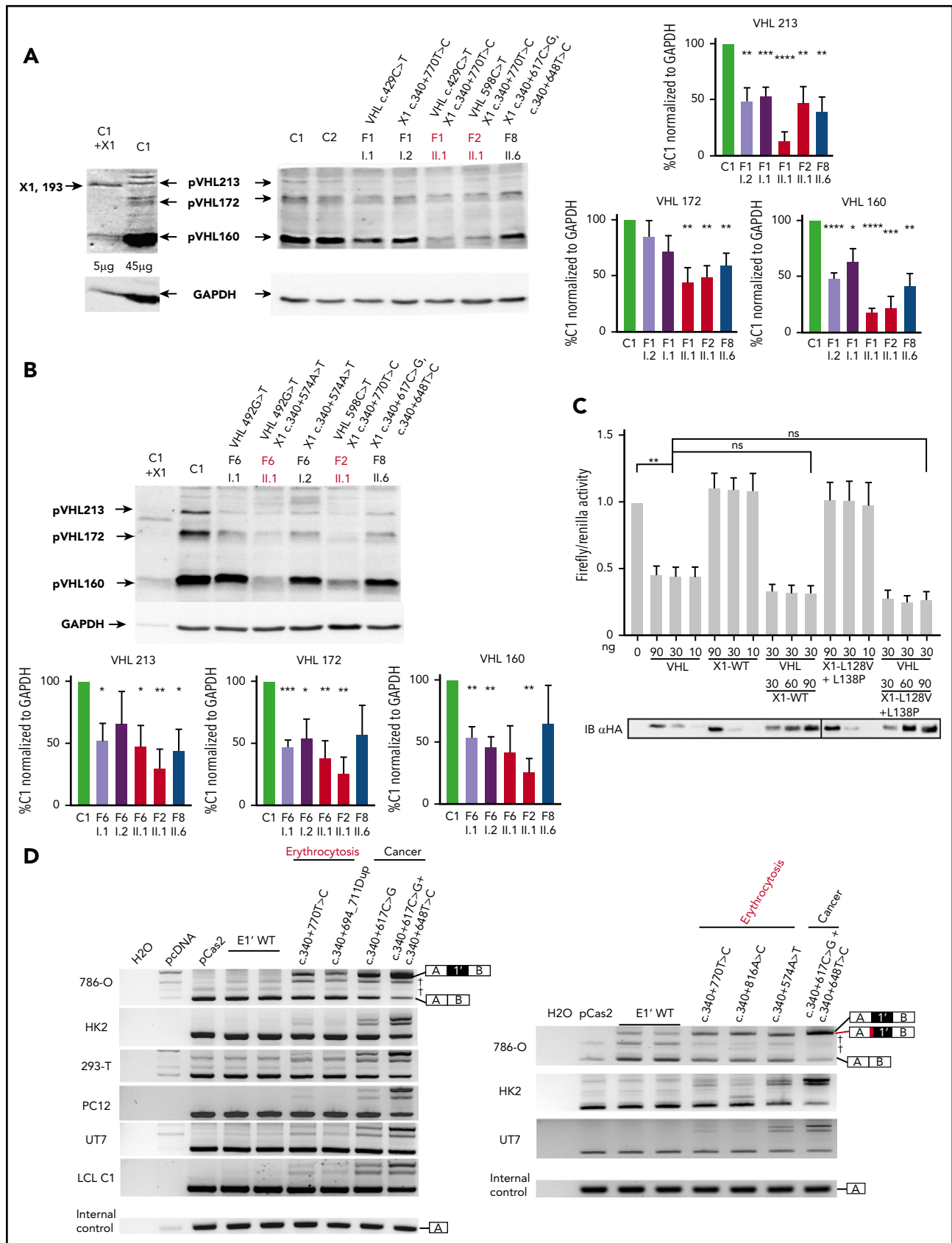


Figure 4. Functional studies of mutations in VHL E1'. (A-B) Immunoblot analysis of patient LCLs. A representative immunoblot (A) and quantification of 3 different immunoblots (B) are presented. Relative gene expression was normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) expression, and results obtained with LCL control (C1) were fixed at 100%. Data are represented as the mean \pm standard error of the mean. A VHL antibody that recognized VHL E1 was able to detect the hypothetical

c.340+1150). Segregation analysis (F4, F5, and F6) or Chuvash core haplotype analysis (F2 and F3) demonstrated that all patients were compound heterozygous, with 1 mutation being inherited from each parent (supplemental Figure 5).

In a parallel independent study, WGS was used to investigate patients with Chuvash polycythemia, heterozygous for the *VHL* R200W mutation (F2 and F3). The intronic variant c.340+770T>C was the only rare variant identified in the *VHL* gene in these patients. Other WGS filtering strategies (supplemental Methods) did not identify any additional significant mutations in biologically relevant genes (supplemental Figure 6). This study also conducted WGS for a trio with congenital erythrocytosis for which prior whole-exome sequencing had not identified any mutations (F7). No rare biologically relevant variants were identified using the filtering strategy described in supplemental Methods. However, further inspection of the new cryptic exon led to the identification of a c.340+816A>C variant in the homozygous state in the proband, with both parents being heterozygous for this mutation.

In addition to these patients with erythrocytosis, in which no variants in *VHL* were initially detected, screening of patients with VHL disease may also fail to detect mutations in canonical *VHL* exons. An example of such a family (F8) with hereditary hemangioblastoma, clear cell renal cell carcinoma, and pheochromocytoma was studied. Microsatellite analysis demonstrated a cosegregation of markers surrounding the *VHL* region with the disease (supplemental Figure 7). Sequencing of the new E1' cryptic exon in this family identified 2 heterozygous variants in E1': a previously unreported c.340+617C>G variant and a c.340+648T>C variant described as a rare polymorphism in databases (Table 1). These variants segregated in 6 patients who developed the disease and were absent in 4 healthy tested descendants, indicating their presence on a single disease-associated allele (Figure 2B). Sequencing of tumor DNA did not display loss of heterozygosity, suggesting that a WT *VHL* deletion (as observed in classic VHL disease) may not be a prerequisite in cells of patients with this specific *VHL* genotype (supplemental Figure 4).

Expression study of the new *VHL* isoforms in patients' cells

RNA sequencing (RNA-seq) analysis of patients' LCLs and tumors demonstrated that genomic variants in E1' were associated with an upregulation of transcripts containing the E1' cryptic exon compared with controls (Figure 3A; supplemental Figure 8). In addition, the mutated allele was preferentially expressed (>70%), suggesting a causal relationship between the genetic variants and E1' retention (supplemental Figure 8C).

We performed quantitative RT-PCR using TaqMan probes specific for the different splicing junctions or for the region upstream of E1' (location of probes shown in Figure 1C), which showed a strong upregulation of a unique isoform containing E1' spliced with E1 in LCLs and tumors from patients of F8 compared with controls (Figure 3B). The study therefore focused on the isoform E1-E1'. Transcripts that retained E1' spliced with E1 contained (in-frame with E1) a premature termination codon (Figure 1C) and could likely be targeted for degradation according to nonsense-mediated mRNA decay (NMD) mechanisms. We therefore investigated the expression of transcripts that contained E1 spliced with E1' in the absence or presence of puromycin, an inhibitor of NMD. Without puromycin, higher levels of expression were seen in samples with mutated E1' (Figure 3C) compared with samples containing WT E1'. Exposure to puromycin resulted in a profound induction of isoforms containing E1 spliced with E1' vs other isoforms (Figure 3C; supplemental Figure 9). These results indicated that isoforms with E1-E1' junction were degraded by NMD and may have failed to produce proteins. We performed RT-PCR on LCL samples of F6 carrying the mutation in the SA site, using primers in *VHL* exons E1 and E3. We observed fragments of larger size in samples from patients carrying the mutation, which were highly visible in the presence of puromycin (Figure 3D). Cloning and sequencing of these fragments demonstrated a dysregulated splicing of E1' with the use of an alternative SA site located 15 nucleotides downstream (supplemental Figure 10).

To measure the impact of the splicing dysregulation on pVHL expression, we performed an immunoblot with an antibody able to recognize the amino acids encoded by E1 and therefore able to detect the different pVHL isoforms.⁸ The antibody detected overexpressed exogenous X1 protein after transfection of the plasmid encoding X1 (Figure 4A). However, it failed to detect the endogenous protein, even in patients' samples that overexpressed the mRNA containing E1-E1'. Instead, the immunoblots showed a lower expression of all the VHL protein isoforms in patients with mutated E1' (Figures 4A-B).

Functional characterization of the mutated *VHL* E1'

Because the hypoxia pathway represents the major pathway involved in the genesis of secondary erythrocytosis and VHL disease,^{13-15,28} we performed functional studies of the hypothetical WT or mutated X1 proteins using hypoxia response element-dependent reporter assays. These functional tests failed to reveal any substantial effects of X1 on this pathway, either alone or in competition with pVHL (Figure 4C; supplemental Figure 11). We next focused our study on a potential impact of the E1' variants on splicing by performing splicing

Figure 4 (continued) X1 protein. In the most left lane, control LCLs were transfected with an expression vector containing the coding sequence for X1; 5 μg of proteins were blotted vs 45 μg for other samples to avoid signal saturation with overexpressed X1. Patients with erythrocytosis are indicated in red. (C) Functional hypoxia response element (HRE)-dependent reporter assays were performed in 786-O cells (ie, *VHL*⁻ cells that constitutively express HIF-2α). The results are expressed as relative firefly luciferase activity with *Renilla* luciferase as an internal control; 1.0 unit denotes the basal activity of endogenous HIF-2α using the HRE luciferase reporter plasmid. The ability of WT and mutated X1 to downregulate firefly luciferase activity (related to HIF activity) was compared with pVHL and in competition with pVHL. An immunoblot using an antibody specific to the hemagglutinin tag (HA) was used to detect HA-VHL and HA-X1. X1-L128V[†]L138P corresponds to a potential impact of the c.340+648T>C and c.340+617C>G mutations on the hypothetical X1 protein. Three independent experiments were performed. (D) Characterization of *VHL* E1' retention by the minigene experiment (representative picture of agarose gel; n = 3). RT-PCR was performed on mRNA obtained from cell lines transfected with a minigene construct containing *VHL* E1' (WT or mutated) flanked by large intronic sequences cloned between the *SERPING1* exons (exons A and B, targeted by the RT-PCR primers). The plasmids were transfected, and the expression of the spliced chimeric transcripts (containing EA and EB ± E1') was analyzed. Two WT constructs containing E1' were used; 1 contained the single-nucleotide polymorphism c.340+1150T>C (rs779808). Bands corresponding to EA and EB spliced together or with *VHL* E1' are indicated on the right. **P* < .05, ***P* < .01, ****P* < .001, *****P* < .0001 based on Student *t* test. †Corresponds to unspecific bands verified by sequencing. The minigene experiment performed with the construction carrying the mutation c.340+574A>T (that targets the SA site of E1' in F6) confirmed the use of an alternative SA site (right panel) with the deletion of 15 nucleotides (represented in red). ns, not significant.

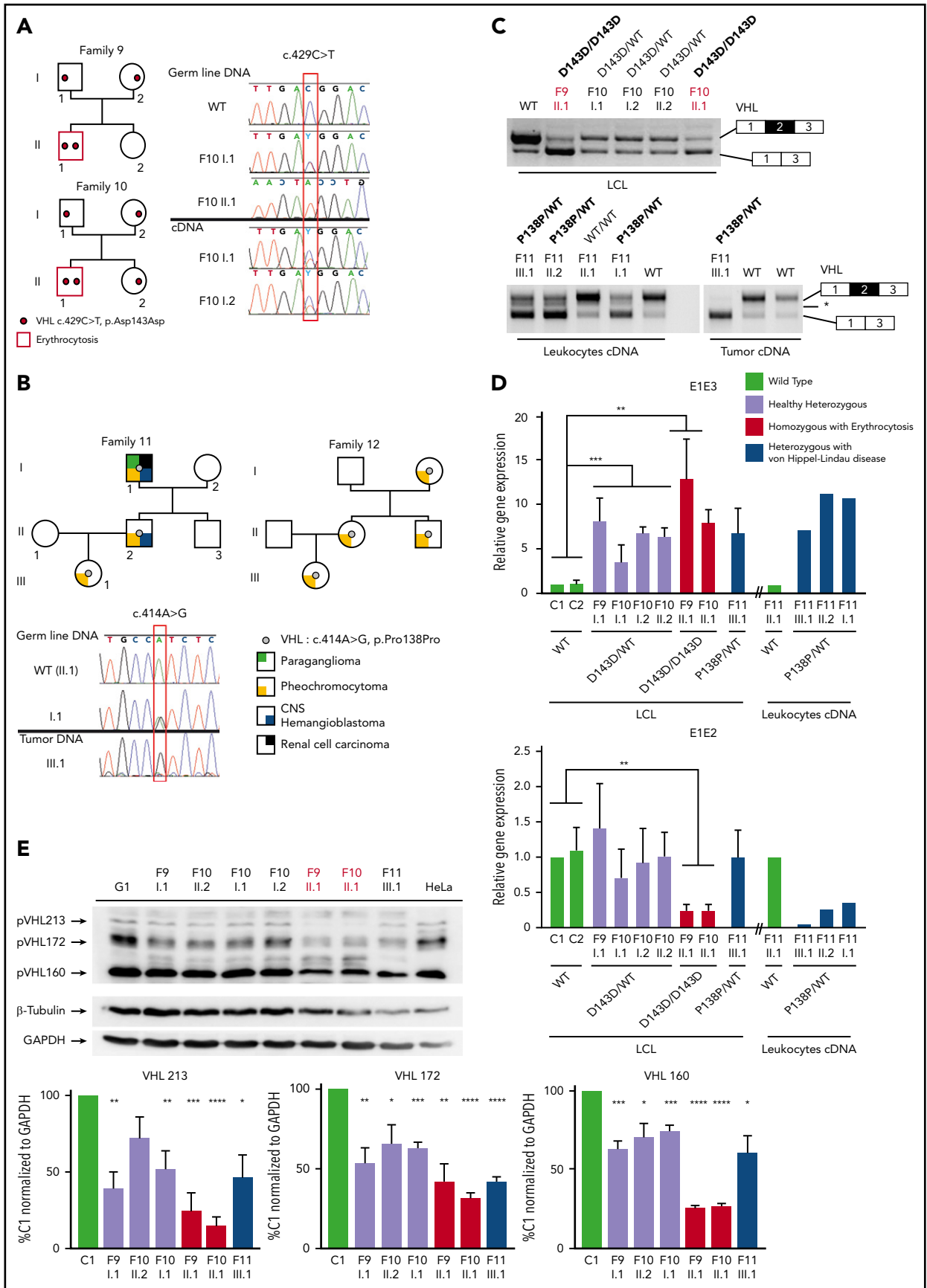


Figure 5.

reporter minigene assays in various cell lines. The experiments showed that splicing of WT E1' was barely detected (Figure 4D). Interestingly, higher molecular weight bands corresponding to the expected spliced isoforms containing E1' appeared in the presence of the mutations. Cloning and sequencing of the isoforms confirmed a retention of E1' at the expected splicing sites. The level of expression of this upper isoform, specific to E1' inclusion, was higher for mutations associated with cancers than for those with erythrocytosis, independently of the cell lines. Notably, the combination of both variants associated with cancer had a more pronounced effect than each variant individually (the single-nucleotide polymorphism showing very low effect), suggesting a synergistic effect of the variants on splicing (supplemental Figure 12). In silico analysis of c.340+617C>G consistently predicted a severe alteration of splicing by the creation of an exonic splicing enhancer site (supplemental Figure 13).

The minigene experiment performed with mutated E1' at the SA site (F6) confirmed the inclusion of E1' during splicing (Figure 4D, right panel), using the same downstream alternative SA site identified in mRNA extracted from patients' LCLs (supplemental Figure 10).

Synonymous mutations in *VHL* E2 induce exon skipping

We then focused our study on the synonymous heterozygous D143D mutation in E2, identified in the proband of F1. This mutation was also identified in the homozygous state in 2 patients with erythrocytosis (F9 and F10; Figure 5A; Table 1). The mRNA extracted from LCLs established from the different members of these 2 families was reverse transcribed and sequenced. Comparison of chromatograms obtained by sequencing of DNA vs cDNA displayed a weaker peak of the mutated allele in cDNA, reflecting a decreased expression of the mRNA transcripts carrying the mutation.

A different synonymous mutation in E2, c.414A>G p.Pro138Pro (P138P), was identified in 2 families (F11 and F12) with VHL disease (Figure 5B). This heterozygous mutation segregated with the disease in 3 generations. Sequencing of DNA extracted from the pheochromocytoma of F11-III.1 showed a loss of the WT allele in the tumor, demonstrating loss of heterozygosity as currently described in classical VHL disease.

Suspecting an effect of the synonymous mutations on splicing, we next assessed the expression of the *VHL* transcripts in patients' samples by RT-PCR (Figure 5C). We found a significant change in the ratio of expressed *VHL* isoforms, with a higher expression of the E1-E3 transcripts in LCLs of patients homozygous for D143D and in the pheochromocytoma with P138P mutation. These results suggest an effect of the mutation on splicing regulatory elements, leading to E2 skipping. In silico analysis of

the synonymous mutations indicated a potential effect on exonic splicing enhancer motifs (supplemental Figure 14).

To specifically quantify the different *VHL* isoforms, we performed quantitative RT-PCR on LCLs using TaqMan probes complementary to the *VHL* E1-E2 or E1-E3 junction (Figure 5D). All mutated samples displayed an increase of the isoform with skipped E2 (E1-E3). The patients carrying a homozygous D143D mutation presented a severe decrease of the WT isoform expression level. These results demonstrated that synonymous mutations in E2 affected mRNA isoform production.

To evaluate a potential impact on protein expression, we performed a western blot analysis. We did not observe any overexpression of the pVHL172 isoform as expected from mRNA quantification studies. However, in patients homozygous for D143D, we detected a strong downregulation of all the pVHL isoforms, which was also observed to a lesser extent in LCL samples from heterozygous patients (Figure 5E).

Proportion of *VHL* E2 skipping is correlated with disease severity in minigene experiments

To study the impact of the synonymous E2 mutations on splicing in different cell lines, we performed minigene assays. We tested the implications of the D143D and P138P mutations on splicing in addition to the nearby mutations described in patients with erythrocytosis (P138L and G144R).^{4,6,29} Indeed, these mutations may also affect splicing rather than VHL protein function. We first evaluated the loss of function of these mutants by reporter assays. We found that these could downregulate HIF in a manner similar to the WT protein, in contrast to VHL proteins lacking E2 (pVHLΔE2), which are described as nonfunctional in terms of their regulation of HIF activity^{8-10,12} (Figure 6A). Minigene experiments were performed in a variety of cell lines relevant to the studied diseases, using LCLs as control (Figure 6B). We demonstrated that all mutations caused E2 skipping in LCLs. In other cell lines, these experiments demonstrated a major impact on the *VHL* E2 splicing of the P138P mutation, which is associated with cancer. The mutations P138L, D143D, and G144R, which are all associated with erythrocytosis, displayed a weaker effect on splicing, with slight variations among cell lines but with a stronger effect in the erythroid cell line (UT7 cultured with erythropoietin).

Splicing dysregulation of *VHL* is causal in the development of disease

RNA-seq confirmed that the synonymous mutations were not silent but instead induced potent E2 skipping (Figure 6C; supplemental Figure 15A). Additional transcriptome analyses of the pheochromocytoma carrying P138P showed an upregulation of HIF target genes (typically seen in *VHL*-related pheochromocytomas) compared with the pheochromocytoma carrying a *RET* mutation (supplemental Figure 15B).³⁰ We then reanalyzed

Figure 5. Genetic and expression study of synonymous mutations in *VHL* E2. (A-B) Pedigree and sequence chromatograms of germline DNA, tumor (pheochromocytoma) DNA, or cDNA prepared from 2 families (F9 and F10) with erythrocytosis (A) and 2 families (F11 and F12) with VHL disease (B). (C) Results of RT-PCR using mRNA extracted from LCLs (F9 and F10) and leukocytes and tumor material (pheochromocytoma) (F11 and F12). (D) TaqMan quantification of the different *VHL* isoforms in LCLs (established from patients of F9, F10, and F11) cultured in the absence or presence of puromycin. TaqMan probes are specific to the *VHL* E1-E2 or E1-E3 junction. Relative gene expression was normalized to LCL control (C1). (E) Immunoblot analysis of patient LCLs. Patients with erythrocytosis are indicated in red. A representative immunoblot (upper panel) and quantification of 4 different immunoblots are displayed (lower panel). Relative gene expression was normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) expression, and results obtained with C1 were fixed at 100%. Data are represented as the mean \pm standard error of the mean. **P* < .05, ***P* < .01, ****P* < .001, *****P* < .0001 based on Student *t* test. CNS, central nervous system.

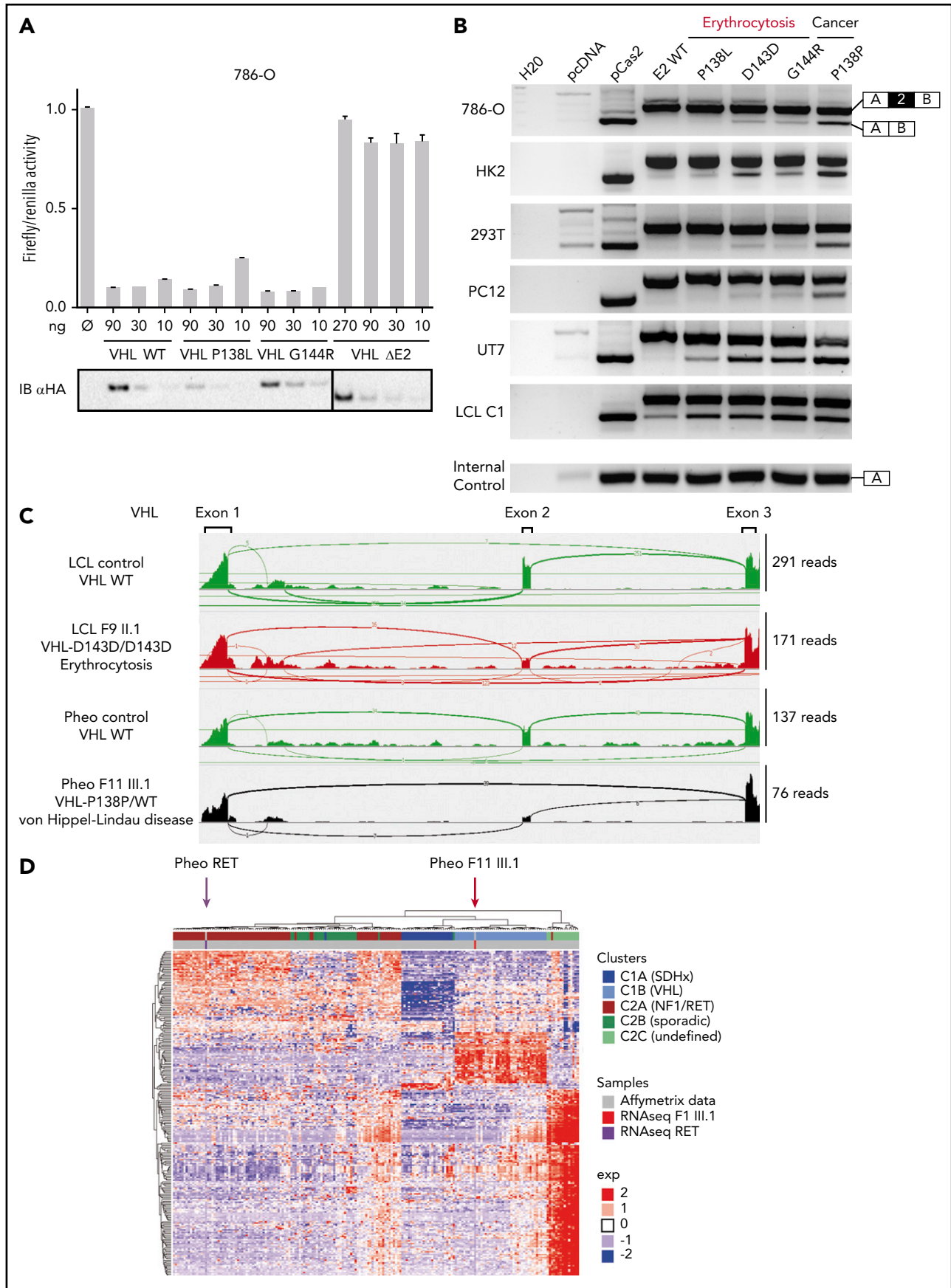


Figure 6.

our independent cohort of pheochromocytomas³⁰ using RNA-seq data from both tumors. After unsupervised classification, we observed a segregation of the P138P pheochromocytoma with other *VHL*-related tumors (C1B cluster), whereas the control pheochromocytoma bearing the *RET* mutation was grouped with other *RET*-related tumors (Figure 6D).

Discussion

The hypoxia pathway plays a central role in erythrocytosis or tumors developed by patients carrying *VHL* mutations. Nevertheless, the full molecular mechanisms at the origin of these different phenotypes remain to be elucidated. To date, functional studies of *VHL* mutants have been performed on missense mutations. We describe here, for the first time, functional studies of *VHL* mutations that did not have an impact on the coding sequence but that influenced *VHL* splicing. We discovered a complex regulation of *VHL* splicing that may help to explain the complexity of genotype/phenotype correlations observed in *VHL*-related disorders. Notably, we demonstrated that synonymous variants (D143D or P138P) could have an impact on *VHL* splicing and should be considered as pathogenic mutations. Our study points to a particular region in E2 that may be considered as a splicing regulatory domain. Therefore, it would be interesting to evaluate the impact of all the nucleotide changes described in *VHL* E2¹⁷ on splicing, in the same way as described for 2 missense mutations (P138L and G144R). We observed that, depending on the mutation in this region, the impact on splicing could be moderate (D143D, G144R, and P138L) or severe (P138P), correlating with the severity of the disease seen in individuals carrying these *VHL* mutations (erythrocytosis vs cancers). This observation confirms the hypothesis of a continuum model of tumor suppression by *VHL*.^{21,31}

Regarding the erythrocytosis developed in patients homozygous for D143D, both probands (F9 II.1 and F10 II.1) presented mutations in the β -globin gene (*HBB*) that induced hemoglobin instability or thalassemia (Table 1). This may have compensated for the strong erythropoiesis associated with a very high serum erythropoietin level associated with the D143D mutation.

More importantly, we discovered a new *VHL* cryptic exon, E1', expressed in healthy tissues. Our study led to the identification of E1' heterozygous mutations occurring in the second allele of 6 families with erythrocytosis previously associated with a heterozygous mutation in *VHL* rather than a homozygous mutation. Our investigations further confirmed that polycythemia associated with *VHL* mutation was definitely an autosomal recessive disease.

In addition, we identified an E1' homozygous mutation in a patient with erythrocytosis of unknown origin. This result demonstrated the causal role of the alterations in this new cryptic exon in the pathophysiology of erythrocytosis. Importantly, we also identified E1' mutations in patients with unexplained *VHL* disease.

This *VHL* E1' exon remained unidentified until this point because of its low expression and the fact that this deep intronic region was never explored or represented in whole-exome sequencing data. Our data showed that these newly described E1'-containing transcripts may have been polyadenylated (because they were captured by polydT in RNA-seq) but were also likely to have been subjected to NMD and may therefore have failed to produce a protein. However, we cannot exclude the possibility that they were translated into a new protein (X1) not expressed in sufficient quantity to be detected by western blot. This potential X1 isoform would contain the *VHL* E1 that encodes the NH₂-terminal part of pVHL, including 16 residues involved in HIF binding from the 17 described.²⁶ Nonetheless, the hypoxia-dependent reporter assays failed to identify a potential direct role of this isoform in the HIF pathway. Instead, our results provided compelling evidence that mutations in E1' induced a severe retention of this E1' cryptic exon, which correlated with a defect in global *VHL* protein expression. Therefore, our study strongly favors a dysregulation of splicing with a consequent downregulation of pVHL expression as the underlying cause of the diseases observed. Here, insufficient pVHL levels but not reduced HIF binding by the mutant pVHL (as seen in the Chuvash polycythemia mutation *VHL* R200W¹) may have led to an impairment of HIF degradation. Of note is the fact that the functional study of mutations identified in E1' in association with erythrocytosis resulted in a less severe impact on splicing than the mutations associated with cancer, confirming that polycythemia was associated with *VHL* hypomorphic mutations.

Our findings may have broad implications for patients with presumed congenital erythrocytosis. First, the underlying cause of congenital erythrocytosis has been identified in only approximately one third of patients in most published studies. However, previous studies have focused on missense and nonsense changes in coding regions and known regulatory domains of candidate genes. This has also been the case in patients with suspected *VHL* disease. Our study shows that synonymous exonic changes, as well as changes within intronic sequences affecting exon splicing, may be responsible for these disorders and should be considered during the diagnostic process. Notably, the *VHL* E1' exon should be added to the list of regions routinely sequenced in patients with congenital erythrocytosis. Second, the detection of molecular changes

Figure 6. Functional study of synonymous mutations in *VHL* E2. (A) Functional hypoxia response element (HRE)-dependent reporter assays were performed in 786-O cells to evaluate the impact of *VHL* mutations in E2 (P138L and G144R) on *VHL* protein activity. The *VHL* protein lacking E2 (pVHL172/*VHL*ΔE2) was used as a negative control. The results are expressed as firefly luciferase activity relative to *Renilla* luciferase as an internal control; 1.0 unit denotes the basal activity of endogenous HIF-2 α using the HRE luciferase reporter plasmid. Immunoblots using an antibody specific for the hemagglutinin tag (HA) were used to detect HA-*VHL*. (B) Characterization of *VHL* E2 skipping by minigene analyses. Minigene experiments were performed in a variety of cell lines relevant to the studied diseases: renal (293T, 786-O, and HK2), pheochromocytoma (phéo; PC12), erythroid cell line (UT7 cultured with erythropoietin), and LCL. RT-PCR was performed on mRNA obtained from cell lines transfected with a minigene construct containing *VHL* E2 flanked by intronic sequences cloned between the *SERPING1* exons (exons A and B; targeted by the RT-PCR primers). Bands corresponding to EA and EB spliced together or with *VHL* E2 are indicated on the right. (C) Sashimi plots from RNA-seq data. The positions of the different *VHL* exons are indicated, with the maximum number of reads for each exon indicated at the right. Splice junctions are denoted by the horizontal links, with details provided in supplemental Figure 15A. (D) Heatmap of pheochromocytoma transcriptome data. A comparison of transcriptome data from the pheochromocytoma from patient F11 III.1 (with P138P mutation) vs Affymetrix data from the largest available cohort of paragangliomas/pheochromocytomas (recruited by the French COMETE network) that identified homogeneous molecular subgroups associated with susceptibility genes (Burnichon et al³⁰).

has implications for the clinical management of patients. For example, phlebotomy in patients with *VHL*- or *HIF2A*-related erythrocytosis may worsen the clinical situation by increasing the risk and severity of pulmonary hypertension in these patients.³² Third, the confirmation of the continuum model of tumor suppression by *VHL* helps us to understand the very low frequency of secondary tumors in patients with *VHL*-related erythrocytosis.³³ However, this also means that later occurrence of such neoplasms cannot be definitely excluded. Therefore, the clinical management of these patients should include regular follow-up to assess the risk for thromboembolic complications, pulmonary hypertension, and cardiovascular disease, as well as regular examinations to check for the presence of typical *VHL*-related tumors. Patients with *VHL* disease resulting from E1' mutations will also benefit from regular screening for tumors. Finally, the detection of these new genetic changes will also allow for appropriate genetic counseling of affected patients and their families.

In conclusion, *VHL* is a major tumor suppressor gene that plays a pivotal role in the oxygen-sensing pathway, which is involved in multiple physiological (eg, angiogenesis and erythropoiesis) and pathological (eg, cancer) processes. Our findings regarding the complex splicing regulation of this gene in erythrocytosis and tumorigenesis may therefore open new avenues for diagnosis of these conditions as well as research on the biological mechanisms related to the hypoxia-signaling pathway. Notably, we suggest further targeted exploration of the *VHL* E1' region in unresolved cases of congenital erythrocytosis, inherited kidney cancers, hereditary paraganglioma/pheochromocytoma syndrome, and hemangioblastomas, in addition to all types of sporadic tumors with altered hypoxia signaling.

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Authorship

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Footnotes

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