

ORIGINAL RESEARCH

Germline mutations in the new E1' cryptic exon of the VHL gene in patients with tumours of von Hippel-Lindau disease spectrum or with paraganglioma

Alexandre Buffet ⁽¹⁾, ¹ Bruna Calsina, ² Shahida Flores, ³ Sophie Giraud, ^{4,5} Marion Lenglet, ^{6,7} Pauline Romanet, ⁸ Elisa Deflorenne, ¹ Javier Aller, ⁹ Isabelle Bourdeau, ¹⁰ Brigitte Bressac-de Paillerets, ^{5,11} María Calatayud, ¹² Caroline Dehais, ¹³ Erwan De Mones Del Pujol, ¹⁴ Atanaska Elenkova, ¹⁵ Philippe Herman, ¹⁶ Peter Kamenický, ¹⁷ Sophie Lejeune, ^{5,18} Jean Louis Sadoul, ¹⁹ Anne Barlier, ^{5,8} Stephane Richard, ^{5,6,20} Judith Favier, ¹ Nelly Burnichon, ^{1,21} Betty Gardie, ^{6,7} Patricia L Dahia, ³ Mercedes Robledo, ² ida Flores,³ Sophie Giraud,^{4,5} Deflorenne,¹ Javier Aller,⁹ lerets,^{5,11} María Calatayud,¹² ol,¹⁴ Atanaska Elenkova,¹⁵ lie Lejeune,^{5,18} Jean Louis Sadoul,¹⁹ h Favier,¹ Nelly Burnichon,^{1,21} Robledo,² responsible for the development in affected patients of renal cysts or clear cell carcinomas, and other features as retinal or central nervous system Anne-Paule Gimenez-Roqueplo^{1,5,21}

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For numbered affiliations see end of article.

Correspondence to

Dr Alexandre Buffet, Université de Paris, PARCC, INSERM, Equipe Labellisée par la Ligue contre le Cancer, F-75015, Paris, France: alexandre.buffet@inserm.fr

BC, SF, SG and ML contributed equally.

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ABSTRACT **Backgrounds** The incidence of germline mutations in the newly discovered cryptic exon (E1') of VHL gene in patients with von Hippel-Lindau (VHL) disease and in patients with paraganglioma or pheochromocytoma

(PPGL) is not currently known.

Methods We studied a large international multicentre cohort of 1167 patients with a previous negative genetic testing. Germline DNA from 75 patients with a single tumour of the VHL spectrum ('Single VHL tumour' cohort), 70 patients with multiple tumours of the VHL spectrum ('Multiple VHL tumours' cohort), 76 patients with a VHL disease as described in the literature ('VHL-like' cohort) and 946 patients with a PPGL were screened for E1' genetic variants.

Results Six different genetic variants in E1' were detected in 12 patients. Two were classified as pathogenic, 3 as variants of unknown significance and 1 as benign. The rs139622356 was found in seven unrelated patients but described in only 16 patients out of the 31 390 of the Genome Aggregation Database (p<0.0001) suggesting that this variant might be either a recurrent mutation or a modifier mutation conferring a risk for the development of tumours and cancers of the VHL spectrum.

Conclusions VHL E1' cryptic exon mutations contribute to 1.32% (1/76) of 'VHL-like' cohort and to 0.11% (1/946) of PPGL cohort and should be screened in patients with clinical suspicion of VHL, and added to panels for Next Generation Sequencing (NGS) diagnostic testing of hereditary PPGL. Our data highlight the importance of studying variants identified in deep intronic sequences, which would have been missed by examining only coding sequences of genes/exomes. These variants will likely be more frequently detected and studied with the upcoming implementation of wholegenome sequencing into clinical practice.

INTRODUCTION

Von Hippel-Lindau (VHL) disease is an autosomaldominant renal cancer predisposition syndrome¹

other features as retinal or central nervous system haemangioblastomas, pancreatic cysts or neuroendocrine tumours, endolymphatic sac tumours and pheochromocytomas and/or paragangliomas (PPGLs). A germline mutation (including gross deletion) is identified in one of the three exons of VHL in almost all affected patients.² Nevertheless, some patients with clinically diagnosed VHL disease, but without identified VHL germline mutation, have been reported.³ One of the tumour types of the VHL tumour spectrum, PPGL, are rare neuroendocrine tumours with a great genetic heterogeneity and the highest heritability rate with about 40% of genetically determined forms.^{4 5} Indeed, to date, approximately 17 susceptibility genes have been **a** reported but two thirds of identified mutations are found in SDHB, SDHD and VHL genes.⁶⁷

Recently, a cryptic exon of VHL gene, named E1', has been discovered. A germline mutation in the first intronic region which results in creation of a cryptic exon designated E1' was found in one large family with a typical VHL disease and without any alteration in the other VHL exons.⁸ VHL gene is one of the major PPGL susceptibility genes but, to date, E1' exon has not been included in PPGL target gene panels.

Hence, our objective was to assess the prevalence of E1' germline mutations in two international cohorts of patients: first, in 221 patients with a single or multiple tumours suggesting a VHL disease and then in 946 with a PPGL but without an identified mutation in the three VHL exons or in the main PPGL susceptibility genes, respectively.

METHODS

Patient's selection

four different groups:

A total of 1167 patients were analysed, divided into



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| Table 1 | Main clinical and tumour characteristics of the different |
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| cohorts | |

| conorts | |
|---|--------------|
| Total patients n=1167 | |
| VHL-like, n=76 patients | |
| Age at first diagnosis mean (min-max) | 45.4 (20–76) |
| Multiple haemangioblastomas | 16 (21%) |
| Haemangioblastoma with another VHL tumour | 28 (37%) |
| One VHL tumour and family history of VHL tumour | 32 (42%) |
| Multiple VHL tumours, n=70 patients | |
| Age at first diagnosis mean, (min-max) | 55 (11–81) |
| Three or more VHL tumours | 3 (4%) |
| Two VHL tumours | 67 (96%) |
| Single VHL tumour, n=75 patients | |
| Age at first diagnosis mean (min-max) | 34.6 (11–78) |
| Clear cell renal cell carcinoma | 3 (4%) |
| Cerebral haemangioblastoma | 27 (36%) |
| Retinal haemangioblastoma | 10 (13.3%) |
| Other tumours | 35 (46.6%) |
| PPGL, n=946 patients | |
| Age at first diagnosis mean (min-max) | 43 (8–94) |
| Benign PPGL | 869 (92%) |
| Single benign PPGL | 771 (82%) |
| Multiple benign PPGL | 98 (10%) |
| Metastatic PPGL | 77 (8%) |
| Single metastatic PPGL | 67 (7%) |
| Multiple metastatic PPGL | 10 (1%) |
| Familial PPGL | 17 (2%) |
| PPGL, paraganglioma; VHL, von Hippel-Lindau. | |

PPGL, paraganglioma; VHL, von Hippel-Lindau.

- ▶ 946 patients with PPGL but without germline mutation in major PPGL susceptibility genes ('PPGL' cohort) (table 1 and online supplementary table \$1).
- ▶ 76 patients with a VHL disease as defined in the literature,²⁹ that is, patients with multiple haemangioblastomas, or a single haemangioblastoma with another tumour of the VHL spectrum, or one tumour of the VHL spectrum (excepted epididymal and renal cysts) and family history of VHL tumour but no germline *VHL* gene mutation ('VHLlike' cohort).
- ▶ 70 patients with multiple tumours of the clinical spectrum of VHL disease but who did not fill the definition of a VHL disease and who had no germline *VHL* mutation ('Multiple VHL tumours' cohort).
- ▶ 75 patients with a single tumour of the VHL spectrum without *VHL* mutation occurring at a young age ('Single VHL tumour' cohort) (table 1 and online supplementary table S2).

Germline DNA from 'VHL-like', 'Multiple VHL tumours' and 'Single VHL tumour' cohorts had been previously tested for *VHL* gene by Sanger sequencing or Next Generation Sequencing and large rearrangements by MLPA or QMPSF. The procedures used for PPGL diagnosis were in accordance with international guidelines.¹⁰¹¹

Moreover, a control cohort of 198 European subjects without VHL manifestation was analysed in order to determine the frequency of variant in the general population.

Each patient signed a written informed consent for genetic analyses.

Direct sequencing of the E1' cryptic exon of VHL on germline DNA

Sanger sequencing on germline DNA of E1' was performed as previously described.⁸ Variants interpretation was performed by using different criterions: ACMG criteria,¹² allele frequency in databases, phenotype of patients and tumour analysis as described below.

VHL gene analysis in tumour

Tumour DNA was extracted from frozen or paraffin embedded tumour by the QIAamp DNA minikit (Qiagen). Loss of heterozygosity (LOH) was evaluated by (1) Sanger sequencing of the E1' cryptic exon of *VHL* by mutation-specific primers and (2) microsatellite analysis on D3S1537, D3S1038, D3S1317 D3S3547, D3S3727 as previously described.^{13 14} *VHL* gene deletion on tumour DNA was assessed with the *SALSA MLPA* P016 *VHL* probemix (MRC-Holland).

CA9 immunochemistry

Immunohistochemistry was performed as previously described on 6 μ m slides cut from paraffin-embedded tumours with anti CA9 antibody (1/1500, ab15086, Abcam).¹⁵ Antigen retrieval was performed by boiling slides in Tris-EDTA buffer (pH9) for 45 min. Revelation was performed using Histogreen as a chromogen. Images were acquired with a Leica DM400B microscope with Leica Application Suite software V.2.8.1 and a Leica DFC420C camera.

PNMT and VHL RT-qPCR

RNA was extracted from paraffin embedded tumours of six control PPGL (3 NF1-related, 2 RET-related and 1 TMEM127tex related PPGL), 5 VHL- related PPGL (all carrying a missense mutation in VHL gene) and patients #3 and #10 PPGL by using and the Maxwell 16 LEV RNA FFPE Purification Kit (Promega). data RNA was quantified and its purity assessed with a NanoDrop ND-1000 spectrophotometer (Labtech). RT PCR was performed on 1000 ng of RNA with iScript cDNA Synthesis Kit iScript Bululu (BioRad). Then, as described in,¹⁶ pre amplification of *PNMT*, VHL, GAPDH and 18S on complementary DNA was performed with SsoAdvanced PreAmp Supermix (BioRad). Because of RNA ⊳ fragmentation, all primers were designed to amplify amplicons smaller than 100 bp. We used two VHL primer sets. The first set amplified the VHL transcript including exons 1 and 2 (E1-E2) (F: 5'- CATCCACAGCTACCGAGGTC-3' overlapping exons 1 and 2 and R: 5'-GTGTGTCCCTGCATCTCTGA-3' located nd on exon 2). The second set amplified the VHL transcript with <u>s</u> exon 1 and the cryptic exon (E1-E1') (F: 5'-GCATCCACAGC-TACCGAGTC-3' overlapping exon 1 and the cryptic exon and R: 5'-AGTCTCCCCAGGAGGAATGT-3' located on the cryptic exon). Quantitative PCR was performed on VHL (E1-E2), nologies PNMT, GAPDH and 18S by SYBR Green Master MixSybrGreen (BioRad) on the C1000 Touch (BioRad) and VHL (E1-E1') was amplified by PCR in parallel with GAPDH. All experiments were performed in duplicate three times.

Statistical analysis

Statistical analysis was carried out with GraphPad software. Differences between allele's frequency in gnomAD and our cohort of patients and relative risk were assessed by χ^2 tests. A p<0.05 was considered significant.

RESULTS

We analysed the germline DNA of 1167 patients from France, Spain, Canada and the USA. We identified a rare germline

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| Table 2 | Table 2 Patients with a genetic variant in the E1' cryptic exon of VHL gene | riant in the E1' cry | ptic exon of VHL | gene | | | | | | | |
|---|--|---|--|---|---|------------------------------------|--------------------------------|-------------------|----------------------------------|----------------------------------|--------------------------|
| | | | | | Allele | | | | PNMT and | | |
| Patient number | Phenotype (age at first diagnosis) | Follow-up duration (years) | Germline genetic variant | dbSNP ID | frequency* (%) | НОН | Somatic <i>VHL</i> mutation | CAIX IHC | <i>VHL (E1-E2)</i> expression | <i>VHL</i> (E1-E1' expression | Classification of VUS |
| #1 | Carotid body PGL (47) | 5 | c.340+563C>T | | NA | Not | ND | Neg | ND | ND | Benign |
| #2 | Carotid body PGL (34) | - | c.340+578C>T | rs139622356 | 0.05 | NA | NA | NA | NA | NA | VUS |
| #3 | PCC (38) | 7 | c.340+578C>T | rs139622356 | 0.05 | No | No | Neg | Normal | Increase | VUS |
| #4 | Carotid body PGL (74) | - | c.340+578C>T | rs139622356 | 0.05 | NA | NA | NA | NA | NA | VUS |
| #5 | Carotid body PGL (56) | - | c.340+578C>T | rs139622356 | 0.05 | NA | NA | NA | NA | NA | VUS |
| 9# | PGL+ccRCC (81–82) | 2 | c.340+578C>T | rs139622356 | 0.05 | NA | NA | NA | NA | NA | VUS |
| L# | bilccRCC (49–51) | NA | c.340+578C>T | rs139622356 | 0.05 | NA | NA | NA | NA | NA | VUS |
| 8# | ccRCC (39) | NA | c.340+578C>T | rs139622356 | 0.05 | NA | NA | NA | NA | NA | VUS |
| 6# | Multiple retinal HMB (36) | 32 | c.340+617C>G‡ | | NA | NA | NA | NA | NA | NA | Pathogenic |
| #10 | PCC (11) | 15 | c.340+682T>C | | NA | No | c.482G>A p.R161Q | Pos | Decrease | Increase | Pathogenic |
| #11 | Multiple HN PGL (12) | - | 340+725A>T | | NA | NA | NA | NA | NA | NA | VUS |
| #12 | PCC (34) | 10 | c.340+866C>A | rs536631685 | 0.02 | NA | NA | NA | NA | NA | VUS |
| *Frequency †loss of the ‡Mutation bilccRCC, b pheochrom | *Frequency in gnomAD or 1000 Genomes, . tloss of the mutated allele. #Mutation described in Lenglet <i>et al.</i> ⁸ bilccRCC, bilateral clear cell renal cell carcinoma; HMB, haemangioblastoma; HN, head and neck; LOH, loss of heterozygosity; NA, none available; ND, not done; Neg, negative immunochemistry; PCC, pheochromocytoma; PGL, paraganglioma; Pos, positive immunochemistry; VHL, von Hippel-Lindau; VUS, variant of uncertain significance. | oma; ccRCC, clear cell 35, positive immunoch | renal cell carcinoma iemistry; VHL, von Hig | ı; HMB, haemangiobl ppel-Lindau; VUS, va | lastoma; HN, head a rriant of uncertain si | and neck; LOH, los: gnificance. | s of heterozygosity; N | A, none available | ; ND, not done; Neg, | negative immunoche | mistry; PCC, |

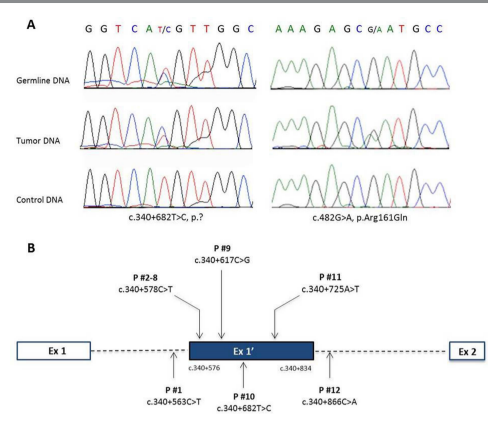


Figure 1 VHL E1' variants identified: mutations identified in patient #10 on the germline DNA and somatic DNA (A); location of germline variants found in VHL E1'(B).

genetic variant (minor allele frequency <1%) in the E1' VHL cryptic exon in 12 patients (1%). One of these patients was classified as 'VHL-like' (1 of 76 patients, 1.3%), 2 as 'Multiple VHL tumours' (2/70 patients, 2.9%), 1 as 'Single VHL tumour' (1/75 patients, 1.3%) and 8 belonged to the 'PPGL' cohort (8/946, 0.8%) (table 2). Among these 12 patients, we identified 6 different variants, 4 in the E1' and 2 at the intron-exon junction and we considered only two variants as pathogenic mutations (figure 1). None of these variants was found in a control cohort of 198 European subjects without VHL manifestations.

Seven patients (patients #2 to #8) (0.6%) carried the same rare variant of uncertain significance (VUS), c.340+578C>T which is referenced in dbSNP as rs139622356 and has been previously reported in the Genome Aggregation Database (gnomAD). The five remaining patients carried different E1' variants. One of them (patient #9) harbours the c.340+617C>G mutation previously described in the original paper.⁸ Patient #12 carried the c.340+866C>A VUS, which is referenced in dbSNP (rs536631685) and 1000 Genomes, but not in the Genome Aggregation Database (gnomAD). Finally, three novel VUS of the E1' VHL cryptic exon were discovered in the three remaining patients. None of the four patients with a PPGL and an E1' VUS have developed VHL spectrum tumour(s) during their follow-up and none of them had family history of VHL disease (table 2); however, segregation analysis was only performed in patient #9. The proband's mother did carry the variant and had a pancreatic cyst and multiple vertebral body haemangiomas which both are evocating of VHL disease.¹⁷

Among the remaining 11 patients, 3 tumours were available, 2 as paraffin embedded samples (patients #1, #10) and 1 as a frozen tumour (patient #3). None of them presented a LOH at VHL locus and the mutated allele was lost as determined by

Protected by copyright, including for uses related to text and Sanger sequencing in tumour #1. In tumour DNA of patient #10, which harbours the c.340+682T>C variant, we identidata fied a second variant in the exon 3 of VHL (c.482G>A; p.Arg161Gln), known to be pathogenic (figure 1). This somatic mutation was previously described in this patient.¹⁸ In the absence of LOH, this exon 3 variant may function as the second VHL hit in this tumour. No other mutation of the VHL gene was given by the transfer data in the transfer data in the transfer data is the transfer data in the transfer data in the transfer data is the transfer data in the transfer data is the transfer data in the transfer data identified in tumour DNA of patient #3.

training To validate and classify these different VUS, we carried out different functional studies on available tumour tissues. We first performed immunohistochemistry to study the expression of CA9, known to be expressed at the membrane of tumour cells in case of VHL inactivation.¹⁹ A membranous positive CA9 immunostaining has been previously reported in VHL-related PPGL, haemangioblastoma, endolymphatic tumours and ccRCC.^{15 20 2} We observed a cluster of tumour cells with a positive membranous CA9 immunostaining only in the PPGL of patient #10 technologies (figure 2A) which can be seen in VHL-related PPGL.¹⁵ Then, we assessed the expression of PNMT gene, which is one of the most downregulated genes in VHL-related PPGL,^{22 23} by RT-qPCR. As expected, the PPGL of patient #10 exhibited a significant low expression of PNMT mRNA, comparable to the VHL-tumours used as controls. On the contrary, the level of PNMT expression was equivalent to control tumours in the PPGL of patient #3, which produced both epinephrine and norepinephrine. Finally, we analysed the expression of VHL gene by RT-qPCR. We assessed the expression of two different VHL mRNA: the mRNA containing the exons 1 and 2 (E1-E2), which will lead with the exon 3 to the expression of the two main VHL proteins (pVHL213 and pVHL160) and VHL mRNA containing the exon 1 and E1' (E1-E1'), which was previously described as increased in tumour or in lymphoblastoid cell lines

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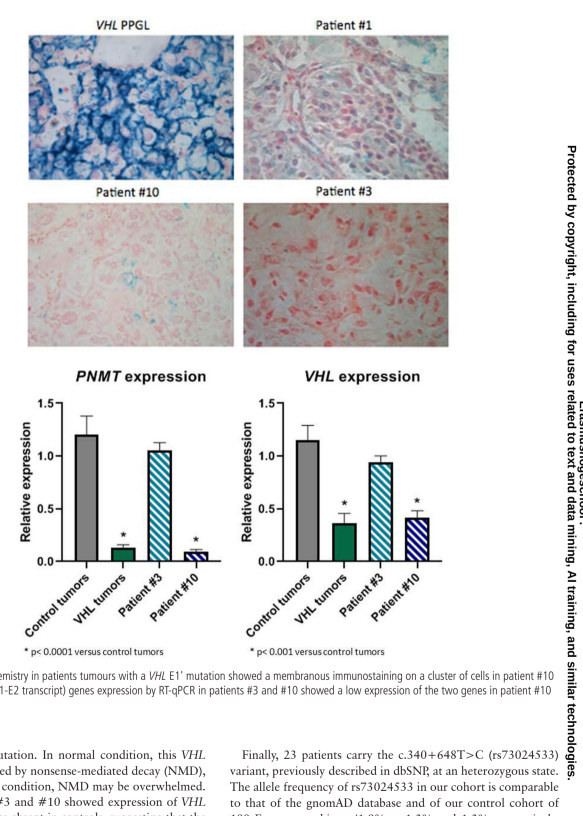


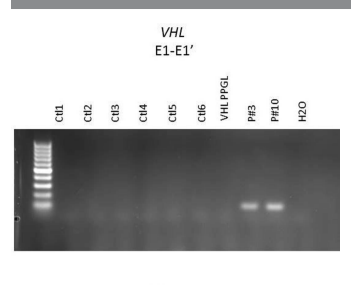
Figure 2 CA9 immunochemistry in patients tumours with a VHL E1' mutation showed a membranous immunostaining on a cluster of cells in patient #10 PPGL (A). PNMT and VHL (E1-E2 transcript) genes expression by RT-qPCR in patients #3 and #10 showed a low expression of the two genes in patient #10 tumours (B).

of patients with E1' mutation. In normal condition, this VHL E1-E1' mRNA is degraded by nonsense-mediated decay (NMD), and in this pathological condition, NMD may be overwhelmed. The PPGL of patients #3 and #10 showed expression of VHL E1-E1' mRNA which was absent in controls, suggesting that the two variants change the VHL mRNA splicing (figure 3). Moreover, the PPGL of patient #10 showed a low expression of VHL E1-E2 mRNA comparable to the VHL-related PPGL used as control (figure 2B). Altogether, these data provide evidence that this VHL E1' mutation (c.340+682T>C) is a pathogenic mutation that combined with the second mutation (c.482G>A; p.Arg161Gln) induce tumorigenesis.

The allele frequency of rs73024533 in our cohort is comparable to that of the gnomAD database and of our control cohort of 198 European subjects (1.9% vs 1.3% and 1.3%, respectively, p = 0.0536).

DISCUSSION

E1' mutations were previously described by Lenglet et al in eight families, either with erythrocytosis or VHL disease. These mutations led to an abnormal VHL mRNA with the insertion of the



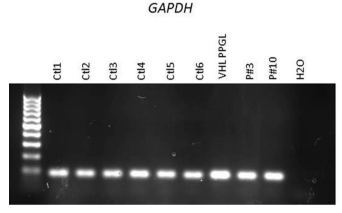


Figure 3 Expression of E1-E1' transcript of *VHL* gene showed expression only in patients #3 and #10.

E1' in the transcript and to mRNA degradation by NMD and to global defect in VHL protein expression.⁸

In our large international study, we identified four new germline variants in E1' VHL gene and we classified two of them as pathogenic, representing 1.3% of 'VHL-like' cohort (1/76 patients) and 0.11% of 'PPGL' cohort (1/946 patients). Our patients did not have all the manifestation of VHL disease. However, in the single patient in whom a familial genetic screening was performed (patient #9), the proband's mother had her first screening (cerebral and medullary MRI and abdominal CT scan) at the age of 70 years old, which diagnosed one pancreatic cyst and multiple vertebral body haemangiomas. Interestingly, multiple vertebral body haemangiomas are rare in VHL disease but have been described in patients with Chuvash polycythemia, a disease secondary to a recurrent germline biallelic mutation in VHL gene (c.598C>T, p.Arg200Trp).²⁴ Our data suggest incomplete penetrance of E1' VHL mutations, as it was previously described for the SDHA gene-another PPGL susceptibility gene- mutations that exhibit a relatively high allele frequency in gnomAD.²¹

We have identified the same variant c.340+578C>T (rs139622356) in seven patients, but our tumour analyses were not able lead to the classification of this variant in a pathogenic variant. Indeed in one tumour with this variant, we identified the E1-E1' mRNA which suggest that the variant is pathogenic. However, epinephrine secretion and *PNMT* expression of this tumour are strong indicator against the diagnosis of *VHL*-related PPGL.²⁶ Moreover, we identified this variant in 0.6% of our

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cohort, which is 10 times more frequent in our cohort than in reference databases. Indeed, this variant is described in 0.05% of gnomAD subjects (7/1167 vs 16/31 390, p<0.0001). It is note-worthy that in Tuscan and Iberian subjects reported in the 1000 Genomes project, the frequency of this rs139622356 is 0.9%. All these data suggest that this variant could be either a pathogenic variant that is not implicated in the PPGL of our patient because of the lack of LOH/second *VHL* mutation, or a modifier variant contributing potentially to an 8.5-fold risk (95% CI 4.4 to 14.3, p<0.0001) for development of PPGL or VHL tumours. Hence, more functional analyses and more tumours analyses will be required to achieve a definitive conclusion.

Our study demonstrates that E1' VHL variants are rare events in 'VHL-like' and 'PPGL' patients, but nearly as frequent as the VHL mutation rate in exons 1 and 2 in patients with PPGL (in g the molecular genetic laboratory of Hôpital Européen Georges Pompidou-Paris-France VHL mutation rate in exon 1 has been reported to be 0.74% (p=0.062), in exon 2: 0.18% (p=0.99) and in exon 3: 0.92% (p=0.0264),²⁷ or as frequent as in exons of other PPGL susceptibility genes (for instance, the mutation rate in exon 1 of SDHD is 0.43%). However, because patients with well-established VHL pathogenic mutations were excluded from our cohort, the current frequency may be an underestimation. As the identification of VHL variants has important implications for management and follow-up of patients and relatives, we suggest that E1' cryptic exon should be added to NGS diagnostic panels. Considering the genetic heterogeneity of PPGLs and the high rate of detectable driver mutations in these tumours,¹⁰ a low frequency of variants in any given new gene/ exons is not unexpected. However, the interpretation of these E1' variants might be difficult and more functional analyses has to be designed in order to validate these variants. Finally, our study underlines the importance of variants identified in deep intronic sequences, which would have been missed by examining only coding sequences of genes/exomes. These variants will likely be more frequently detected and studied in the next future with the upcoming implementation of whole-genome sequencing into clinical practice.

Author affiliations

¹Université de Paris, PARCC, INSERM, Equipe Labellisée par la Ligue contre le Cancer, F-75015, Paris, France

²Hereditary Endocrine Cancer Group, Spanish National Cancer Research Centre (CNIO), Madrid, Spain; and Centro de Investigación Biomédica en Red de Enfermedades Raras (CIBERER), Madrid, Spain

³Division of Hematology and Medical Oncology, Department of Medicine, Mays Cancer Center, University of Texas Health Science Center at San Antonio, San Antonio, Texas, USA

⁴Genetics Department, Hospices Civils de LYON (HCL), University Hospital, East Biology and Pathology Center, B-A3, 59 Bld Pinel, 69677, Bron, France

⁵Réseau National pour Cancers Rares PREDIR labellisé par l'Institut National contre le Cancer, Hôpital Bicêtre, Assistance Publique-Hôpitaux de Paris, Le Kremlin-Bicêtre, France

⁶École Pratique des Hautes Études, PSL Research University, Paris, France

⁷L'Institut du Thorax, INSERM, Centre National de la Recherche Scientifique (CNRS), Université de Nantes, Nantes, France

⁸Aix Marseille Univ, APHM, INSERM, MMG, Laboratory of Molecular Biology Hospital La Conception, Marseille, France

⁹Endocrinology and Nutrition Service. Hospital Universitario Puerta de Hierro, 28222, Majadahonda, Spain

¹⁰Division of Endocrinology, Department of Medicine, Centre de recherche du Centre hospitalier de l'Université de Montréal (CRCHUM), Université de Montréal, Montréal, Quebec, Canada

¹¹Gustave Roussy, Université Paris-Saclay, Département de Biopathologie and INSERM U1186, Villejuif, F-94805, France

 ¹²Department of Endocrinology and Nutrition, Hospital Universitario 12 de Octubre, 28041 Madrid, Spain
¹³Assistance Publique-Hôpitaux de Paris, Hôpitaux Universitaires La Pitié Salpêtrière-

¹³Assistance Publique-Hôpitaux de Paris, Hôpitaux Universitaires La Pitié Salpêtrière-Charles Foix, Service de Neurologie 2-Mazarin, Paris, France

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¹⁴Service ORL, Hôpital Pellegrin, CHU de Bordeaux, Bordeaux, France ¹⁵Department of Endocrinology, USHATE "Acad. Ivan Penchev", Medical University Sofia, Sofia, Bulgaria ¹⁶Assistance Publique, Hôpitaux de Paris, Service ORL-CCF, hôpital Lariboisière,

université Paris VII. Paris. France

⁷Assistance Publique-Hôpitaux de Paris, Hôpital de Bicêtre, Service d'Endocrinologie et des Maladies de la Reproduction, Centre de Référence des Maladies Rares de l'Hypophyse, Le Kremlin Bicetre, France

¹⁸Department of Clinical Genetics, Centre Hospitalier Régional et Universitaire de Lille, Lille, France

⁹Service d'Endocrinologie, Hôpital de L'Archet, CHU de Nice, Nice, France

²⁰Génétique Oncologique EPHE, INSERM U1186, Gustave Roussy Cancer Campus, Villejuif, France

¹Assistance Publique-Hôpitaux de Paris, Hôpital Européen Georges Pompidou, Service de Génétique, Paris, France

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Contributors APGR conceived and supervised the study, ABu, APGR, MR, PLD and BG designed the study and analysed the results. JF and NB participated to the data analysis. ABu., BC, SF, SG, ML and ED designed and performed the experiments. PR, JA, IB, BBdP, MC, CD EdM, AE, PH, PK, SL, JLS, ABa, SR, BG, PLD, MR and APGR collected subjects and clinical parameters. ABu and APGR wrote the manuscript. ABu prepared the figures and the tables. All the authors discussed the results and commented the manuscript.

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ORCID iD

Alexandre Buffet http://orcid.org/0000-0002-4742-2708

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