

A SUMOylation-defective MITF germline mutation predisposes to melanoma and renal carcinoma

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So far, no common environmental and/or phenotypic factor has been associated with melanoma and renal cell carcinoma (RCC). The known risk factors for melanoma include sun exposure, pigmentation and nevus phenotypes1; risk factors associated with RCC include smoking, obesity and hypertension². A recent study of coexisting melanoma and RCC in the same patients supports a genetic predisposition underlying the association between these two cancers3. The microphthalmia-associated transcription factor (MITF) has been proposed to act as a melanoma oncogene⁴; it also stimulates the transcription of hypoxia inducible factor⁵ (HIF1A), the pathway of which is targeted by kidney cancer susceptibility genes⁶. We therefore proposed that MITF might have a role in conferring a genetic predisposition to co-occurring melanoma and RCC. Here we identify a germline missense substitution in MITF (Mi-E318K) that occurred at a significantly higher frequency in genetically enriched patients affected with melanoma, RCC or both cancers, when compared with controls. Overall, Mi-E318K carriers had a higher than fivefold increased risk of developing melanoma, RCC or both cancers. Codon 318 is located in a small-ubiquitin-like modifier (SUMO) consensus site (YKXE) and Mi-E318K severely impaired SUMOylation of MITF. Mi-E318K enhanced MITF protein binding to the HIF1A promoter and increased its transcriptional activity compared to wild-type MITF. Further, we observed a global increase in Mi-E318Koccupied loci. In an RCC cell line, gene expression profiling identified a Mi-E318K signature related to cell growth, proliferation and inflammation. Lastly, the mutant protein enhanced melanocytic and renal cell clonogenicity, migration and invasion,

consistent with a gain-of-function role in tumorigenesis. Our data provide insights into the link between SUMOylation, transcription and cancer.

MITF encodes a member of the Myc supergene family of basic helix-loop-helix zipper transcription factors. It has a complex intron-exon structure, yielding protein products with different amino termini⁷. The M-isoform (expressed in the melanocyte lineage) of MITF regulates expression of a large set of genes promoting proliferation and invasion. For instance, it controls the expression of MET⁵ and CDKN2A/p16INK4A⁵, which have key roles in melanoma development. The role of the A-isoform of MITF (expressed in kidney) in renal cell transformation has yet to be elucidated. However, MITF belongs to the same family of transcription factors (the MiT family) as the TFEC, TFE3 and TFEB genes; the two latter were identified as targets of somatic recurrent translocations in a RCC subtype found predominantly in children and young adults⁸.

We sequenced the *MITF* gene in 62 patients with melanoma and RCC. Five patients exhibited a germline heterozygous missense substitution, p.E318K (c.952G>A, in MITF isoform 4, NCBI accession NM_000248.3) (Fig. 1a). The frequency of this variant was significantly higher in patients with both melanoma and RCC (melanoma + RCC) than in 1,659 population-based cancer-free controls ($P=1.3\times10^{-4}$). Carriers of the p.E318K variant (Mi-E318K) exhibited a 14-fold higher risk than controls for developing melanoma + RCC (Table 1). To investigate the effect of Mi-E318K on susceptibility to melanoma alone, we genotyped 603 affected patients that had undergone oncogenetic testing for melanoma susceptibility genes¹ and were negative for *CDKN2A* and *CDK4* mutations (Table 1). Mi-E318K

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Figure 1 | Mi-E318K mutation impairs MITF SUMOylation. a, MITF sequence data of the c.952G>A, p.E318K germline substitution (Mi-E318K). b, HEK293 cells were co-transfected with empty pcDNA3 vector (EV) or pcDNA3 encoding wild-type (Mi-WT) or mutant (K182R, Mi-E318K or K182R+E318K)

frequency was significantly higher in these patients than in controls $(P = 7.8 \times 10^{-5})$. The carriers of Mi-E318K in this group exhibited >4-fold higher melanoma risk than controls (Table 1). In addition, Mi-E318K co-segregated with melanoma in three melanoma-prone families that had DNA available in relatives of index cases carrying Mi-E318K (Supplementary Fig. 1a). We also found that Mi-E318K was more strongly associated with multiple occurrences of primary melanomas than with single occurrence, irrespective of family history (P = 0.02, Supplementary Table 1). To investigate the effect of Mi-E318K in RCC susceptibility, we genotyped 164 patients with RCC referred to oncogenetic clinics, who were enriched for rare histological subtypes and who were wild type for the known RCC-predisposing genes⁶. These patients also exhibited a higher frequency of Mi-E318K than controls (P = 0.008) (Table 1). After correcting for multiple testing, the association between Mi-E318K and melanoma + RCC, melanoma only or RCC only remained significant, although marginally so for the RCC-only group (P = 0.02). The three groups showed similar Mi-E318K allele frequencies (P = 0.10). When all 829 patients with melanoma and/or RCC were pooled, Mi-E318K frequency was significantly different from controls ($P = 1.2 \times 10^{-6}$). The pooled group of Mi-E318K carriers had a greater than fivefold risk of developing melanoma, RCC or both cancers, as compared to controls (Table 1). The clinical and pathological features of the 27 Mi-E318K carriers are described in Supplementary Table 2. Principal component analysis (PCA) of single nucleotide polymorphism (SNP) data across the genome in more than 75% of cases and controls showed appropriate clustering of cases and controls after exclusion of a few subjects of non-European ancestry (Supplementary Fig. 1b). Association analysis of melanoma and/or RCC with Mi-E318K provided similar results when performed with or without adjusting for principal components (Supplementary Table 3). To assess whether Mi-E318K predisposed to the coexistence of melanoma and a second primary neoplasm other than RCC, we gathered data from European countries, but these data were insufficient for separate analyses of each type of cancer. Overall, we concluded from our findings that Mi-E318K is a rare substitution that confers an increased risk for developing melanoma and/or RCC. However, given the selection of patients who were part of case series undergoing genetic testing, further investigation of Mi-E318K in large series of unselected sporadic renal cancer and melanoma cases is merited.

Myc-tagged MITF and either with an empty vector or His-SUMO1 pSG5 vectors. Western blot analysis with antibodies to MITF, SUMO1 and ERK2 used as a loading control. c, Western blot analysis with antibodies to MITF and ERK2 of protein extracts before (Input) and after Ni-NTA affinity purification.

Previous studies have shown that MITF is expressed in a majority of melanomas and in a subset of kidney tumours9. Accordingly, our immunohistochemical analyses showed that MITF was expressed in both melanomas from patients carrying the germline mutation Mi-E318K (n = 8) and in melanomas from wild-type patients (n = 8)(Supplementary Fig. 2a, b). In addition, two out of seven RCCs from Mi-E318K carriers were positive for MITF (Supplementary Fig. 2c, d), suggesting that Mi-E318K might have a role in renal transformation. None of the six wild-type MITF RCC samples analysed in this study showed MITF labelling.

Mi-E318K occurs at a conserved position in MITF within a consensus motif (IKQE) that matches perfectly with the consensus sequence ΨKXE for covalent binding of SUMO¹⁰ (Supplementary Fig. 3a, b). Because SUMOylation is critically dependent on the acidic residue at +2 (E) of the acceptor lysine (K), we tested whether Mi-E318K affects MITF SUMOylation. Co-expression of histidine-tagged (His)-SUMO1 with wild-type MITF and western blot analysis revealed a 120-kDa band and a doublet above 95 kDa, in addition to the 55-65 kDa native doublet (Fig. 1b). The p.K182R mutation introduced within the second MITF SUMOylation site led to complete disappearance of the 120-kDa MITF form, but had little effect on the 95-kDa doublet. When glutamic acid residue 318 was replaced by a lysine, mimicking the germline mutation, we observed a strong decrease in the levels of all the SUMOmodified forms of MITF. Lastly, no SUMOvlated forms were seen with the double-mutant [p.K182R;p.E318K]. Similar observations were made with co-expression of haemagglutinin-tagged (HA)-SUMO2 (Supplementary Fig. 4a), demonstrating that Mi-E318K affected both SUMO1 and SUMO2 modifications. Western blot analysis with anti-MITF antibody, after affinity purification of His-SUMO1-containing proteins on Ni-NTA columns, confirmed that the high molecular weight bands were indeed SUMO-modified MITF proteins and that Mi-E318K markedly reduced all the MITF SUMOylated forms (Fig. 1c). Altogether, these results demonstrated that codon 316 was the major SUMO acceptor site in MITF and that Mi-E318K severely impaired SUMO conjugation to MITF.

SUMOylation has been shown to orchestrate a variety of cellular processes, in part through the control of nucleo-cytoplasmic signal transduction. However, both Mi-E318K and wild type were detected

Table 1 | Frequency of the germline Mi-E318K substitution

Subjects	Number of non-carriers	Number of carriers‡	Total	Frequency of p.E318K	FET P value†	OR (95%CI)*
Controls	1,649	10	1,659	0.003	-	Reference
Patients with melanoma and/or RCC	802	27	829§	0.016	1.2×10^{-6}	5.55 (2.59-12.91)
Melanoma + RCC	57	5	62	0.040	1.3×10^{-4}	14.46 (3.74–48.04)
Melanoma only	586	17	603	0.014	7.8×10^{-5}	4.78 (2.05–11.75)
RCC only	159	5	164	0.015	0.008	5.19 (1.37–16.87)

^{*} OR (95% CI) is the odds ratio (with 95% confidence interval) associated with the Mi-E318K carrier status

[†] FET, Fisher's exact test for the difference in Mi-E318K allele frequency between each group of patients and controls

[‡] All carriers are heterozygotes for the Mi-E318K substitution. Clinical and pathological features of the 27 carriers are described in Supplementary Table 2.

[§] A full description of the 829 patients is given in Methods

^{||} The Fisher's exact test showed no significant differences (P = 0.10) in allele frequency among the three groups of patients (melanoma + RCC, melanoma only, RCC only).

by immunofluorescence in the nucleus of transfected melanoma cells. This indicated that Mi-E318K did not affect MITF nuclear localization (Supplementary Fig. 4b, c).

Previous reports indicated that SUMOylation of MITF repressed its transcriptional activity, particularly when the targeted promoter contained multiple E-boxes¹⁰. Indeed, on a synthetic reporter gene comprising three E-boxes, Mi-E318K showed two- to threefold greater transcriptional activity than that of wild type (Fig. 2a). The double MITF mutant, [p.E318K;p.K182R], was more effective than Mi-E318K, and the single p.K182R mutant exhibited activity comparable to that of wild type. We next tested whether Mi-E318K affected MITF activity on physiological target promoters, such as MET and HIF1A, two MITF-regulated genes involved in cell survival and in melanocyte and kidney tumorigenesis. We also tested *CDKN2A* and *TYR* promoters, which are involved in melanocyte proliferation and differentiation processes, respectively. Wild type and mutant Mi-E318K showed similar activity on the MET (Fig. 2b), TYR and CDKN2A promoters (Supplementary Fig. 5), but Mi-E318K more efficiently activated the HIF1A promoter than wild type (Fig. 2c). This observation suggests that Mi-E318K alters MITF transcriptional activity on a subset of its target genes. To determine the global transcriptional effects of Mi-E318K, we performed pan-genomic expression profiling of A375 melanoma and RCC4 cells infected with adenoviruses that encoded either wild-type MITF or Mi-E318K (Supplementary Fig. 6a, b, c, d). Analysis of the genes regulated differentially by wild type and Mi-E318K in

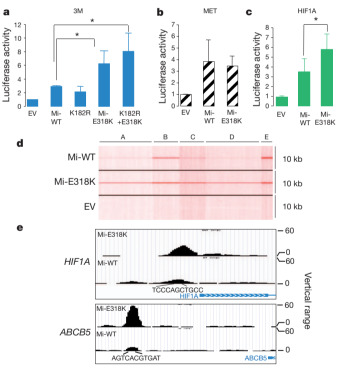


Figure 2 | Mi-E318K increases MITF binding and transcriptional regulation of a subset of its target genes. a-c, Reporter plasmids containing the luciferase gene under the control of different MITF target promoters were co-expressed with empty pcDNA3 (EV) or pcDNA3 encoding wild-type (Mi-WT) or mutant (K182R, Mi-E318K, K182R+E318K) MITF. a, HEK293 cells were transfected with a 3M-box reporter. b, c, 501mel human melanoma cells were transfected with the *c-MET* promoter reporter (**b**) or the *HIF1A* promoter reporter (c). The results were expressed as the fold stimulation over basal luciferase activity. Data represent the mean + s.d. of three independent experiments. Asterisk indicates a statistically significant difference (Student's *t*-test, $P \le 0.05$). **d**, MITF-occupied loci in genomic DNA extracted from $501 mel \, melanoma \, cells$ expressing Mi-WT or Mi-E318K. MITF tag density was compared in the region of ± 5 kb around the MITF-occupied loci. **e**, University of California Santa Cruz (UCSC) view of MITF occupancy of HIF1A and ABCB5 loci. The blue bar with arrowheads below each panel indicates the transcription start site; the E-box nucleotide sequence is shown under peaks.

A375 cells failed to identify a specific Mi-E318K signature. However, in RCC4 cells, we identified a Mi-E318K signature composed of 32 genes (Supplementary Table 4). On the basis of the bioinformatic analysis using Ingenuity Pathways Analysis (IPA) software, these genes were associated with cell growth, proliferation and inflammation (Supplementary Fig. 7). Among the genes downregulated by Mi-E318K compared to wild type, IRAK2 (ref. 11), EGR1 (ref. 12) and *IL6* (ref. 13) have tumour suppressor functions. The upregulated genes included CCR7, a HIF-1α target gene¹⁴ and activator of the NFκB pathway¹⁵; ABCB5, one of the most well-documented markers for melanoma initiating cells¹⁶; GADD45G, a member of the GADD45 family of stress sensors, which connects NFkB to the MAP kinase pathway¹⁷; and TRIM63, which was one of the top five upregulated genes and was the major gene found to be differentially expressed in patients with TFE3/TFEB translocation RCC as compared to other subtypes of RCC8. These experiments confirmed the different transcriptional potentials of wild type and Mi-E318K, particularly on genes involved in proliferation and inflammation.

To investigate whether SUMOylation influenced genome-wide MITF occupancy of its target sites, we performed chromatin immunoprecipitation coupled to high-throughput sequencing (ChIP-seq) experiments in 501mel human melanoma cell lines expressing wild type or Mi-E318K. Data analysis shows that 22,157 sites are occupied by Mi-E318K, whereas only 9,107 were detected for wild type, the difference being the presence of a large number of weakly occupied sites in the Mi-E318K data set. Comparative analysis of the ChIP-seq data sets using read density matrix clustering identified sites that are occupied by both wild type and Mi-E318K (Fig. 2d, sets B and E), whereas some promoter sites are uniquely or preferentially occupied by Mi-E318K (Fig. 2d, sets A and D; Supplementary Table 5). For example, a higher occupancy of the HIF1A promoter by Mi-E318K compared to wild type is observed (Fig. 2e). Mi-E318K also showed increased binding to the ABCB5 locus (Fig. 2e), the expression of which is increased in RCC4-expressing Mi-E318K. Of note, Mi-E318K binds more efficiently than wild type to the *HMOX1* promoter (Supplementary Table 5), a gene involved in both kidney cancer¹⁸ and melanoma cell growth¹⁹. Although the HIF1A promoter was better occupied and activated (in a gene promoter assay) by Mi-E318K than by wild type, we observed equal amounts of HIF1A messenger RNA in Mi-E318K- or wild-type-infected A375 and RCC4 cells. HIF1A transcripts are subjected to post-transcriptional regulation that controls their stability²⁰; this may mask the transcriptional effect of Mi-E318K. Alternatively, the cellular context or the chromatin landscape in RCC4 or A375 cells may not permit increased transcription. Indeed, this was observed for MITF target genes such as TYR, TYRP1 and DCT, which were not upregulated by MITF overexpression in A375 cells. Taken together, our data indicate that the naturally occurring Mi-E318K severely impaired MITF SUMOylation and showed higher global transcriptional activity, in agreement with the currently accepted model of SUMO-mediated transcriptional repression²¹. The global increase in Mi-E318K-occupied loci coupled with the existence of sites exclusively bound by the mutant protein indicate that SUMOylation regulates the repertoire of MITF target genes. SUMOylation-deficient Mi-E318K protein may therefore result in the regulation of distinct sets of genes, hence leading to gain-of-function properties.

To evaluate the tumorigenic potential of Mi-E318K, we tested its effect on migration, invasion and colony formation of stable melanoma cells (501mel) expressing wild-type MITF or Mi-E318K (Fig. 3a). We observed increased migration (Fig. 3b) and invasion (Fig. 3c) in two different Mi-E318K clones compared to wild-type clones. Similar results were obtained in VHL-deficient RCC4 cells (Fig. 3d, e) infected with adenoviruses encoding wild type or Mi-E318K. In contrast, Mi-E318K caused barely significant effects in A375 cells (used also for expression profiling) (Supplementary Fig. 8). Lastly, in a colony-forming assay with immortalized murine melanocytes, Melan-a (Fig. 3f), and RCC4 cells (Fig. 3g), the number of colonies after transfection of

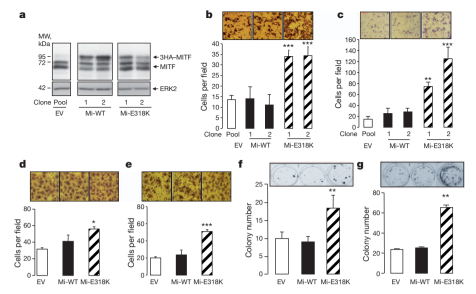


Figure 3 | Mi-E318K-enhanced migration, invasion and clonogenicity of melanoma and renal cancer cells. a, Western blot analysis with antibodies to MITF or ERK2 of 501mel cells stably transfected with empty vector (EV), or vector encoding Mi-WT or Mi-E318K (clones 1 and 2). MW, molecular weight. b, c, Migration (b) and invasion (c) assays with 501mel melanoma cells described earlier. d, e, Migration and invasion (e) assays in RCC4 cells transduced with empty adenovirus (EV) or adenoviruses encoding either Mi-WT or Mi-E318K. b-e, Photographs above each bar graph show a representative field of the underside of filter after cell migration or invasion; magnification, ×200. f, g, Mi-E318K increased the number of colonies formed with Melan-a or RCC4 cells, respectively. Photographs of the colonies were taken at 2 weeks. \mathbf{b} - \mathbf{g} , Bars show the mean + s.d. of three replicate assays. **b**–**g**, Student's *t*-test compared Mi-WT to Mi-E318K. *P < 0.05, **P < 0.01 and ***P < 0.001.

Mi-E318K was enhanced compared to wild-type-transfected or control cells. Overall, these data indicated that Mi-E318K was more potent than wild type for promoting invasive and tumorigenic behaviours in melanoma and RCC cells. However, Mi-E318K did not significantly stimulate melanoma or RCC cell growth (data not shown). These features are reminiscent of melanoma cell populations with increased invasive and division potential but with a slow growth rate, which are considered to be melanoma-initiating cells²². Thus, Mi-E318K might favour a phenotypic switch of melanoma cells towards a tumour-initiating cell phenotype, possibly in synergy with hypoxia²³.

Indeed, both the epidermis (where melanocytes are located) and the inner renal medulla are physiologically hypoxic tissues^{24,25}. Hypoxia is capable of rapidly inducing HIF1A to initiate a cell-survival response²⁶. Of note, several genes identified by expression profiling or ChIP-seq (HIF1A, CCR7, HMOX1) function in the hypoxia pathway. Both ultraviolet exposure and hypoxia generate radical oxygen species (ROS). In turn, oxidative stress activates inflammatory pathways that can lead to cellular transformation, proliferation and stem cell survival, among other features²⁷. Our IPA analysis associated the Mi-E318K signature with inflammation, cell proliferation and cancer. Various environmental stresses, including hypoxia and ROS, were previously shown to induce global protein SUMOylation²⁸. In this context, Mi-E318K, which prevents MITF SUMOylation, could impair the adaptation of cells to stress and initiate tumour formation. Our study has wide implications for understanding the role of MITF and its SUMOylation in the physiology and tumorigenesis of melanocytes and kidney cells.

We have identified a rare oncogenic germline substitution, Mi-E318K, that predisposes to both melanoma and RCC. Interestingly, MITF loss-of-function mutations are responsible for an inherited disorder in neural crest cell development, the type 2a Waardenburg syndrome⁷. This is reminiscent of the RET oncogene, in which activating germline mutations predispose to medullary thyroid carcinoma, but loss-of-function germline mutations predispose to Hirschsprung's disease, a congenital absence of enteric neurons in the gastrointestinal tract²⁹. Our data indicate that mutation screening in genetically enriched patients is a powerful strategy to identify rare genetic variation that confers a moderate risk of cancer³⁰. Lastly, it highlights the relevance of hereditary tumour models for shedding light on cell-growth-related signals and identifying cancer driver genes.

METHODS SUMMARY

Participants were included with the approval of an institutional review board. Sanger sequencing of *MITF* isoforms was performed in 62 melanoma + RCC cases. Mutations numbering referred to NCBI accession NM_000248.3. Wild-type or Mi-E318K genotyping was performed in 1,659 controls and 829 cases by TaqMan; positive samples were checked by Sanger sequencing. Genome-wide

SNP data were genotyped by the Centre National de Génotypage (CNG, CEA) using Illumina technology. PCA was applied to the SNP data using EIGENSTRAT software. Immunohistochemistry was performed on melanoma and kidney fixed samples with anti-MITF antibody (Abcam, clone D5). Mutations in the two SUMO1 fixation sites, p.K182R and p.E318K, were generated with the QuickChange method in wild-type MITF-M and MITF-A isoforms and verified by sequencing. Plasmids were transfected into HEK293, 501mel, Melan-a and RCC4 cells with FuGENE 6 reagent (Roche Applied Science). Western blotting and immunofluorescence were performed with antibodies to MITF (Abcam, clone C5), HA-tag, SUMO1 or ERK2 (Santa Cruz Biotechnology). Reporter assays were performed by transient transfection in triplicate with lipofectamine (Invitrogen), and pCMVBGal was included to control transfection efficiency. Gene expression microarray assays were performed with Agilent technology by the Institut de Cancérologie Gustave Roussy (IGR) genomics platform. ChIP-seq experiments were performed by the Institut de Génétique et de Biologie Moléculaire et Cellulaire (IGBMC) microarray and sequencing platforms. Colony forming assays were performed after 14 days of culture; cells were fixed, stained with 0.4% crystal violet, and photographed. Statistical analyses for comparison of allele frequencies between cases and controls were performed with Fisher's exact test. All computations were performed with Stata software, version 11. For functional assays, statistical significance was evaluated with the Student's t-test. Results were considered significant when the Student's *t*-test, **P* value \leq 0.05, ***P* \leq 0.01, ****P* \leq 0.001.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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profiling analysis. M.-F.A. initiated the collection of melanoma and RCC cases. S.R. initiated the collection of RCC families. M.-F.A and F.D. initiated the MELARISK collection. H.M. and V.C. contributed to the management of the MELARISK database.

Author Information Genome data has been deposited at the European Genome-Phenome Archive (EGA; http://www.ebi.ac.uk/ega), which is hosted at the EBI, under accession number EGAS0000000048. Gene expression data related to this paper have been submitted to the Array Express repository at the European Bioinformatics Institute (http://www.ebi.ac.uk/arrayexpress/) under the accession number E-TABM-1198. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of this article at www.nature.com/nature. Correspondence and requests for materials should be addressed to B.B.-d.P. (brigitte.bressac@igr.fr).

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METHODS

Study participants. Patients with melanoma and RCC. Sixty-two patients that had developed both melanoma and RCC were included in the MELARISK collection at the Institut de Cancérologie Gustave Roussy (IGR) and other French hospitals, 71% were males. None of these patients carried a germline *CDKN2A*, *CDK4* or *VHL* pathogenic mutation.

Patients with melanoma only. The patients with melanoma were enrolled through a nationwide network of French dermatology and oncogenetic clinics that constituted the Familial Melanoma Project and the MELARISK collection. The present study investigated a total of 371 index cases from independent pedigrees with a family history of melanoma (at least two melanoma cases) and/or pancreatic cancer, and 232 sporadic cases diagnosed with multiple primary malignant melanomas. A subset of the families (N=34) included a melanoma patient and relatives with pancreatic cancer. These were part of the CDKN2A mutation testing procedure; carriers of the CDKN2A mutation have high risk of developing pancreatic cancer³¹. All cases had confirmed diagnoses of malignant melanoma through medical records, review of pathological material, and/or pathological reports. For all these patients, mutation screening for CDKN2A (exon 1 β , 1 α , 2 and 3) and CDK4 (exon 2) had been undertaken for molecular diagnosis purposes. None of these patients carried a known CDKN2A or CDK4 pathogenic mutation.

Patients with RCC only. Overall, 164 patients included in this study had developed RCC only. Those patients had been recruited through French urology and oncogenetic clinics, within the French National Cancer Institute's 'Inherited Predisposition to Kidney Cancer' network. They were considered 'genetically enriched' based on the following criteria: familial aggregation, young age of onset and rare histological subtypes. Out of the total of 164 patients, 79 had sporadic ccRCC, 54 had papillary renal cell carcinoma (PRCC; with 6 type I PRCC, 19 type II PRCC, 1 mixed type PRCC and 28 unknown histological subtypes of PRCC), 1 had a mixed RCC phenotypes (that is, papillary and ccRCC), 5 had juvenile RCC and 25 had RCC of unknown histological subtype. None of the 79 patients with ccRCC carried a *VHL* pathogenic germline mutation; none of the 19 patients with PRCCII carried a *FH* germline mutation; and none of the 6 patients with PRCCI carried a *MET* germline mutation.

Control subjects. Controls were supplemented by a population-based sample of 1,659 French subjects that had participated in the Supplementation in Vitamins and Mineral Antioxidants (SU.VI.MAX) study³². We confirmed that these subjects had not developed any cancer at the time of the present study.

The study was approved by an institutional review board (CCPPRB no. 01-09-05, Paris Necker) for the MELARISK collection and by the Ethical Committee of Le Kremlin-Bicêtre University Hospital, for the kidney cancer collection. It was conducted according to the Declaration of Helsinki Principles. All participating subjects signed informed consent and provided blood samples. After informed consent had been obtained from the subjects, DNA was extracted from peripheral blood lymphocytes with the QIAamp DNA Blood mini kit (QIAGEN), according to manufacturer guidelines.

<code>MITF</code> gene sequencing. Sanger sequencing of <code>MITF</code> coding sequence, the intronexon boundaries and the 8 alternative promoters was performed in DNA extracted from the blood of 62 patients affected with melanoma + RCC (primers available upon request) on a 3730 DNA Analyser (Applied Biosystems; ABI). Nucleotide number refers to the wild-type cDNA sequence of <code>MITF</code> (NM_000248.3) as reported in GenBank.

Genotyping of Mi-E318K. Genotyping was carried out with Taqman according to the manufacturer recommendations. Primers and probes were supplied in the Assay-by-Design by Applied Biosystems (ABI). PCR reactions were performed with 10 ng genomic DNA and 0.2 μmol l⁻¹ TaqMan MGB probes. Probe 5′-VIC-ATCAAGCAAGAACCCG-3′ was designed to match the wild-type allele, and probe 5′-6-FAM-CAAGCAAAACCCG-3′ was designed to match the mutant allele (underlined bold indicates the nucleotide at the mutation site). PCR thermocycling was performed on ABI thermocyclers, as follows: 95 °C for 15 min; 30 cycles of 95 °C for 15 s and 60 °C for 1 min. Assays were carried out in 96-well plates that included a negative control (with no DNA) and positive controls (DNA from subjects with Mi-E318K). Plates were read on a 7900HT Fast Real-Time PCR System (ABI) with Sequence Detection Software (ABI). Carriers of Mi-E318K were confirmed by Sanger sequencing with the above protocol and primers for *MITF* exon 9.

PCA of genome-wide SNP data. To verify that the association of Mi-E318K with melanoma and/or RCC was not influenced by population stratification, we carried out PCA of SNP data across the genome in cases and controls that had been genome-wide genotyped and satisfied stringent quality control criteria. A total of 1,628 controls and 569 cases affected with melanoma, RCC or both cancers were genotyped by the Centre National de Génotypage (CNG) using Illumina HumanHap300 Beadchip version 2 duo array for controls and Illumina Humancnv370k and Human660W-Quad arrays for cases. Samples were excluded

for any of the following reasons: (1) a call rate of less than 97% of the total number of SNPs on the chip (222 samples); (2) sex as ascertained by genotyping not matching reported sex (10 samples); (3) heterozygosity on autosomes departing from the estimated expected value (7 samples); (4) relatedness with another sample (4 samples). This resulted in exclusion of 243 samples. PCA was then applied to 1,954 subjects that passed quality control using EIGENSTRAT software³³. To identify individuals of non-European ethnicity, SNPs were thinned to reduce linkage disequilibrium and combined with the HapMap data of wide-ranging ethnicity. The first two principal components (PCs) clearly separated the HapMap data into distinct clusters according to ethnicity and identified 29 study samples of non-European ancestry who were excluded. The remaining 1,925 European ethnicity samples (1,389 controls, 536 cases) were analysed similarly without the Hapmap data. Plotting the first two principal components showed appropriate clustering of cases and controls (Supplementary Fig. 1).

MITF immunohistochemistry. Briefly, 4-µm tissue sections were cut from paraffin-embedded blocks, deparaffinized and rehydrated. RCC (patient ID numbers 10276, 24976, 11473, 21309, 21939, 25220, 26534; all in Supplementary Table 2) immunohistochemistry was performed with a three-step, avidin-biotin-peroxidase method. Staining was performed in an automated Dako Autostainer. Heat-mediated antigen retrieval was performed in 0.1 mol l⁻¹ pH 6.0 citrate, and samples were heated in a water bath for 30 min in a Dakolink processor (Dako). Monoclonal mouse anti-human MITF (Clone D5, Dako, 1/100) was incubated with the samples for 30 min at room temperature (22 °C). SSM immunohistochemistry (patient ID numbers 19525, 24976, 10254, 15168, 15012, 21000, 22112, 27708 all in Supplementary Table 2 except number 27708 in Supplementary Fig. 1) was performed with a three-step avidin-biotin-alkaline-phosphatase method. Staining was performed in a Roche Diagnostics Benchmark XT automated stainer. Heatmediated antigen retrieval was performed in the automat in cell conditioning solution 1 (Roche Diagnostics) for 1 h (Dako). Monoclonal mouse anti-Human MITF (Clone D5, Roche Diagnostics, pre-diluted) was incubated with samples for 16 min at room temperature. All slides were then counterstained with haematoxylin. Appropriate negative and positive controls were prepared in parallel.

Functional assays. Plasmids. The pcDNA3-Mi construct, which carried the M-MITF isoform and the 3M vector have been described previously³⁴. The A-MITF isoform was a gift from H. Arnheiter (NIH). The *MITF* mutations, p.K182R and/or Mi-E318K, were generated with the QuickChange method (Stratagene) with the following sense primers and their reverse complements: Mi-K182R 5'-CTTCCCAACATAAGAAGGGAGCTCACAGC-3'; MI-E318K 5'-GGATCATCAAGCAAAAACCAGTTCTTGAG-3'. Mutations were confirmed by DNA sequencing. His-SUMO1 and HA-SUMO2 were a gift from A. Dejean and have been described elsewhere³⁵.

Cell cultures, transfections, and immunoblots. Human 501mel and A375 melanoma cells, human RCC4 cells and HEK293 cells were grown in DMEM supplemented with 7% FBS. Melan-a cells were cultured in RPMI 1640, 7% FCS, 200 nM TPA and 200 pM cholera toxin. All the cells were maintained at 37 $^{\circ}$ C in a humidified atmosphere containing 5% CO $_2$. HEK293 cells were cultured in 6-well dishes (10 4 cells per well) and transfected with the above-mentioned plasmids (2 μg of total DNA per well) and FuGENE 6 (Roche Applied Science). Forty-eight hours later, the cells were washed with PBS and lysed at 95 $^{\circ}$ C in 1× loading buffer (41.6 mM Tris, pH 6.8, 1.5% SDS, 6.7% glycerol). Proteins were resolved by electrophoresis in 10% SDS–polyacrylamide gels and transferred to PVDF membranes. Proteins were detected with ECL (Amersham) and antibodies to MITF (Abcam), HA-tag (Abcam), SUMO1 (Santa Cruz Biotechnology) or ERK2 (Santa Cruz Biotechnology).

Reporter assays. Human 501mel melanoma cells and HEK293 cells were seeded in 24-well dishes $(25\times 10^3$ cells per well). Subsequently, cells were transiently transfected with 0.3 µg reporter plasmid (3M, pHIF1A, pMET), 0.05 µg of MITF-encoding plasmids or empty, control pcDNA3 (EV), 2 µl of lipofectamine reagent (Invitrogen) and 0.05 µg of pCMV β Gal for controlling the variability in transfection efficiency. Cells were lysed 48 h later and assayed for luciferase and β -galactosidase activities. Transfections were repeated at least three times.

Immunofluorescence. HEK293 cells were seeded on glass coverslips (100×10^3 cells) in 6-well dishes, and transfected with 3 µg of the different MITF mutants and 10 µl lipofectamine. Forty-eight hours later, cells were fixed for 10 min with 4% paraformaldehyde, permeabilized for 2 min with 0.1% Triton X-100/1% bovine serum albumin (BSA), and treated for 2 min with NH $_4$ Cl 50 mM . Then, samples were washed three times in PBS and stained for 1 h with monoclonal anti-MITF antibody (Abcam, clone D5) in 1% BSA/PBS. Next, samples were washed three times in PBS for 5 min each and then stained secondarily for 1 h with Alexa-594-conjugated goat-anti-mouse antibody (Molecular probes) in 1% BSA. Cells were counterstained with 4,6-diamidino-2-phenylindole (DAPI), mounted with Fluoromount-G (Southern Biotech), and examined with a Zeiss Axiophot microscope equipped with epifluorescent illumination.

Colony forming assay. Immortalized Melan-a mouse melanocytes, A375 human melanoma cells and RCC4 human cells (8 \times 10 4 cells per well) were transfected with wild type or Mi-E318K (3 µg total DNA per well and 10% pBABE-puro) with the FuGENE 6 Transfection reagent (Roche Applied Science). Puromycin (1 µg ml $^{-1}$) was added to media at 48 h post-transfection. Fourteen days later, the cells were fixed and stained with 0.4% crystal violet, and the plates were photographed.

Establishing stable 501mel cells that expressed tagged MITF. 501mel cells were transfected with FuGENE 6 reagent, a vector encoding puromycin resistance, and a pCMV vector that was either an empty, control vector or a vector encoding 3×HA-tagged MITF (wild type or Mi-E318K). Transfected cells were selected with puromycin; the expression of MITF was verified by western blot analysis with anti-MITF (Abcam, C5) or anti-HA (Roche, 12CA5) antibodies.

Migration and invasion assays. Migration (on uncoated filters) and invasion (on coated filters with matrigel) were investigated in a Boyden chamber system that comprised 24-well plates and 8- μm pore filter inserts (BD Bioscience). Stable 501mel melanoma cells $(2\times10^5$ cells), A375 melanoma cells $(1.5\times10^5$ cells) and RCC4 renal cancer cells $(5\times10^4$ cells) were infected with control vector or adenovirus that encoded wild type or Mi-E318K at a multiplicity of infection (MOI) of 20 for 24 h, were then resuspended in serum-free DMEM and seeded on the upper chamber inserts. DMEM with 7% FCS was placed into the lower chamber. Twenty-four hours (501mel and RCC4) or 6 h (A375 cells) later, cells adherent to the underside of the filters were fixed with 4% PFA and stained with 0.4% crystal violet. Adherent cells were counted in five random fields at $\times 200$ magnification. Results represented the average of duplicate samples from three independent experiments.

Gene expression arrays. Three replicates were performed for both RCC4 and A375 cells. mRNA was isolated with Trizol (Invitrogen) from A375 melanoma cells and from RCC4 cells infected with either control, wild type or Mi-E318K, according to standard procedures. Briefly, probes were synthesized from 500 ng of total RNA in two steps, according to the manufacturer's instructions. The two samples to be compared were labelled separately with different fluorescent dyes, cyanine-3 (Cy3) and cyanine-5 (Cy5). For each sample, 1 µg of purified cRNA labelled in cv5 was mixed with the same amount of cRNA labelled in cv3. Label incorporation was checked on a NanoDrop spectrophotometer. Hybridizations were performed with a dye-swap strategy on whole-human-genome dual colour 8 × 60K oligonucleotide microarrays (design 028004; Agilent Technologies). Feature extraction software provided by Agilent (version 10.7.3.1) was used to quantify the intensity of fluorescent images and to apply a linear/lowess normalization to correct for artefacts caused by nonlinear rates of dye incorporation and inconsistent relative fluorescence intensities between some green and red dyes. All data were imported into Resolver software (Rosetta Biosoftware) for database management, quality control, computational re-

combination of dye-swaps and statistical analysis. Mi-E318K specific signature was generated using the following parameters: intensity >50; fold change >1.5; P value <0.05. Functional analysis was carried out with the Ingenuity Pathway Analysis (Ingenuity System, http://www.ingenuity.com).

ChIP-seq. ChIP-seq was performed as previously described³⁶. Briefly, chromatin was prepared from native 501mel cells or cells that stably expressed HA-tagged wild type or Mi-E318K. The isolated chromatin was HA-immunoprecipitated and sequenced with an Illumina GAIIx sequencer. The raw data were analysed with the Illumina Eland pipeline V1.6 programme. Peak detection was performed with MACS software (http://liulab.dfci.harvard.edu/MACS/), and the peaks were annotated with GPAT software (http:// bips.u-strasbg.fr/GPAT/Gpat_home.html). Peak annotations were performed with a window that included $\pm 20\,\mathrm{kb}$ from the coordinates at the beginning and end of RefSeq transcripts. The total number of reads for the wild-type data set was 1.7 fold higher than for the Mi-E318K data set. To facilitate quantitative comparisons, an appropriate number of reads was randomly removed from the wild-type data set to match the number present in the Mi-E318K data set. Subsequently, a quantitative comparison of the ChIP-seq data sets was performed with seqMINER³⁷. Clustering was performed by counting the number of tags in a 36bp sliding window for each ChIP-seq data set based on the coordinates of wild-type or Mi-E318K binding sites or on a reference sequence (RefSeq TSS). Only peaks with ≥10 reads were used for comparison. A matrix of binding sites and densities was generated and subjected to K-means clustering with the seqMINER programme.

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CORRECTIONS & AMENDMENTS

CORRIGENDUM

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Corrigendum: A SUMOylationdefective MITF germline mutation predisposes to melanoma and renal carcinoma

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In this Letter, one image was mistakenly duplicated during preparation of the artwork. In the original Fig. 3d, the left image illustrating migration of RCC4 cells transduced with empty adenovirus (EV) at 24 h is a duplicate of the middle image showing migration of RCC4 cells transduced with an adenovirus encoding Mi-WT. The corrected images and migration graph are shown in Fig. 1 of this Corrigendum. This correction does not alter any of the conclusions, and the authors apologize for any confusion this may have caused. *Nature* has not received a response from the following authors to approve this Corrigendum: V.M., T.M.-D., A.Ro., P.B., E.C. and V.C., and C. Becuwe., J.-L.B., J.C.-B., S.D., C.D., J.L., and K.M. from The French Familial Melanoma Study Group (L.D. is deceased).

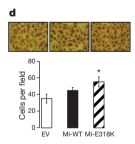


Figure 1 | This is the corrected Fig. 3d of the original Letter.