Role of BRAFV600E in the First Preclinical Model of Multifocal Infiltrating Myopericytoma Development and Microenvironment


*Authors contributed equally to this work.

Manuscript received December 23, 2013; revised May 14, 2014; accepted May 20, 2014.

Correspondence to: Carmelo Nucera, MD, PhD, Harvard Medical School, Laboratory of Human Thyroid Cancers Preclinical and Translational Research, Division of Cancer Biology and Angiogenesis, Department of Pathology, Beth Israel Deaconess Medical Center, Harvard Medical School, 99 Brookline Avenue, Boston 02215, MA, USA. (e-mail: cnucera@bidmc.harvard.edu).

Myopericytoma (MPC) is a rare tumor with perivascular proliferation of pluripotent stem-cell–like pericytes. Although indolent, MPC may be locally aggressive with recurrent disease. The pathogenesis and diagnostic biomarkers of MPC are poorly understood. We discovered that 15% of benign MPCs (thyroid, skin; 3 of 20 samples) harbored BRAFV600E; 33.3% (1 of 3 samples) of BRAFV600E-MPCs were multifocal/infiltrative/recurrent. Patient-MPC and primary MPC cells harbored BRAFV600E, were clonal and expressed pericytic-differentiation biomarkers crucial for its microenvironment. BRAFV600E-positive thyroid MPC primary cells triggered in vitro (8.8-fold increase) and in vivo (3.6-fold increase) angiogenesis. Anti-BRAFV600E therapy with vemurafenib disrupted angiogenic and metabolic properties (~3-fold decrease) with down-regulation (~2.2-fold decrease) of some extracellular-matrix (ECM) factors and ECM-associated long non-coding RNA (LincRNA) expression, with no effects in BRAFV600E-MPCs. Vemurafenib also inhibited (~3-fold decrease) cell viability in vitro in and BRAFV600E-positive thyroid MPC patient-derived xenograft (PDX) mice (n = 5 mice per group). We established the first BRAFV600E-dependent thyroid MPC cell culture. Our findings identify BRAFV600E as a novel genetic aberration in MPC pathogenesis and MPC-associated biomarkers and imply that anti-BRAFV600E agents may be useful adjuvant therapy in BRAFV600E-MPC patients. Patients with BRAFV600E-MPC should be closely followed because of the risk for multifocality/recurrence.


Myopericytoma (MPC) describes rare, nodular tumors characterized by a radial and multilayered perivascular growth of ovoid and spindled-shaped cells with mesenchymal stem-cell–like features, often with associated blood vessels arranged in an irregular, “staghorn” pattern (1–3). Most MPCs are benign (2), but some are malignant with metastatic potential and poor survival (4,5). MPCs arise over a wide age range, primarily affecting subcutaneous tissues of the distal extremities (2), and lesions occasionally present with skin ulceration, pain or tenderness; additionally, symptomatic MPCs arise in the proximal extremities, head region and internal organs (1). Most MPC are treated surgically, and complete excision should prevent recurrent/persistent disease, although negative surgical margins are difficult to achieve outside of the extremities (2).

Biomarkers and oncogenic events driving MPC development are not well understood. In this study, we performed a comprehensive genomic and functional validation of associated MPC biomarkers with clinical implications. We have applied a high-throughput genotyping assay on 29 MPCs from available, formalin-fixed paraffin-embedded (FFPE), discarded/ unidentified samples, using an Institutional Review Board-approved protocol (Beth Israel Deaconess Medical Center, Boston, MA) to unravel oncogenic events driving pathogenesis (Supplementary Methods and Supplementary Table 1, available online). The patient with thyroid MPC provided written informed consent for genetic analysis. For all other patients, we used discarded/ unidentified tissue specimens and consent for genotyping test. Disease stage was assessed by radiologic imaging. We also used and immunohistochemistry, primary cell cultures, dynamic functional assays, shRNA, and developed an MPC-patient-derived xenograft (PDX) mouse model (for detailed methods, please see the Supplementary Methods, available online). All animal work was done in accordance with federal, local, and institutional guidelines at the Beth Israel Deaconess Medical Center (Boston, MA), and all experiments were performed with four-month-old Crl:NU(NCr)-Foxn1nu female, athymic, immunodeficient, nude mice (strain code: 490) (Charles River, Wilmington, MA) (n = 5 per group).

Statistical analysis was carried out using GraphPad Prism 6 software (version Prism 6, GraphPad Software Inc., San Diego, CA). Mann-Whitney test was used to analyze the statistical significance of differences between two groups. For categorical data, Fisher’s exact test was used. All reported P values were two sided. Data are reported as the averaged value, and error bars represent the standard deviation of the average for each group in duplicate or triplicate. Results with P values below .05 were considered statistically significant.

We developed the first translational model to date of multifocal and infiltrative thyroid MPC (Figure 1; Supplementary Figures 1 and 2, available online). The
Figure 1. A multimodality perioperative evaluation of patient with multifocal infiltrating thyroid myopericytoma harboring the heterozygous BRAF WT/V600E mutation. A1) Multinodular thyroid disease, 15 mm follicular variant papillary thyroid microcarcinoma (PTCFV) in the left thyroid lobe and a left thyroid mass (myopericytoma [MPC]) with mediastinal extension in a 44-year old woman. A2) Neck Doppler ultrasound shows a 4.0 x 3.4 cm irregular, hypo-echoic, hyper-vascular mass in the left thyroid lobe with mediastinal extension. A3) Contrast enhanced T1-weighted magnetic resonance imaging before thyroid surgery shows a four-cm, hyper-intense mass (arrowhead) localized in the left thyroid lobe with mediastinal extension, impingement, rightward tracheal (asterisk) deviation and possible extension into soft tissue (arrow highlights a right thyroid nodule). A4) Total body 18F-FDG (fluorodeoxyglucose) PET (Positron emission tomography)/CT (Computed Tomography) exam reveals high 18F-FDG uptake in the left persistent/recurrent mass (arrow). A5) Elastography analysis shows a non-deformable pattern (the arrow highlights a reduction of the elasticity, blue color) in the vast majority part of the left persistent/recurrent MPC. B1-B7) Histology of the thyroid and perithyroidal soft tissue with multifocal and infiltrating thyroid MPC (hematoxylin and eosin staining). (B1, scale bar = 500 µ) Macrol follicular thyroid tissue. (B2-B3, scale bar = 200 µ) The thyroid MPC is present in several foci (asterisk) in the thyroid gland. (B2-B3, arrowhead highlights thyroid follicles). (B4, scale bar = 500 µ) The MPC cells show cytoplasmic to membranous staining with antibodies against α-smooth muscle actin (αSMA) and platelet-derived growth factor receptor beta (PDGFRB). C5) MPC cells also show positivity (punctate) for NG2 (chondroitin sulfate proteoglycan 4) but are negative for desmin (C6). C7) Trichrome staining highlights the abundant amount of collagen deposition (blue staining, asterisk) surrounding the MPC (purple staining) or the intratumoral deposition of collagen (arrows). D1-D2) DNA genotyping analysis by mass spectrometry (MS) traces of formalin-fixed paraffin embedded thyroid (n = 1) (D1) and (D2) skin (eg, leg) (n = 28) MPC tissue samples reveals a heterozygous BRAF WT/V600E allele (long arrows) (A > T). The intensity of the signal vs mass of the analyte is plotted in the background, while the inset shows the cluster plot with all samples analyzed in the run (intensity of the signals of the two alleles). The long arrow links the actual sample (circle in the inset) to the mass of the expected mutant V600E allele. Color coded by the software is automatic allele calls. Cyan squares for heterozygous samples, orange triangles for homozygous. Ambiguous calls are identified by a red dot. Allele frequencies deviating from the expected values are assigned ambiguous or homozygous calls by the software.

initial pathologic diagnosis in October 2010, reported by the referring international institution, was undifferentiated (anaplastic) thyroid carcinoma with two consequent cycles of chemotherapy given to the patient. By patient request, outside review of the pathology was performed at the Massachusetts General Hospital (Boston) the following month, and the
diagnosis was amended to reflect a multifocal and infiltrative left thyroid MPC (Figure 1; Supplementary Figures 1 and 2, available online), with no clinically suspicious lymph nodes or distant metastases. Completion thyroidectomy was performed in June 2011 (Supplementary Figure 1F, available online). Histologic evaluation confirmed residual thyroid MPC (Supplementary Figure 1, A-D, available online).

The MPC was about two-fold more metabolically active (standardized uptake value [SUV] = 4.5) compared with the non-tumoral thyroid hyperplastic nodules (SUV = 2.4) (Figure 1A4), but 18-FDG (Fluorodeoxyglucose) PET (positron emission tomography)/CT (computed tomography) did not note any non-regional tissue involvement (Supplementary Figure 1D, available online). Twelve months post-operatively, imaging revealed persistent disease in the thyroid bed (Figure 1A5; Supplementary Figure 1, G-J, available online). MPC lacked features of malignancy (increased number of mitoses, necrosis, vascular invasion), showed apparent differentiation towards pericyte lineage (Figure 1, B2-B7) and was characterized by perivascular growth (Figure 1, B5 and B6, and 1C2). A higher vascular density (CD31+) was found in the thyroid MPC (mean vessels/field = 20.2, SD = 0.4 vessels/field) compared with the adjacent normal thyroid (mean = 5.2 vessels/field, SD = 0.2, P = .007) (Figure 1C2). MPC cells are arranged circumferentially around smaller vessels (Figure 1, B5 and B6, and 1C2; Supplementary Figure 2B5, available online). The ratio of MPC cells to endothelial cells is about 3:1, quantified based on the number of platelet-derived growth factor receptor beta (PDGFRβ)-positive MPC cells and CD31-positive endothelial cells (Supplementary Methods, available online). MPC biomarkers (alpha-smooth muscle actin [αSMA], PDGFRβ, NG2 [neuron-glial antigen 2], and extracellular-matrix [ECM] molecules, eg, CollagenIα1 [COLIa1]), as well as desmin and p16/Ink4A immunoexpression are described in Figure 1, C3-C7; Supplementary Figure 2, B1-B19, and Supplementary Table 2, available online.

As MPC is a poorly characterized “orphan” disease, we performed in-depth genotyping. We analyzed 29 FFPE tissues (20 benign MPC, four benign intravascular MPC, and five malignant MPC) (Supplementary Table 1, available online) by Mass Spectrometry genomic technology, which interrogates about 1000 mutations in 112 validated oncogenes and tumor suppressors. Results were validated by performing Sequenom (6) (Figure 1, D1-D2; and Supplementary Figure 2, C1-C4, available online), pyrosequencing and Sanger sequencing (data not shown). We found that 3/20 (15%) benign MPCs (ie, one thyroid and two cutaneous MPCs) from three different patients (Figure 1, D1-D2; Supplementary Figure 2, C1-C4, and Supplementary Table 1, available online) harbored the heterozygous BRAFWT/V600E mutation in exon-15 hot-spot T1799A of the BRAF gene sequence. None of the four benign intravascular MPCs or five malignant MPCs harbored BRAFWT/V600E. Two out of three (66.6%) BRAFWT/V600E-MPCs (thyroid, skin) derived from two different patients’ infiltrated adjacent soft tissue; two of four (50%) multifocal MPCs harbored BRAFWT/V600E, compared with 1/25 (4%) unifocal MPCs with wild type (WT) BRAF (P = .04) (Supplementary Table 1, available online). One out of three (33.3%) MPCs (ie, thyroid) with BRAFV600E showed recurrent/persistent disease after one year of follow-up (Figure 1A5; Supplementary Figure 1, G-J, available online). Furthermore, our analysis of X-chromosome inactivation and methylation profile from the female patient with thyroid BRAFV600E-MPC (Figure 1D1 and Figure 2, A1 and A2) revealed that this tumor was monoclonal compared to the adjacent, uninvolved thyroid tissue (Figure 2A3) (7).

To provide a translational application for our study, we established early and late passages of primary cells cultured in vitro from human thyroid BRAFWT/V600E-MPC. These cells also harbored BRAFWT/V600E and expressed pericyte lineage-specific differentiation biomarkers (Figure 2A1). BRAFWT/V600E is the most frequently mutated oncogenic kinase. Vemurafenib is the first orally available selective inhibitor of BRAFV600E approved by the US Food and Drug Administration for the treatment of BRAFWT/V600E-melanoma (8–11). We tested the effects of vemurafenib on BRAFWT/V600E-MPC cells and BRAFWT pericytes (Figure 2, A-G). Vemurafenib substantially reduced phospho(p)ERK1/2 and pMEK1/2 protein levels in BRAFWT/V600E-MPC cells as compared with controls (Figure 2C). As a result, this treatment statistically significantly (P < .001) suppressed BRAFWT/V600E-MPC cell viability, with no effect on the growth of BRAFWT-pericytes (Figure 2B), suggesting its high specificity for the BRAFV600E vs BRAFWT. Furthermore, we hypothesized that BRAFWT/V600E plays a role in MPC angiogenic and metabolic properties; we found that BRAFWT/V600E-MPC cells substantially grew as large cell aggregates on Matrigel (Supplementary Figure 3A, available online), and, when cocultured with human microvascular endothelial cells, statistically significantly (8.8-fold, P = .002) triggered in vitro angiogenesis as compared with controls (Figure 2, D1-D2). Vemurafenib treatment statistically significantly (about 3-fold, P = .002) disrupted this effect (Figure 2, D1 and D2); additionally, treatment with shRNA that targeted BRAFV600E statistically significantly reduced MPC cell adhesion and migration (Supplementary Figure 4C, available online) with no effect in BRAFWT-pericytes (data not shown). Remarkably, BRAFWT/V600E-MPC cells also statistically significantly (P = .002) increased vascular density in MPC-PDX mice (Supplementary Figure 5, available online), and vemurafenib therapy statistically significantly (3-fold, P = .007) suppressed both BRAFWT/V600E-MPC cell viability (Figure 2G; Supplementary Figure 5, available online) and vascular density/angiogenesis (3.6-fold, P = .002) (Supplementary Figure 5, available online) without any obvious toxicity. Subsequently, we found that COLIa1 (5-fold, P = .007), PDGFRB (1.3-fold increase, P = .02), integrin-β1 (ITGB1) (1.7-fold increase, P = .02), ID2 (DNA-binding protein inhibitor) (6.5-fold increase, P = .002), and the long intergenic non-coding RNA (LinRNA) ID2 (2.8-fold increase, P = .02) were statistically significantly (with moderate or high copy number) BRAFV600E-dependent in MPC cells as compared with control cells with BRAFVT (Figure 2F; Supplementary Table 3, available online). Their expression levels (ID2 = 2.2-fold; COLIa1 = 1.5-fold; PDGFRB = 1.5-fold; ITGB1 = 1.5-fold; and ID2 LinRNA = 1.6) were statistically significantly reduced by vemurafenib treatment as compared with BRAFWT-pericytes (Figure 2F). Furthermore, MPC cell adhesion (P = .002, Supplementary Figure 3B, available online) and migration (P = .002,
Figure 2. In vitro and in vivo preclinical model of human primary thyroid myopericytoma harboring the heterozygous \( \text{BRAF}^{WT/V600E} \) mutation. 

A1) Immunocytochemistry of established non-immortalized primary human thyroid myopericytoma (MPC) cells with the heterozygous \( \text{BRAF}^{WT/V600E} \) mutation. Phase image of MPC cells, scale bar = 100 \( \mu \). Immunohistochemistry staining (scale bar = 400 \( \mu \)) in the thyroid MPC cells show cytoplasmic to membranous staining with antibodies against \( \alpha \)-smooth muscle actin (\( \alpha \)-SMA), platelet-derived growth factor receptor beta (PDGFRB), and CD44 (stem cell and cell adhesion marker). Thyroid MPC cells also show positivity for NG2 (chondroitin sulfate proteoglycan 4), phospho(p)-ERK1/2, and vimentin. CD31, CD45, pan-keratin, desmin, and thyroglobulin (Tg) immunostains are negative.

A2) Exon 15, T1799A ACAG T/A GAAA

B) Cell viability assay

C) DNA genotyping analysis by mass spectrometry (MS) traces of the human primary thyroid MPC cells (n = 1) reveals a heterozygous \( \text{BRAF}^{WT/V600E} \) allele (long arrows) (\( A > T \)). The intensity of the signal vs mass of the analyte is plotted in the background, while the inset shows the cluster plot with all samples analyzed.
Figure 2E) were also statistically significantly reduced by vemurafenib treatment compared with BRAF\(^{WT}\)-pericytes. MPCs continue to be under recognized, resulting in inappropriate treatment and patient anxiety (12). Benign MPCs are generally indolent tumors (1); however, when they involve internal organs, they may be locally aggressive, with recurrent/persistent disease because of difficulty of complete excision. We discovered that BRAF\(^{V600E}\) is a distinct genetic alteration seen in benign MPCs, often multifocal and infiltrative. Patients with multifocal MPCs present a treatment dilemma, often requiring multiple surgeries for lesions that may be quite painful, even if not malignant. BRAF\(^{V600E}\) is a prognostic biomarker of tumor recurrence and aggressiveness (13–24) and also facilitates tumorigenesis (25–29).

BRAF\(^{WT/V600E}\)-induced MPC cell adhesion, migration, and angiogenesis associated with upregulation of molecules (eg, COL1A1, PDGFR\(B\), ID2) (30–33) crucial for ECM remodeling, angiogenesis, and for autocrine and paracrine communication in the tumor microenvironment (Supplementary Figure 6, available online), which ultimately may lead to MPC aggressiveness. High doses of vemurafenib therapy were effective to inhibit BRAF\(^{V600E}\)-MPC cell viability and angiogenesis, suggesting that this therapy blocked BRAF\(^{V600E}\)-dependent pro-migratory pathways and so diminished pro-angiogenic capabilities of MPC cells. Please see the Supplementary Results (available online) for additional findings that may be of interest.

Collectively, using anti-BRAF\(^{V600E}\) therapy as a surgical adjuvant may provide a novel advancement in the therapeutic strategy and treatment of locally aggressive multifocal BRAF\(^{V600E}\)-positive MPCs or possibly serve as a therapeutic alternative for cases in which surgical options are limited by location and extent of disease, or in medically poor surgical candidates. Our study is limited by sample size (29 available MPCs), precluding optimal evaluation of MPC pathological features with/without BRAF\(^{V600E}\). However, our results were substantiated by our integrated in vitro and in vivo approaches. It is also possible that MPC heterogeneity reduced the sensitivity for detection of the BRAF\(^{V600E}\) mutation in some cases; therefore, we cannot exclude the possibility that the percentage of BRAF\(^{V600E}\)-positive MPC cases is higher.

In conclusion, our results demonstrate a subset of MPC harbor BRAF\(^{V600E}\) that drives tumor development. We report the first MPC arising in the thyroid. Anti-BRAF\(^{V600E}\) therapy effectively suppresses viability in the only currently available MPC short-term cell culture that harbors the BRAF\(^{V600E}\) mutation. BRAF\(^{V600E}\) plays a role in the MPC microenvironment (Supplementary Figure 6, available online) and might ultimately lead to aggressive behavior. Finally, we report a multiplex panel of diagnostic markers for MPCs in the run (intensity of the signals of the two alleles). The long arrow links the actual sample (circle in the inset) to the mass of the expected mutant V600E allele. Color coded by the software is automatic allele calls. Cyan squares for heterozygous sequences, orange triangles for homozygous. Ambiguous calls are identified by a red dot. Allele frequencies deviating from the expected values are assigned ambiguous or homozygous calls by the software. These findings were validated by two independent experiments. A3) Methylation-sensitive PCR analysis of the X-chromosome locus was assessed on DNA isolated from manually dissected female thy-roid MPC (n = 1) or non-malignant adjacent thyroid tissue (n = 1) formalin fixed paraffin embedded specimen. Screenshot of electropherogram displaying discordant patterns of X-chromosome inactivation (arrowheads) in the thyroid MPC vs the benign thyroid tissue. For each sample, the corresponding plot is a quantitative representation of the size and amount of fluorescent PCR products amplified from undigested DNA or digested DNA with the methylation-sensitive enzyme HpaII when analyzed on an automated genotyping platform. The peak indices were estimated using (base pairs) that is the corresponding peak height (amount of PCR product), as quantified by DNA Genotyping software. These findings were validated by triplicate sample assays. B) Thyroid MPC cells with heterozygous BRAF\(^{WT/V600E}\) or normal pericytes with BRAF\(^{WT}\) were treated with the indicated concentrations of vemurafenib for 48 hours, and viability was determined using the Cell Titer-Glo ATP-based luminescence assay, with Dimethyl sulfoxide (control)–treated cells as the control. Cell growth curves were determined following two to three days of treatment with DMSO or vemurafenib. Arrows highlight change of cell shape in MPC cells vs control. All scale bars are =10\(\mu\) MPC cells images) and 50\(\mu\) (pericytes images). These data represent the average ± standard deviation (error bars) of eight independent replicate measurements (**P < 0.05, ***P < 0.01, ****P < 0.001, Mann-Whitney test, two-sided). C) A parallel plate similar to (B) was set up and corresponding phospho(p)ERK1/2 and pMEK1/2 protein levels measured from BRAF\(^{WT/V600E}\), MPC cells or BRAF\(^{WT}\) pericytes protein samples using western blotting assays. These data are representative of three independent experiments. D1-D2) BRAF\(^{WT/V600E}\)-thyroid MPC cells cocultured with human microvascular endothelial cells (HMVECs) induce tubule formation (asterisk, control) (magnifications in all images: 20×; scale bars = 200\(\mu\)) on growth factors reduced Matrigel in the presence of very low serum concentration (0.2%) within six hours compared to control. Please see the Supplementary Results (available online) for additional findings that may be of interest. D) Cells were treated with D1-D2) BRAF\(^{WT/V600E}\)-MPC cell viability and angiogenesis were classified as “low copy number” if it was between 1 mRNA copies/10\(^6\) copies, “moderate copy number” if it was between 1 and 15 mRNA copies/10\(^6\) copies, and “high copy number” if it was greater than 15 mRNA copies/10\(^6\) copies. These data represent the average ± standard deviation (error bars) of six independent replicate measurements (**P = 0.02, Mann-Whitney test, two-sided). E) Cell migration assays within five hours in human primary thyroid MPC cells with BRAF\(^{WT/V600E}\) or in pericytes with BRAF\(^{WT}\) in the presence of vehi-cl (control) or vemurafenib treatment. These data represent the average ± standard deviation (error bars) of six independent replicate measurements (**P = 0.02, Mann-Whitney test, two-sided). F) Quantitation of mRNA expression levels (copy number) by multi-gene transcriptional real-time reverse transcriptase PCR (RT-PCR) analysis of genes fundamental for extracellular matrix (ECM) remodeling and angiogenesis in human primary thyroid MPC cells with BRAF\(^{WT/V600E}\) or in pericytes with BRAF\(^{WT}\) in the presence of vehicle (control) or vemurafenib treatment for 24 hours (0.2% serum cell growth medium). Gene expression was classified as “low copy number” if it was below 1 mRNA copies/10\(^6\) copies, “moderate copy number” if it was between 1 and 15 mRNA copies/10\(^6\) copies, and “high copy number” if it was greater than 15 mRNA copies/10\(^6\) copies. These data represent the average ± standard deviation (error bars) of four to six independent replicate measurements (**P < 0.05, ***P < 0.01, ****P < 0.001, Mann-Whitney test, two-sided). G) A patient-derived xenograft (PDX) mouse model of BRAF\(^{WT/V600E}\)-human primary thyroid MPC cells engineered to express GFP (green fluorescent protein, scale bar=100\(\mu\)) injected in the subcutaneous compartment of the left ear of nude mice (n = 10) to visually follow perivascular growth of MPC cells over time in vivo. Mice were treated starting four days after BRAF\(^{WT/V600E}\)-MPC cells implantation (defined ‘baseline’) with vemurafenib (100 mg/kg twice daily by oral gavage) (n = 5) or vehicle (n = 5) for two consecutive weeks. MPC cell viability was determined by the CSS.1 histogram software calculating the absolute number of pixels in the GFP-positive areas (arrows) of the left ear divided by the total number of pixels (247,545) in the highlighted selec-tion area. These data represent the average ± standard deviation of two independent experiments (control-treated mice vs vemurafenib-treated mice, P = .007, Mann-Whitney test).
We propose that genetic testing for the BRAFV600E mutation is part of the pathological evaluation for multifocal and infiltrative MPMs, and patients with BRAFV600E-positive MPMs should be periodically re-evaluated for recurrence, especially with available targeted drug therapies.

References


Funding

Carmelo Nucera (Principal Investigator, Human Thyroid Cancers Preclinical and Translational Research at the Beth Israel Deaconess Medical Center/Harvard Medical School) was awarded grants by the National Cancer Institute/National Institutes of Health (IR21CA165039-01A1 and IR01CA181813-01A1) the American Thyroid Association (ATA), and ThyCa:Thyroid Cancer Survivors Association Inc. for Thyroid Cancer Research. Carmelo Nucera was also recipient of the Guido Berlucchi “Young Investigator” research award 2013 (Brescia, Italy). Beth Israel Deaconess Medical Center was recipient of a research grant by Roche (2011). CP was awarded a grant by the Tuberosclerosis Complex Research Program (TSCR) of the USA Department of Defense (DOD)/(W81XWH-13-1-0262).

Notes

The study funders had no role in the design of the study, the collection, analysis, or interpretation of the data, the writing of the manuscript, or the decision to submit the manuscript for publication.

We thank Drs. Yutaka Kawakami (Keio University, Tokyo, Japan) for providing HIV-U6 vectors. We also thank Drs. Shiva Gautam (statistical consultant), Alexander Gimelbrant, Janice Nagy, Gideon Bollag, Riccardo Taulli and Francesca Ianni, as well as Mrs. Sudeepa Syamala, Mrs. Mei Zheng, Miss Nina Hu, Mr. Bhavik Padmani and Neal Smith for technical assistance. We thank Professor William Aird for critical suggestions.

Authors’ contributions: Peter Sadow: performed pathologic diagnosis, reviewed immunohistochemical data, and edited manuscript. Carmen Priolo: edited manuscript, performed in vitro experiments, analyzed data, and designed experiments. Simona Nanni: performed in vitro experiments. Florian A. Karreth: edited manuscript, performed in vitro experiments, and analyzed data. Mark Duquette: performed in vitro experiments. Roberta Martinelli: performed in vivo experiments. Amjad Hussain: performed
in vivo experiments. **John Clohessy**: performed in vivo experiments. **Heinz Kutzner**: performed clinical-pathological diagnosis. **Thomas Mentzel**: performed clinical-pathological diagnosis. **Christopher Carman**: analyzed data. **Antonella Farsetti**: analyzed data. **Elizabeth Petri Henske**: edited manuscript and analyzed data. **Emanuele Palescandolo**: analyzed data. **Laura E. Macconail**: analyzed data. **Seum Chung**: performed surgical therapy. **Guido Fadda**: performed pathological analysis. **Celestino Pio Lombardi**: performed surgical therapy. **Antonella De Angelis**: analyzed data. **Oreste Durante**: analyzed data. **John A. Parker**: analyzed data, edited manuscript and reviewed radiologic data. **Alfredo Pontecorvi**: contributed with inpatient clinical exams. **Harold Dvorak**: edited manuscript and designed experiments. **Christopher Fletcher**: edited manuscript, performed pathologic diagnosis, and reviewed histo-pathological data. **Pier Paolo Pandolfi**: analyzed data and edited manuscript. **Jack Lawler**: analyzed data and edited manuscript. **Carmelo Nucera**: wrote and edited manuscript, reviewed clinical and experimental data, analyzed data, collected data, analyzed clinical exams, performed in vitro and in vivo experiments, and designed experiments. **Affiliations of authors**: Department of Pathology, Massachusetts General Hospital (PMS), Harvard Medical School, Boston, MA; Division of Pulmonary and Critical Care Medicine, Department of Medicine, Brigham and Women's Hospital (CP, EPH), Harvard Medical School, Boston, MA; Unit of Endocrinology, Department of Medicine, A. Gemelli, Catholic University, Roma, Italy (SN, AP); Division of Cancer Genetics, Department of Medicine (FAK, JC, PPP), Cancer Research Institute, Harvard Medical School, Boston, MA; Laboratory of Human Thyroid Cancers Preclinical and Translational Research, Division of Cancer Biology and Angiogenesis, Department of Pathology, Cancer Research Institute, Center for Vascular Biology Research (MD, CN), Beth Israel Deaconess Medical Center, Harvard Medical School, Boston, MA; Department of Medicine (RM, CVC) and Department of Pathology (AH, HD, JL), Center for Vascular Biology Research, Beth Israel Deaconess Medical Center, Boston, MA; Dermatopathologische Gemeinschaftspraxis, Siemensstrasse, Friedrichshafen, Germany (HK, TMI); National Research Council (CNR-IBCN) and Department of Experimental Oncology, Regina Elena Cancer Institute, Rome, Italy (AF); Department of Medical Oncology, Dana Farber Cancer Institute, Harvard Medical School, Boston, MA (EP, LEM); MIT/Broad Institute, Cambridge, MA (EP, LEM); Department of Plastic and Reconstructive Surgery, National Health Insurance Service Ilsan Hospital, Ilsan, Korea (ISI); Department of Pathology (GF) and Department of Surgery (CPL), A. Gemelli, Catholic University, Roma, Italy; Department of Experimental and Clinical Medicine (AMDA) and Unit of Radiotherapy (OD), University of Catanzaro, Italy; Department of Nuclear Medicine, Beth Israel Deaconess Medical Center (JAP); Department of Pathology, Brigham and Women's Hospital, Harvard Medical School, Boston, MA (CFI).