

## MYELOID NEOPLASIA

Recurrent *ETNK1* mutations in atypical chronic myeloid leukemia

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## Key Points

- Whole-exome sequencing reveals the presence of recurrent somatic mutations of *ETNK1* in patients with atypical chronic myeloid leukemia.
- *ETNK1* mutations impair the catalytic activity of the enzyme, causing a decrease in the intracellular levels of phosphoethanolamine.

Despite the recent identification of recurrent *SETBP1* mutations in atypical chronic myeloid leukemia (aCML), a complete description of the somatic lesions responsible for the onset of this disorder is still lacking. To find additional somatic abnormalities in aCML, we performed whole-exome sequencing on 15 aCML cases. In 2 cases (13.3%), we identified somatic missense mutations in the *ETNK1* gene. Targeted resequencing on 515 hematological clonal disorders revealed the presence of *ETNK1* variants in 6 (8.8%) of 68 aCML and 2 (2.6%) of 77 chronic myelomonocytic leukemia samples. These mutations clustered in a small region of the kinase domain, encoding for H243Y and N244S (1/8 H243Y; 7/8 N244S). They were all heterozygous and present in the dominant clone. The intracellular phosphoethanolamine/phosphocholine ratio was, on average, 5.2-fold lower in *ETNK1*-mutated samples ( $P < .05$ ). Similar results were obtained using myeloid TF1 cells transduced with *ETNK1* wild type, *ETNK1*-N244S, and *ETNK1*-H243Y, where the intracellular phosphoethanolamine/phosphocholine ratio was significantly lower in *ETNK1*-N244S ( $0.76 \pm 0.07$ ) and *ETNK1*-H243Y ( $0.37 \pm 0.02$ ) than in *ETNK1*-WT ( $1.37 \pm 0.32$ ;  $P = .01$  and  $P = .0008$ , respectively), suggesting that *ETNK1* mutations may inhibit the

catalytic activity of the enzyme. In summary, our study shows for the first time the evidence of recurrent somatic *ETNK1* mutations in the context of myeloproliferative/myelodysplastic disorders. (*Blood*. 2015;125(3):499-503)

## Introduction

Atypical chronic myeloid leukemia (aCML) is a clonal disease belonging to the myeloproliferative/myelodysplastic (MPN/MDS) group of hematological disorders according to the World Health Organization 2008 classification.<sup>1,2</sup> aCML closely resembles the classical chronic myeloid leukemia, with the main difference being that aCML is invariably negative for any *ABL1* rearrangement. The molecular lesions responsible for the onset of aCML remained

unknown since 2013, when, by applying whole-exome sequencing, our group demonstrated the presence of recurrent, somatic mutations of the *SETBP1* gene, showing that *SETBP1* mutations are associated with an increase in white blood cell count and with poor prognosis.<sup>3,4</sup> Despite this finding, a complete description of the somatic lesions responsible for the onset of this disorder is still lacking. To identify additional somatic abnormalities occurring in aCML, we extended

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our previous study to a larger aCML cohort comprising a total of 15 matched cases. The results are described here.

## Methods

### Patients

Diagnosis of aCML and related diseases was performed according to the World Health Organization 2008 classification.<sup>1,2</sup> All patients provided written informed consent, which was approved by the institutional ethics committee. This study was conducted in accordance with the Declaration of Helsinki. Patient samples were processed as previously described.<sup>3</sup>

### Exome sequencing

Genomic DNA (1 µg) was fragmented (500 bp), using a Diagenode Bioruptor Next Gen sonicator and processed according to the standard Illumina protocol. Genomic libraries were enriched for the exonic regions (Illumina TruSeq Exome Enrichment kit) and sequenced on an Illumina Genome Analyzer IIx with 76-bp paired-end reads using Illumina TruSeq chemistry.

### RNA sequencing

All RNA libraries were generated from 2 µg total RNA extracted with TRIzol (Life Technologies) following the standard Illumina protocol with minor modifications. Libraries were sequenced on an Illumina Genome Analyzer IIx with 76-bp paired-end reads, using Illumina TruSeq chemistry.

### Bioinformatics

Bioinformatic processing has been performed as previously described.<sup>3</sup> Sequences and alignments were quality-tested, using SAM-Profiler.<sup>5</sup> Copy number abnormalities were analyzed using CEQer.<sup>6</sup> The presence of fusions was assessed using FusionAnalyser,<sup>7</sup> and splicing sites mutations using SpliceFinder.<sup>8</sup>

### Validation of mutations and targeted resequencing

All variants were validated by Sanger sequencing. To confirm *ETNK1* variants, genomic DNA was polymerase chain reaction-amplified (100 ng per reaction) and sequenced using the following primers: ETNK1-FOR8 CATCAAAGTAAGGGGTTTCTGG and ETNK1-REV8 TCTGCCCTTGCTGTGTAG.

### Liquid chromatography–mass spectrometry

For liquid chromatography–mass spectrometry experiments, peripheral blood samples (buffy coat) were collected at diagnosis in individuals with aCML. Phenotype was evaluated by fluorescence-activated cell sorting analyses. Only samples with myeloid cells (positive for CD33, CD13, or CD117 staining) greater than 80% of the total were used. As a source of normal cells, we used lymphocytes obtained by culturing patient cells with 2.5 µg/mL phytohemagglutinin-M (PHA-M, Roche) and 200 UI/mL interleukin 2 (Aldesleukin, Novartis) for 3 to 4 days and then incubating cells for 2 to 3 weeks with interleukin 2 only, as previously described.<sup>3</sup> Phenotype was evaluated by fluorescence-activated cell sorter analysis. Only samples with lymphoid cells (positive for CD3, CD4, CD5, CD8, or CD19 staining) greater than 80% of the total were used.

Cell lysis and extraction were performed on a total of  $5 \times 10^6$  cells, as previously described,<sup>9</sup> with minor modifications. High-performance liquid chromatography (HPLC)-mass spectrometry quantitation was performed on a Shimadzu-Nexera HPLC chromatograph, interfaced with an AbSciex-3200 QTRAP mass spectrometer (AB SCIEX). HPLC analyses were performed on a Phenomenex Jupiter Proteo 1  $\times$  250 mm. MRM analyses were performed as follows: ion spray voltage, 5500 V; source temperature, 150°C; declustering potential, 35 V; collision energy, 20 eV; GAS1, 15 L/minute; curtain gas, 30 L/minute. Transitions 184.1/86 (phosphocholine) and 142.6/85 (phosphoethanolamine) were used for quantitation purposes. With this experimental setup, the response ratio of phosphocholine/phosphoethanolamine was 15.05.

**Table 1. Recurrent somatic lesions identified in 15 aCML samples by whole-exome sequencing**

Genes	Patients			
SETBP1	CMLPh-003	CMLPh-005	CMLPh-013	CMLPh-015
NRAS	CMLPh-003	CMLPh-004	CMLPh-013	CMLPh-014
EZH2	CMLPh-004	CMLPh-008	CMLPh-009	—
ASXL1	CMLPh-005	CMLPh-013	CMLPh-014	—
ETNK1	CMLPh-008	CMLPh-013	—	—
U2AF1	CMLPh-007	CMLPh-011	—	—

### Methylcellulose colony assay

From a peripheral blood sample (buffy coat; patient CMLPh-019), 150 000 cells were seeded in Methocult H4034 (StemCell Technologies), using the standard protocol. After 2 weeks of incubation at 37°C, 5% CO<sub>2</sub>, individual colonies were picked, washed with 200 µL phosphate-buffered saline, and lysed in 20 µL of the following buffer: 10 mM Tris-HCl, 50 mM NaCl, 6.25 mM MgCl<sub>2</sub>, 0.045% NP40, and 0.45% Tween-20 at pH 7.6. After adding 1 µL of 20 µg/mL proteinase K, the lysate was incubated at 56°C for 1 hour and at 95°C for 15 minutes. Subsequently, the sample was polymerase chain reaction-amplified by using the following primer pairs: ETNK1-p19-For: CATCAAGAATCTTTGTCAGGAG, ETNK1-p19-Rev: CAAATCCTGTGGGAATGAGAGAG, SETBP1-IntFor: AACTATGCCAAATCTCCAGC, and SETBP1-IntRev: GAAAGTTGTCCACAATGAGATG.

The amplicons were purified on agarose gel, using the QIAquick gel extraction kit (Qiagen), following the standard protocol and eluted in 40 µL H<sub>2</sub>O. Nine microliters of purified amplicon + 1 µL 25 µM forward primer were used for Sanger sequencing (GATC Biotech).

### Mutagenesis

pDONR233 plasmids encoding ETNK1 H243Y and N244S were generated using the following protocol: primers ETNK1\_H243Y\_For (CCATGCTATTCATGCATACAATGGCTGGATCCCC), ETNK1\_H243Y\_Rev (GGGGATCCAGCCATGTATGCATGAATAGCATGG), ETNK1\_N244S\_For (CTATTCATGCACACAGTGGCTGGATCCCCAAATC), and ETNK1\_N244S\_Rev (GATTTGGGGATCCAGCCACTGTGTGCATGAATAG) were used to mutagenize the pDONR223-ETNK1 entry vector (Addgene) with the Pfu Ultra High Fidelity enzyme (Agilent). The product was digested with DpnI (Roche), and 2 µL were used to transform the competent TOP10 bacterial strain (Life Technologies). The presence of the mutations was subsequently confirmed by Sanger sequencing. The coding sequence for *ETNK1* variants was then subcloned into the pLX304 vector, using Gateway technology (Life Technologies).

### Lentiviral infection

Two hundred ninety-three 293FT packaging cells (Life Technologies) were transfected with 7 µg psPAX2, 1.4 µg pCMV-VSV-G (Addgene), and 7 µg pLX304 (encoding V5-flagged wild type, H243Y, or N244S ETNK1), using jetPRIME (Polyplus-transfection SA). Lentivirus was collected 2 and 3 days after transfection. To generate TF1 cells stably infected with lentiviruses,  $5 \times 10^5$  cells were transduced by spin infection in lentiviral supernatants supplemented with 4 µg/mL polybrene (Sigma-Aldrich) and 20% RPMI 1640. After 48 hours, the cells expressing wild type and mutated ETNK1 were resuspended in complete medium and selected for 3 weeks in the presence of Blasticidin 10 µg/mL. Western blot was performed using anti-V5 (Abcam) 1:2000 in milk 5% for 2 hours at room temperature (RT) and anti-actin (Sigma) 1:2000 in milk 5% for 2 hours RT; secondary antibodies were anti-mouse (Bio-Rad) 1:2000 incubated for 1 hour RT and anti-rabbit (Bio-Rad) 1:2000 for 1 hour RT, respectively.

## Results

To gain insight into the somatic variants responsible for the onset of aCML, we generated whole-exome and transcriptome sequencing

**Table 2. Frequency of *ETNK1* mutations in 515 patient samples and 32 healthy donors**

Disease	Samples	Mutated samples	Mutated samples, %
AML	64	0	0.0
ALL	53	0	0.0
CLL	51	0	0.0
MDS	77	0	0.0
<b>MPN</b>			
CML	38	0	0.0
PV	18	0	0.0
ET	39	0	0.0
CNL	1	0	0.0
PMF	14	0	0.0
<b>MDS/MPN</b>			
aCML	68	6	8.8
CMML	77	2	2.6
JMML	5	0	0.0
MDS/MPN-u	10	0	0.0
Healthy donors	32	0	0.0

aCML, atypical chronic myeloid leukemia; ALL, acute lymphoblastic leukemia; AML, acute myeloid leukemia; CLL, chronic lymphocytic leukemia; CML, chronic myeloid leukemia; CMML, chronic myelomonocytic leukemia; CNL, chronic neutrophilic leukemia; ET, essential thrombocytosis; JMML, juvenile myelomonocytic leukemia; MDS, myelodysplastic syndromes; MDS/MPN-u=myelodysplastic/myeloproliferative neoplasm-unclassifiable; MPN, myeloproliferative neoplasms; PMF, primary myelofibrosis; PV, polycythemia vera.

data on a cohort of 15 matched case/control aCML samples. A total of 151 nonsynonymous and 42 synonymous single-nucleotide somatic variants were identified. Of these events, 140 were transitions and 53 were transversions. Of the 151 nonsynonymous variants, 141 were missense and 10 were nonsense mutations (supplemental Table 1, available on the *Blood* Web site). Globally, we identified a total of 6 recurrently mutated genes: *SETBP1*, *NRAS*, *EZH2*, *ASXL1*, *ETNK1*, and *U2AF1* (Table 1). Surprisingly, no evidence of somatic variants occurring on *CSF3R* could be found, despite the good sequencing coverage of the whole *CSF3R* coding region, which suggests *CSF3R* mutations are significantly more frequent in chronic neutrophilic leukemia than in aCML,<sup>10,11</sup> and therefore confirming the different biological basis of the 2 disorders.

In 2 (13.3%) of 15 samples, we identified the presence of missense, single-nucleotide variants occurring in the *ETNK1* gene (supplemental Figure 1A). Both *ETNK1* mutations appeared to be present as heterozygous variants in the dominant clone (mutant relative frequency >30%). Interestingly, the mutations affected 2 contiguous residues: H243Y and N244S. Sanger sequencing confirmed the presence and the somatic nature of the 2 variants (supplemental Figure 1B). No fusions or recurrent copy number abnormalities were detected.

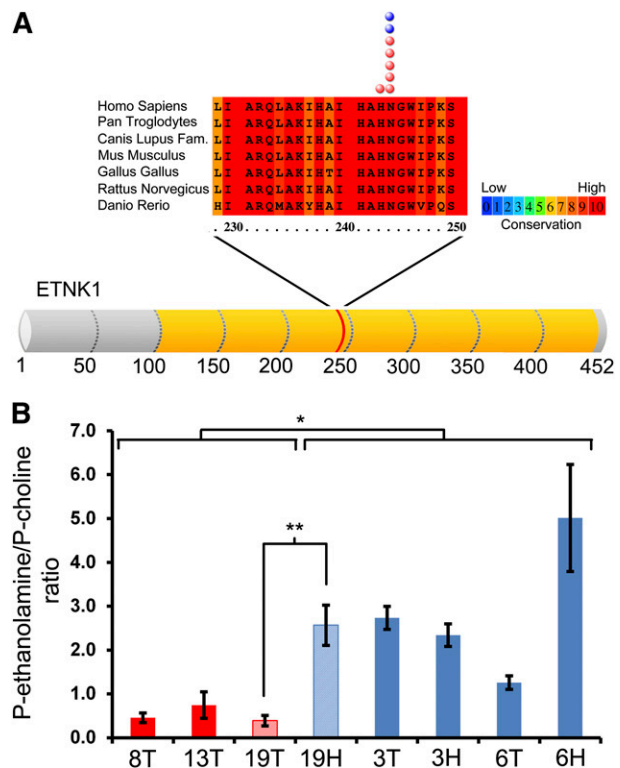
The evidence of recurrent, somatic *ETNK1* mutations has never been reported in cancer, suggesting these variants could be restricted to aCML and, possibly, to closely related clonal diseases. Targeted resequencing of 515 clonal hematological disorders (Table 2) showed evidence of mutated *ETNK1* in 6 of 68 aCML (8.8%; 95% confidence interval, 4%-18%) and in 2 of 77 chronic myelomonocytic leukemia (2.6%; 95% confidence interval, 0.2%-9.5%) samples. In all the cases, *ETNK1* variants appeared to be present as a heterozygous, dominant clone. No evidence of *ETNK1* mutations was found in the other clonal disorders examined, nor in healthy donors. All variants clustered in the same, highly conserved region<sup>12</sup> within the kinase domain (1/8 H243Y and 7/8 N244S) and were heterozygous.

In 2 of 6 (33%; 95% confidence interval, 9%-70%) *ETNK1*-mutated aCML cases, we detected the presence of a coexisting

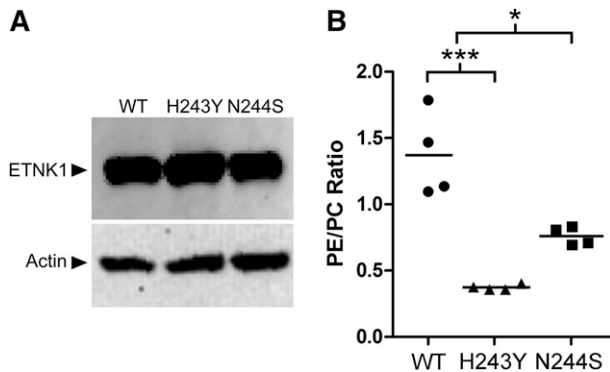
*SETBP1* variant. The fraction of *SETBP1* mutations within this group was perfectly in line with the overall frequency of *SETBP1* mutations in aCML, suggesting that mutations occurring in *ETNK1* and *SETBP1* are not mutually exclusive.

To assess whether *SETBP1* and *ETNK1* mutations coexist in the same clone or are present in separate cells, we performed a colony assay in a patient in whom both mutations had been detected (CMLPh-019). Targeted resequencing of both genes on gDNA from 40 individual colonies demonstrated the coexistence of the 2 variants as heterozygous mutations in all the clones tested (supplemental Figure 2).

To further characterize the pattern of *ETNK1* mutations in cancer, we tested the presence of somatic *ETNK1* variants in leukemias, as well as solid tumors, in a total of 60 paired whole-genome and more than 600 exomes making up 276 paired tumor/germline primary samples and 344 cancer cell lines ([http://cancer.sanger.ac.uk/cancergenome/projects/cell\\_lines/](http://cancer.sanger.ac.uk/cancergenome/projects/cell_lines/))<sup>13-19</sup> by reevaluation of the raw sequencing data. No evidence of recurrent *ETNK1* mutations could be found, which suggests that *ETNK1* mutations are more frequent in aCML and closely related disorders. Further studies with larger



**Figure 1. *ETNK1* mutations in patients with aCML/chronic myelomonocytic leukemia.** (A) Outline of the *ETNK1* protein. The yellow region highlights the catalytic domain of the protein. The red band corresponds to the mutational hotspot. In the upper part, a conservation analysis of the mutated locus is shown. The red and blue circles highlight the position of individual mutations identified in aCML and chronic myelomonocytic leukemia, respectively. (B) Intracellular phosphoethanolamine/phosphocholine ratio in patients with aCML. The red bars show the intracellular phosphoethanolamine/phosphocholine ratio of leukemic (myeloid) cells in 3 *ETNK1* mutated patients (8T, 13T, and 19T, corresponding to patients CMLPh-008, CMLPh-013, and CMLPh-019). The blue bars show the intracellular phosphoethanolamine/phosphocholine ratio of myeloid (3T, 6T) or lymphoid (3H, 6H) cells negative for *ETNK1* mutations (patients CMLPh-003 and CMLPh-006). The light red and light blue bars indicate the intracellular phosphoethanolamine/phosphocholine ratio of a matched myeloid/lymphoid sample whose leukemic (myeloid) cells are positive for *ETNK1* mutation. The error bars represent the standard deviation of 3 experiments. The asterisks indicate the significance level: \**P* < .05; \*\**P* < .005.



**Figure 2. Phosphoethanolamine/phosphocholine ratio in a TF1-ETNK1 cellular model.** (A) Western blot anti-ETNK1 showing the intracellular level of ETNK1 protein in TF1-WT, N244S, and H243Y. (B) Intracellular phosphoethanolamine/phosphocholine ratio as assessed by HPLC–mass Spectrometry in TF1 cells transduced with ETNK1-WT, N244S, and H243Y.

cohorts will be required to thoroughly assess the presence and the frequency of *ETNK1* mutations in other clonal disorders. Conservation analysis revealed that the mutated locus is highly conserved across many species (Figure 1A), which indicates it may play a critical, yet unknown, functional role.

To assess the functional effect of *ETNK1* mutations, the intracellular level of phosphoethanolamine, the product of the *ETNK1* kinase, was assessed on *ETNK1*-positive and *ETNK1*-negative aCML primary samples by using liquid chromatography–mass spectrometry techniques. The analyses were performed on a total of 3 primary peripheral blood myeloid samples carrying *ETNK1* mutations (CMLPh-008T, CMLPh-013T, CMLPh-019T), on 4 *ETNK1*-WT aCML samples, and on *ETNK1*-WT autologous lymphoid cells from *ETNK1*-mutated patient 19 (CMLPh-019H). Of the mutated samples, one carried the H243Y and 2 the N244S variant. All the quantifications were performed at least in triplicate. The results showed that the intracellular phosphoethanolamine/phosphocholine ratio was, on average, 5.2-fold lower in *ETNK1*-mutated samples than in the nonmutated ones ( $P < .05$ ) and suggested that *ETNK1* mutations may damage the catalytic activity of the enzyme (Figure 1B). A similar analysis performed using myeloid TF1 cells transduced with *ETNK1*-WT, *ETNK1*-N244S, or *ETNK1*-H243Y revealed a similar pattern (Figure 2), with an intracellular phosphoethanolamine/phosphocholine ratio significantly lower in *ETNK1*-N244S ( $0.76 \pm 0.07$ ) and *ETNK1*-H243Y ( $0.37 \pm 0.02$ ) than in *ETNK1*-WT ( $1.37 \pm 0.32$ ;  $P = .01$  and  $P = .0008$ , respectively).

## Discussion

*ETNK1* encodes an ethanolamine kinase, which catalyzes the first step of the de novo phosphatidylethanolamine (PE) biosynthesis pathway (supplemental Figure 3).<sup>20</sup> It is responsible for the phosphorylation of ethanolamine to phosphoethanolamine, which is then further processed to PE by a cytidylyltransferase (ECT1) and by the amino alcohol phosphotransferase EPT1.<sup>20</sup> PE is involved in many biochemical processes, among them defining membrane architecture,<sup>21</sup> regulation of the topology of transmembrane domains of membrane proteins,<sup>22</sup> anchoring specific PE-binding proteins to the cell membrane, ensuring the proper progression of cytokinesis during cell division,<sup>23</sup> and optimal activity of the respiratory complexes in the inner membrane of mitochondria.<sup>23</sup> Given the pleiotropic role

of PE, dissecting the biological effects of the *ETNK1* mutations will be a complex task. However, the occurrence of all the variants in a highly conserved region and the demonstration that they impair the catalytic activity of *ETNK1* suggest these mutations may play a significant functional role.

Within myeloid neoplasms, major differences in disease phenotype can be driven by different combinations of commutated genes.<sup>24</sup> Interestingly, aCML and chronic myelomonocytic leukemia share not only clinical and morphological features but also mutant genes.<sup>3,25</sup> In this study, targeted resequencing of patients with various myeloid neoplasms revealed somatic *ETNK1* mutations only in subjects with aCML or chronic myelomonocytic leukemia, confirming that these disorders have a common genetic basis.

In summary, our study shows for the first time the evidence of recurrent somatic mutations of the *ETNK1* gene in the context of MDS/MPN clonal disorders. Further studies will be required to thoroughly dissect the functional effects of these mutations.

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## Authorship

Contribution: C.B.G.-P. conceived of research, critically revised data, and wrote the manuscript; C.D., A. Parmianai, and D.F. generated Sanger sequencing data and validated somatic variants; A. Pirola generated next-generation sequencing (NGS) data; S.R. generated Sanger sequencing data and critically revised the manuscript; G. Signore generated and analyzed liquid chromatography–mass spectrometry data; V.P. analyzed liquid chromatography–mass spectrometry data and critically revised the manuscript; L.M. provided vital reagents and critically revised the manuscript; R.S. analyzed NGS data; V.M. and M.P. generated Sanger sequencing data; G.G., A.M., C.P., G. Saglio, E.U., D.-W.K., D.R., K.Z., N.V., A.S., H.B., J.B., L.C., M. Carrabba, E.E., P.J.C., and M. Cazzola provided vita reagents and critically revised the manuscript; M.P. generated Sanger sequencing data; G.R.B. analyzed cell lines and NGS data and revised the manuscript; E.P. provided vital reagents, analyzed NGS data, and critically revised the manuscript; and R.P. conceived research, generated data, and wrote the manuscript.

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