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ORIGINAL ARTICLE

Novel recurrent mutations in ethanolamine kinase 1 (*ETNK1*) gene in systemic mastocytosis with eosinophilia and chronic myelomonocytic leukemiaTL Lasho¹, CM Finke¹, D Zblewski¹, M Patnaik¹, RP Ketterling², D Chen³, CA Hanson³, A Tefferi¹ and A Pardanani¹

Although *KITD816V* occurs universally in adult systemic mastocytosis (SM), the clinical heterogeneity of SM suggests presence of additional phenotype-patterning mutations. Because up to 25% of SM patients have *KITD816V*-positive eosinophilia, we undertook whole-exome sequencing in a patient with aggressive SM with eosinophilia to identify novel genetic alterations. We conducted sequencing of purified eosinophils (clone/tumor sample), with T-lymphocytes as the matched control/non-tumor sample. In addition to *KITD816V*, we identified a somatic missense mutation in ethanolamine kinase 1 (*ETNK1*N244S) that was not present in 50 healthy controls. Targeted resequencing of 290 patients showed *ETNK1* mutations to be distributed as follows: (i) SM ($n = 82$; 6% mutated); (ii) chronic myelomonocytic leukemia (CMML; $n = 29$; 14% mutated); (iii) idiopathic hypereosinophilia ($n = 137$; < 1% mutated); (iv) primary myelofibrosis ($n = 32$; 0% mutated); and (v) others ($n = 10$; 0% mutated). Of the 82 SM cases, 25 had significant eosinophilia; of these 20% carried *ETNK1* mutations. The ten mutations (N244S = 6, N244T = 1, N244K = 1, G245A = 2) targeted two contiguous amino acids in the *ETNK1* kinase domain, and are predicted to be functionally disruptive. In summary, we identified novel somatic missense *ETNK1* mutations that were most frequent in SM with eosinophilia and CMML; this suggests a potential pathogenetic role for dysregulated cytidine diphosphate-ethanolamine pathway metabolites in these diseases.

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INTRODUCTION

Systemic mastocytosis (SM) is a chronic myeloproliferative neoplasm (MPN) characterized by an accumulation of morphologically and immunophenotypically abnormal mast cells (MC) in extra-cutaneous organs.¹ Although >90% of adult SM patients can be demonstrated to harbor the *KITD816V* mutation,² which constitutively activates KIT kinase activity and is thought to represent the driver mutation for this malignancy, the clinical presentation of SM is extremely heterogeneous. Patients with indolent SM have symptoms primarily related to MC degranulation with a normal life expectancy; in contrast, patients with other variants (aggressive SM (ASM), SM-AHD (SM associated with an associated hematological neoplasm) and MC leukemia) have an aggressive clinical course with features of organ dysfunction related to MC infiltration and shortened life expectancy.³ Possible explanations for the diverse clinical presentation include the lineage differential potential of the stem/progenitor cell targeted by the *KITD816V* mutation (reviewed by Pardanani⁴), and/or the co-occurrence (with *KITD816V*) of other pathogenetically relevant mutations that modify the disease phenotype in a given SM patient.^{5,6}

Approximately 15–20% of SM patients present with significant peripheral blood (PB) and bone marrow eosinophilia;³ we have previously demonstrated that the eosinophils in such cases are derived from the *KITD816V*-mutated neoplastic clone.⁷ In the current study, we pursued an exome-sequencing strategy using DNA isolated from (clonal/tumor) eosinophils that were purified to near-homogeneity from a patient with *KITD816V*-mutated ASM

with associated eosinophilia, in order to identify genetic alterations of potential pathogenetic significance. T-lymphocytes enriched from PB served as the matched control/non-tumor sample in our experimental approach. On the basis of the initial findings from this index case, targeted resequencing of additional samples from patients with SM and other MPNs were undertaken to analyze the prevalence of disease-relevant mutations across a broader cross-section of myeloid malignancies.

METHODS

The current study was approved by the Mayo Clinic institutional review board. All patients provided written informed consent for research sample collection. Diagnosis of SM and other MPNs was as per the 2008 World Health Organization (WHO) criteria.⁸ Granulocytes and PB mononuclear cells were enriched from whole blood using standard double-density Ficoll-gradient centrifugation. T-lymphocytes were enriched from the PB mononuclear cells fraction by positive selection of CD3 antibody-labeled cells by magnetic bead separation using standard protocols (StemCell Technologies, Vancouver, CA, USA). Similarly, a relatively pure population of eosinophils was isolated by negative selection of the CD16-labeled granulocyte fraction using the same methodology. We have previously validated the latter method as yielding eosinophils of $\geq 97\%$ purity.⁷

DNA was isolated from eosinophil and T-lymphocyte fractions using standard methods. Matched genomic libraries enriched for exonic regions (Agilent SureSelect XT kit; Agilent Technologies, Santa Clara, CA, USA) were prepared and sequenced on an Illumina HiSeq 2000 Analyzer using Illumina TruSeq chemistry (Illumina, San Diego, CA, USA). Data were analyzed using the GeneSifter Software (Perkin Elmer, Seattle, WA, USA). Post-processed alignments from the clonal fraction were compared with

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alignments from the control to make calls for somatic variants. Candidate mutations were validated by Sanger sequencing (see Supplementary Data for detailed experimental methods regarding library preparation, target capture, paired-end sequencing and data analysis).

RESULTS

Our experimental strategy was validated by identification of heterozygous *KITD816V* in eosinophils (70 reads, 47.1% mutant), but not in T-lymphocytes (66 reads, 0.0% mutant) in the index patient. A detailed list of somatic coding region single-nucleotide variant calls is presented in Supplementary Table 1. Among the non-synonymous single-nucleotide variants of interest (novel variants, as well as known oncogenic mutations) ranked by percentage mutant burden in clonal eosinophils were: enhancer of zeste homolog 2 (*EZH2*), *EZH2*H694R (tumor: 38 reads, 97.4% mutant; control: 32 reads, 3.1% mutant), signal transducer and activator of transcription 5 α (*STAT5a*), *STAT5a*V707E (tumor: 30 reads, 67.0% mutant; control: 33 reads, 0.0% mutant) and ethanolamine kinase 1 (*ETNK1*), *ETNK1*N244S, (tumor: 38 reads, 50.0%; control: 72 reads, 1.4%; Figure 1).

We conducted targeted re-sequencing of *STAT5a* and *ETNK1* in healthy controls and additional patient samples to assess whether the aforementioned novel genomic variants in these genes represented recurrent molecular abnormalities and whether they were associated with a specific MPN. Neither of the aforementioned genetic variants was identified in granulocyte-derived DNA from 50 healthy controls. In the first stage, we re-sequenced 125 patients, including 50 patients with SM, and 25 each with primary myelofibrosis, chronic myelomonocytic leukemia (CMML) and primary/idiopathic hypereosinophilia. No additional patients harboring the aforementioned *STAT5a* variant were identified. We did however identify *ETNK1* mutations in exon 3 in nine additional patients (i.e., ten patients in total); of these, six patients (including the index case) exhibited significant eosinophilia (four with SM-AHD (the latter was 'MPN-NOS' (MPN-not otherwise specified) and 'MDS/MPN-NOS' (myelodysplastic syndrome/MPN-NOS) in two cases each) and one each with ASM and idiopathic

hypereosinophilia; Table 1; Figure 1). The remaining four patients with *ETNK1* mutations were diagnosed with CMML and notably, none of these cases had associated eosinophilia. Consequently, *ETNK1* mutation prevalence in the initial screen was 10/125 (8%). We also screened the subgroup with wild-type *ETNK1* ($n = 115$), for mutations in exon 3 of its paralogous gene, *ETNK2*, however no additional mutations were identified in the latter gene. In the second stage, we conducted targeted re-sequencing of 165 additional MPN patients for *ETNK1* mutations. Of the total 290 patients screened, the disease distribution (percentage *ETNK1* mutated) was as follows: (i) SM ($n = 82$; 6% mutated), (ii) CMML ($n = 29$; 14% mutated), (iii) idiopathic hypereosinophilia ($n = 137$; < 1% mutated), (iv) primary myelofibrosis ($n = 32$; 0% mutated) and (v) Others ($n = 10$; 0% mutated). Of the 82 total SM patients studied, 36 had indolent SM, 26 had SM-AHD and 20 had ASM. Of the 82 SM cases, 25 had significant associated eosinophilia; of these 5/25 patients (20%) carried *ETNK1* mutations.

We identified 4 novel *ETNK1* mutations involving a total 10 patients (N244S=6, N244T=1, N244K=1 and G245A=2; Figure 1). In every case, the mutation was heterozygous and targeted one of two contiguous amino acid residues (N244 or G245) within the predicted kinase domain. *ETNK1* is 452 amino acids long, of which residues 100–444 comprise the protein kinase-like domain containing the putative nucleotide-binding site and the catalytic-site in an inter-lobe cleft. There is a high degree of sequence conservation between species in this region (Supplementary Figure 1), indicating a high degree of functional conservation. Computational tools for predicting the functional consequence of sequence changes targeting N244 and G245 suggested that the aforementioned *ETNK1* mutations were functionally deleterious (<http://snps.biofold.org/phd-snp/phd-snp.html> and <http://genetics.bwh.harvard.edu/pph2/index.shtml>). Four of five patients with SM harbored a concurrent *KITD816V* mutation; interestingly, one patient was found to have wild-type *KIT* sequence at this locus despite use of an allele-specific PCR for mutation detection (sensitivity $\leq 0.01\%$; Table 1).³ Although not formally analyzed, aside from presence of significant concurrent eosinophilia in non-CMML patients in our study, there were no

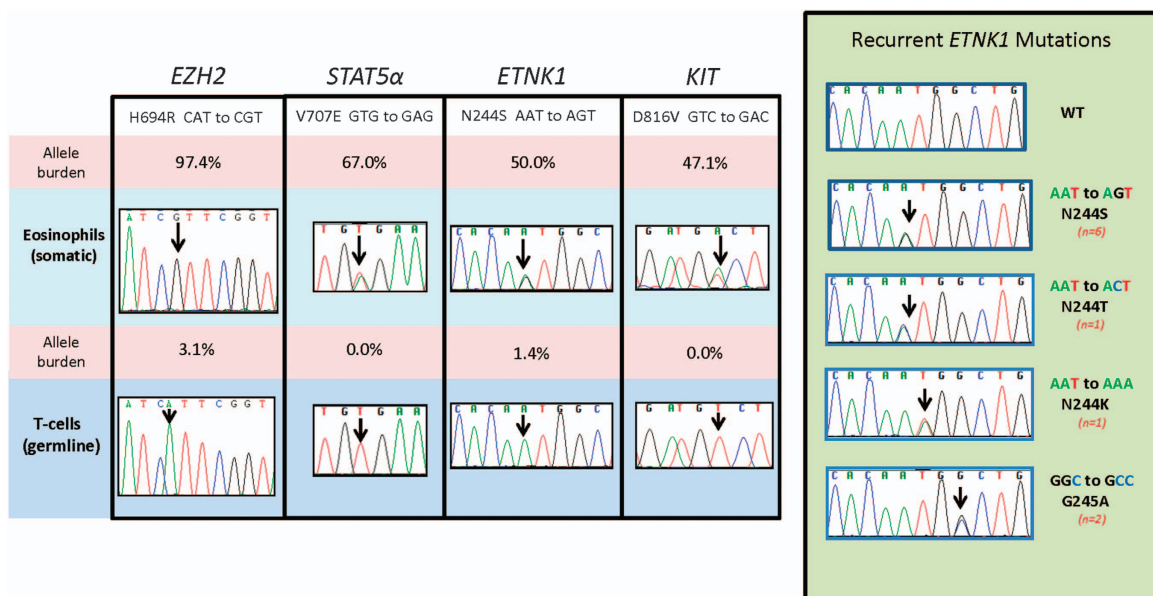


Figure 1. Left panel: targeted resequencing confirming exome-sequencing results in index case with aggressive systemic mastocytosis (ASM) with associated eosinophilia. Sanger sequencing confirmed the presence of coding region non-synonymous sequence variants in *KIT*, *EZH2*, *STAT5a* and *ETNK1* genes in the eosinophil fraction but not in the T-lymphocyte fraction. The percentage mutated alleles of total reads for each fraction and nucleotide/amino acid nomenclature for the observed genetic variation are shown. Right panel: Sanger sequencing traces of various somatic *ETNK1* mutations (four mutation types, total ten mutations) identified in our cohort, showing nucleotide and amino acid change for every variant.

Table 1. Clinical characteristics of *ETNK1*-mutated patients with significant eosinophilia

Patient characteristics (at the time of <i>ETNK1</i> testing)						
Diagnosis	ASM+ eosinophilia	SM-AHD MDS/MPN-NOS+ eosinophilia	SM-AHD MPN-NOS+ eosinophilia	SM-AHD MDS/MPN-NOS+ eosinophilia	SM-AHD MPN-NOS+ eosinophilia	Idiopathic hypereosinophilia
Age (years)	80	84	69	61	53	44
Gender	Female	Female	Male	Male	Male	Male
Bone marrow karyotype	46,XX	46,XX	46,XY	Unbalanced t(1;7) all metaphases	46,XY	46,XY
<i>BCR-ABL1</i> FISH	Negative	Negative	Not done	Negative	Negative	Not done
<i>FIP1L1-PDGFR</i> A	Negative	Negative	Negative	Not done ^a	Not done ^a	Negative
<i>KITD816V</i>	Positive	Positive	Positive	Negative ^b	Positive	Negative
<i>JAK2V617F</i>	Negative	Not done	Negative	Negative	Negative	Not done
<i>ETNK1</i> mutation	N244S	G245A	N244S	N244S	G245A	N244S
Hemoglobin (g/dl)	10.7	10.7	9.6	9.6	8.8	15.9
Leukocyte count (x10 ⁹ /l)	40.1	13.5	40.8	23.6	73.4	12.0
Platelet count (x10 ⁹ /l)	388	143	740	90	54	261
Absolute eosinophil count (x10 ⁹ /l)	24.5	6.66	0.02 (6.0 at diagnosis)	20.83	2.94	5.76
Circulating blasts (%)	0	0	0	0	0	0
Serum tryptase level (ng/ml)	1050	439	212	Not done	674	7.5
Red cell transfusion required	No	No	Yes	No	No	No
Prior splenectomy	No	No	Yes	No	No	No
Prior treatment	None	None	HU, ANA, Imatinib	HU	None	None
Subsequent treatment	Imatinib, IFN+ prednisone	IFN+prednisone + bisphosphonate+ XRT spinal lesion	Cladribine, IFN+ prednisone	IFN	Imatinib	NA

Abbreviations: ANA, anagrelide; ASM, aggressive systemic mastocytosis; *ETNK1*, ethanolamine kinase 1; FISH, fluorescence *in situ* hybridization; HU, hydroxyurea; IFN, interferon α ; MDS, myelodysplastic syndrome; MPN, myeloproliferative neoplasm; NA, not available; NOS, not otherwise specified; SM, systemic mastocytosis; SM-AHD, SM with associated hematological disorder; XRT, external beam radiotherapy. ^aPre-dated discovery of *FIP1L1-PDGFR*A mutation. ^bEvaluated by allele-specific polymerase chain reaction for *KITD816V* on archived bone marrow aspirate sample.

other clinical, molecular/cytogenetic, or bone marrow histologic characteristics that were unique to *ETNK1*-mutated patients as compared with disease-specific non-mutated patients (Table 1). Limited functional studies were conducted wherein human *ETNK1* protein (wild-type and N244S mutant) was overexpressed in murine Ba/F3 cells by means of a murine stem cell virus retroviral expression vector. In these studies, no discernable effect on cell proliferation, survival or apoptosis, regardless of ambient interleukin-3 concentrations was noted, as compared with empty vector control (data not shown).

DISCUSSION

We originally reported our discovery of recurrent *ETNK1* mutations in MPN patients in June 2014.⁹ In the current study, we found *ETNK1* mutations to be largely restricted to patients with SM with associated eosinophilia (20%) and CMML (14%). Although the association of *ETNK1* mutations with eosinophilia was intriguing, it appeared to be restricted to those patients with clonal myeloproliferative eosinophilia. In contrast, in our screening of a large cohort of patients with idiopathic eosinophilia as per WHO criteria (i.e., those lacking histologic, cytogenetic or molecular characteristics of clonal hematopoiesis), only one of 137 patients was found to harbor an *ETNK1* mutation. The association of *ETNK1* mutations with CMML was somewhat less intriguing given our current knowledge which suggests a high degree of genetic instability inherent to CMML, underscored by the high frequency of somatic mutations in diverse genes and oncogenic pathways identified in genetic screening studies.¹⁰ Nevertheless, as with most other recently identified pathogenetically relevant mutations in *BCR-ABL1*-negative MPNs, *ETNK1* mutations were not uniformly associated with a specific MPN in the current study.

ETNK1 is an ethanolamine-specific kinase that catalyzes the phosphorylation of ethanolamine by ATP to yield phosphoethanolamine, which is the first and possibly rate-limiting step committing ethanolamine to the CDP-ethanolamine pathway for biosynthesis of phosphatidylethanolamine (PE) in mammalian tissues.¹¹ PE and phosphatidylcholine (PC) are zwitterionic glycerophospholipids and represent the most abundant phospholipids in eukaryotic cells that are synthesized by two related branches of the Kennedy pathway referred to as the CDP-ethanolamine and CDP-choline pathways, respectively.¹² PE has diverse roles including a major role in maintaining cell-membrane architecture and the topology of transmembrane proteins (summarized by Gibellini and Smith¹³). Further, PE has a major role during cell division in ensuring the proper progression of cytokinesis, and is the precursor for important biologically active molecules such as diacylglycerols, fatty acids and phosphatidic acid. PE also provides for phosphoethanolamine capping of glycoposphatidylinositol that anchors many proteins to the cell surface. Finally, PE phospholipids can exist as plasmalogens that have a key role in membrane dynamics, scavenging of membrane localized free radical species and as a precursor for prostanoid synthesis and intracellular signaling. Disruption of the *Drosophila* ethanolamine kinase gene, *eas*, revealed a phenotype of seizure, neuronal failure and paralysis consistent with altered membrane phospholipid composition.¹⁴ In murine models, disruption of the ethanolamine kinase-2 and phosphoethanolamine cytidyltransferase genes were associated with neonatal lethality (summarized by Vance and Vance).¹⁵ Abnormalities in choline metabolism have emerged as a metabolic hallmark of cancer; overexpression of several enzymes in the PC biosynthetic pathway have been identified and may have a role as prognostic markers in various cancers (summarized by Glunde *et al.*).¹⁶ Further, recent studies

have indicated a reciprocal interaction between hyperactive choline metabolism, leading to increased levels of phosphocholine/total choline, and enhanced oncogenic signaling via RAS, PI3K-AKT, and HIF-1 α pathways. There is an ongoing phase 1 study of a choline kinase- α inhibitor for the treatment of patients with advanced solid tumors (Clinicaltrials.gov, study# NCT01215864). The aforementioned data make it highly plausible that abnormalities in PE metabolism induced by mutations in relevant enzymes in its biosynthetic pathway such as ETNK1 have a similar role in oncogenesis as seen with altered PC metabolism.

Although the current study provides novel insights into the potential role of altered glycerophospholipid metabolism in MPN pathogenesis, significant additional work remains to be done. First, the mechanism by which altered levels of CDP-ethanolamine pathway intermediates lead to a growth or survival advantage for the affected clone remains to be determined. Further, it is unclear based upon current data as to whether the aforementioned ETNK1 mutations constitute the driver mutation for the underlying MPN or MDS/MPN. On the basis of the known co-occurrence of other pathogenic mutations in SM (i.e., KITD816V) and CMML (e.g., ASXL1, DNMT3A, CBL), a reasonable assumption is that ETNK1 mutations provide a phenotype-patterning effect (e.g., eosinophilia) and/or incremental growth/survival advantage such as the co-operation seen between TET2 and KITD816V or JAK2V617F mutations.^{17–19}

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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AUTHOR CONTRIBUTIONS

AP, TLL and AT designed the study and analyzed the data. TLL conducted primary analysis of the exome-sequencing data, cloning of ETNK1 and functional experiments. TLL and CMF performed Sanger sequencing for mutation validation. DZ and MP provided patients for the study. RPK analyzed molecular/cytogenetic data. DC and CAH reviewed bone marrow histology. TLL and AP wrote the first draft of the manuscript. All authors reviewed the draft, provided input for revisions and approved the final version of the manuscript.

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Supplementary Information accompanies this paper on Blood Cancer Journal website (<http://www.nature.com/bcj>)