

Next-generation sequencing of the *TET2* gene in 355 MDS and CMML patients reveals low-abundance mutant clones with early origins, but indicates no definite prognostic value

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Mutations in the *TET2* gene are frequent in myeloid disease, although their biologic and prognostic significance remains unclear. We analyzed 355 patients with myelodysplastic syndromes using "next-generation" sequencing for *TET2* aberrations, 91 of whom were also subjected to single-nucleotide polymorphism 6.0 array karyotyping. Seventy-one *TET2* mutations, with a relative mutation abundance (RMA) $\geq 10\%$, were identified in 39 of 320 (12%) myelodysplastic syndrome and 16 of 35 (46%) chronic myelomonocytic leukemia patients ($P < .001$).

Interestingly, 4 patients had multiple mutations likely to exist as independent clones or on alternate alleles, suggestive of clonal evolution. "Deeper" sequencing of 96 patient samples identified 4 additional mutations (RMA, 3%-6.3%). Importantly, *TET2* mutant clones were also found in T cells, in addition to CD34⁺ and total bone marrow cells (23.5%, 38.5%, and 43% RMA, respectively). Only 20% of the *TET2*-mutated patients showed loss of heterozygosity at the *TET2* locus. There was no difference in the frequency of genome-wide aberrations, *TET2* ex-

pression, or the JAK2V617F 46/1 haplotype between *TET2*-mutated and nonmutated patients. There was no significant prognostic association between *TET2* mutations and World Health Organization subtypes, International Prognostic Scoring System score, cytogenetic status, or transformation to acute myeloid leukemia. On multivariate analysis, age (> 50 years) was associated with a higher incidence of *TET2* mutation ($P = .02$). (*Blood*. 2010;116(19):3923-3932)

Introduction

The myelodysplastic syndromes (MDSs) are clonal disorders of hematopoiesis characterized by dysplasia, ineffective hematopoiesis, and peripheral cytopenias. Evolution to acute myeloid leukemia (AML) occurs in approximately 25% of cases.¹ Based on the International Prognostic Scoring System (IPSS), patients in low-risk, intermediate-1, intermediate-2 and high-risk subgroups have a median survival of 5.7, 3.5, 1.2, and 0.4 years, respectively.² The early genetic lesions that lead to clonal hematopoiesis in MDS remain poorly understood,³ with a low frequency of point mutations in genes, such as *TP53* (5%-10%), *RUNX1* (2-10%), *NRAS* (10%-15%), and *FLT3* (2%-5%).⁴

Several groups have used single-nucleotide polymorphism (SNP) genotyping to identify genomic aberrations, including uniparental disomy (UPD), in MDS patients, which were undetectable by conventional metaphase cytogenetic analysis.⁵ We identified a common region of UPD spanning chromosome 4q24 in 8% of MDS patients that has been confirmed by other groups and led to the discovery of mutations in the *TET2* gene.⁶⁻¹³ The TET family consists of 3 proteins (TET1, TET2, and TET3) sharing 2 highly conserved domains. TET1 protein has been implicated in leukemia as a fusion partner of the *MLL* gene product in the t(10;11)(p12;q23) chromosomal translocation and has been shown

to be capable of hydroxymethylation and thereby possibly contributing to cytosine methylation patterns and epigenetic gene regulation.¹⁴⁻¹⁸

Mutations in the *TET2* gene have been identified in 19%-26% of MDS and 50% of chronic myelomonocytic leukemia (CMML) patients, making it one of the most frequently mutated genes in these myeloid neoplasms.^{9-11,13} Although, *TET2* mutations have been shown to be an early-disease event and pre-*JAK2* lesion¹¹ in myeloproliferative neoplasms (MPNs), recent data from colony assays, in some MPN patients, show that *TET2* and *JAK2* mutations can occur independently of each other.¹⁹ This is highlighted in MPN cases in whom *TET2* mutations were only detected at the time of myeloid leukemic transformation.²⁰⁻²² The prognostic significance of *TET2* aberrations in MDS remains unclear, with conflicting reports on the effect of mutant *TET2* on overall survival.^{10,23} The majority of reported *TET2* mutations are likely to cause inactivation of the TET2 protein, suggesting a tumor-suppressor role for this protein. Furthermore, *TET2* mRNA is highly expressed in normal myeloid progenitor cells, granulocytes, and erythroid cells, whereas granulocytes isolated from MDS cases have reduced *TET2* expression, highlighting the importance of wild-type levels of TET2 protein in myelopoiesis. Decreased *TET2*

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expression is also noted in CD34⁺ cells, CD13⁺/33⁺ myelomonocytic cells, and CD71⁺ erythroid cells isolated from bone marrow, and mature granulocytes obtained from patients with no detectable mutations, suggesting alternative mechanisms of *TET2* dysregulation.¹¹

The advent of next-generation “deep” sequencing technology has enabled cost-effective, high-throughput sequencing, with the added advantage of increased sensitivity more than the conventional Sanger methodology. To examine the prevalence of *TET2* mutations and, especially, lower abundance *TET2* mutant clones, we used ultrasensitive “454-deep sequencing” to investigate the integrity of the *TET2* gene in a well-characterized cohort of 355 de novo MDS and CMML patients. The presence of *TET2* mutations existing as major/minor clones or single/multiple mutants was analyzed for association with conventional clinical and prognostic parameters as well as *TET2* gene expression, JAK2V617F mutation, and 46/1 haplotype status. We also analyzed DNA from CD34⁺, CD3⁺ cells and constitutional DNA to determine mutant clone size and to ascertain the stage of acquisition of the *TET2* aberration. In 91 patients, high-resolution Affymetrix SNP6 microarrays were used to detect cryptic genomic aberrations and their relationship to *TET2* mutations.

Methods

Patients and sample preparation

A total of 355 patients with a median age of 64 years (range, 17-96 years) were identified between 2 hematology centers (Kings College Hospital, London, United Kingdom, and Haematologie, Onkologie und Klinische Immunologie, Heinrich-Heine-Universität, Düsseldorf, Germany). Diagnosis, relevant details, and sample preparation are detailed in supplemental Methods (available on the *Blood* Web site; see the Supplemental Materials link at the top of the online article).

TET2 mutation screen

TET2 (NM.001127208) mutations and polymorphisms were analyzed from bone marrow DNA using the Roche GS FLX sequencing platform (supplemental Table 1). Methodology has been described previously.¹² We sequenced 31 amplicons for 355 patients across the entire *TET2* coding region. Patient-specific “barcodes,” adapted to polymerase chain reaction (PCR) primers in the second round of amplification, were derived from Roche GS Titanium protocol. (supplemental Table 2A). Average sequencing coverage (X) across the entire *TET2* coding region was 130X and > 95% of the sequenced amplicons had sequence coverage more than 40X (> 98% with more than 10X coverage). This reliably allowed detection of mutations down to 10% relative mutation abundance (RMA), defined as the proportion of sequence reads containing the mutation. Independent PCR and Genome Sequencer (GS) FLX sequencing experiments were performed for the confirmation of mutations in all cases (giving > 500X total coverage, on average). These independent experiments reduce sampling errors for RMA calculations owing to a higher combined sequencing coverage. Sanger sequencing confirmation was performed in all cases where the mutation had a RMA of ≥ 25%. To detect low-abundance mutant clones, we resequenced 96 patient samples, including *TET2* mutant and nonmutant cases, at much greater depth (450X vs 130X), and for 13 of these patients at an average coverage of 950X. Mutations found at ≤ 10% RMA were validated in 2 additional independent PCR and GS FLX sequencing experiments, giving a combined coverage of > 800X. In 12 cases, where we had identified *TET2* mutations in the total bone marrow cells, the mutation status was also assessed in paired bone marrow, CD34⁺, and CD3⁺ T cells, and in 8 of these cases, *TET2* mutational status was also assessed in the skin biopsy.

Affymetrix SNP6 analysis

Genotyping was performed using the Affymetrix SNP6 platform and processed according to the manufacturer’s instructions (Affymetrix). Further details are available in supplemental Methods. All microarray data are available at the Gene Expression Omnibus public database less than accession number GSE2330.

JAK2 genotyping

One informative SNP (RS12345895) within the 46/1 *JAK2* haplotype was analyzed by pyrosequencing using primers and conditions described previously.²⁴

Quantitative PCR for JAK2V617F and TET expression

JAK2 V617F allele burden was determined by real-time PCR using MGB-hydrolysis probes (Applied Biosystems).²⁵ *TET* gene expression was analyzed using probes from the Universal Probe Library (Roche) and primers specific for *TET1* (NM_030625.2), *TET2* var 1 (NM_017628.3), *TET2* var 2 (NM_001127208.1), and *TET3* (NM_144993.1), as detailed in supplemental Table 2B.

Statistical analysis

Statistical analysis was performed using SPSS version 17.0 (SPSS Inc) as previously described and expanded in supplemental Methods.¹² A *P* value of less than .05 was set as the threshold of clinical significance.

Results

Classifying and mapping mutations

Samples from 320 MDS and 35 CMML patients (Table 1) were screened for *TET2* mutations at an average sequence coverage of 130X across the gene-coding region using the Roche GS FLX “next-generation sequencing” (NGS) platform. In total, 82 variants were initially identified in the *TET2* gene in 65 (18.3%) patients from total bone marrow DNA, where the proportion of sequence reads containing the variant (RMA) was ≥ 10% (supplemental Table 1). Of these, 71 *TET2* mutations predicted to alter protein structure/function were identified in 39 of 320 (12%) MDS and 16 of 35 (46%) CMML patients (*P* < .001; Table 2; Figure 1). These functionally relevant mutations were used as the primary dataset for all subsequent analyses and consisted of 43 nonsense mutations or Indel mutations, 3 splice-site mutations, and 25 nonsynonymous amino acid changes within the protein-conserved domains.^{9,11,12,23} Independent experiments confirmed mutations in all cases and allowed accurate estimates of RMA. Likewise, Sanger sequencing was used to confirm mutations where RMA was > 25%. For 2 common SNPs (rs34402524 and rs2454206), the respective heterozygous SNP sequence frequency (mean ± SD) was 47.8% ± 2.54% and 48.5% ± 2.18% within the first 96 patient samples, indicating the RMA estimates were accurate. Known SNPs and novel polymorphisms that were identified in more than 2 patients and confirmed in paired constitutional DNA, were excluded from the analysis. Significantly, 90% of the mutations identified here were previously uncharacterized.

Fourteen patients had multiple mutations (Table 2). Three patients had 2 mutations that were associated with the same amplicon being sequenced (unique patient codes [UPCs] 55, 682G, and 2644). Of these 3, UPC 682G codes had 2 mutations, which were found in the same DNA molecule and therefore resided in the same mutant clone, whereas the other 2 patients had mutations

Table 1. Clinical and biological characteristics of the MDS cohort and relationship to *TET2* mutations found at > 10% RMA

Patient characteristics	All n (%)	<i>TET2</i> mutated	<i>TET2</i> nonmutated	<i>P</i>
Number	355 (100)	55 (15.5)	300 (84.5)	
Median age, y	64.4	68.3	64.0	< .01
Range, y	(17.0-96.4)	(42.9-96.0)	(17.0-96.4)	
Sex, M/F	214/141	34/21	180/120	.06
WHO classification				< .01
RA/RCMD	131 (36.9)	17 (30.9)	114 (38.0)	
RARS/RCMD-RS	33 (9.3)	6 (10.9)	27 (9.0)	
RAEB-I/II	105 (29.6)	12 (21.8)	93 (31.0)	
sAML	35 (9.8)	4 (7.3)	31 (10.3)	
CMML-I/II	35 (9.8)	16 (29.1)	19 (6.3)	
5q- syndrome	8 (2.3)	0 (0)	8 (2.7)	
MDS/MPN-U	8 (2.3)	0 (0)	8 (2.7)	
Karyotype				.09
Favorable	192 (54.1)	32 (58.2)	160 (53.3)	
Intermediate	60 (16.9)	12 (21.9)	48 (16.0)	
Poor	71 (20.0)	7 (12.7)	64 (21.3)	
Missing	32 (9.0)	4 (7.2)	28 (9.4)	
IPSS				.25
Low/Int-1	192 (61.5)	40 (72.7)	152 (59.1)	
Int-2/High	97 (31.1)	13 (23.9)	84 (32.7)	
Missing	23 (7.4)	2 (3.4)	21 (8.2)	
Treatment				.08
None/BSC	211 (59.5)	41 (74.6)	170 (56.7)	
EPO/GCSF	29 (8.2)	1 (1.8)	28 (9.3)	
5-azacitidine	14 (3.9)	1 (1.8)	13 (4.3)	
Intensive chemo ± transplant	82 (23.1)	11 (20.0)	71 (23.7)	
Treatment data not available	19 (5.3)	1 (1.8)	18 (6.0)	

found in different DNA molecules and therefore resided in different clones or on alternate alleles, as illustrated for UPC 55 in Figure 2. Furthermore, 2 additional patients (UPCs 1190 and 2719) had multiple mutations where the higher abundance mutation was present at a frequency > 1.9-fold the lower abundance mutation, suggesting distinct mutation events in the evolution of the disease. In the remaining 9 patients, mutant clones were found at a similar RMA.

Nine mutations were found at ≤ 25% RMA in 8 patients (Table 2; Figure 1). One of these patients (UPC 55) had 2 low-level mutations (14% and 17% RMA) that resided in different sequenced molecules. Five of these patients (UPCs 750, 783, 1161, 1288, and 2545) had single mutations, and the 2 remaining patients (UPCs 2222 and 2719) had multiple mutations where 1 was above 25% RMA.

In 45 patients (82%), *TET2* mutations ranged between 15% and 61% RMA (median, 41%). In the remaining 10 patients (18%) with mutational abundance of ≥ 65% RMA, the increased mutation burden was consistent, with a genomic aberration at the *TET2* locus (see below).

Deep sequencing for low-abundance disease clones

To assess the frequency of very low-abundance disease clones, we resequenced the first 96 patients at much greater depth (450× vs 130×). We identified and verified 4 additional mutations with an RMA of 3%-6.3% (Figure 3A). Nonsense mutation Q1138X was found in a patient (UPC 2306) with refractory cytopenia with multilineage dysplasia (RCMD), for whom an additional high-frequency (39% RMA), nonsynonymous mutation had already been identified, implicating distinct mutant clones (Figure 3B). The other 3 mutations were found in 1 case of refractory anemia with excess blasts (RAEB-II; UPC 1590), one case of CMML-I (UPC 2844) and one case of RAEB-I (UPC 682), in whom no previous mutations were detected. Importantly, these additional mutations

were all located within the conserved domains of *TET2*, suggesting that these are not functionally silent events. To determine whether we were missing any frequent low-level mutation events, additional sequencing experiments of 13 *TET2* mutation-negative patients within this subgroup were performed to increase the average sequence depth to > 900×. No additional mutations were discovered at this increased sensitivity. In addition to this, no mutations were found in a control group of bone marrow DNA from 12 normal healthy persons after sequencing at depth (average, 500×).

46/1 JAK2V617F status

The frequency of the 46/1 JAK2V617F mutation predisposition genotype was analyzed in 344 of 355 patients and identified in 148 (43%) patients with an allelic frequency of 24% (130 heterozygous, 18 homozygous). This was not significantly different to levels reported in the general population.²⁴ Furthermore, there was no significant difference in the incidence of the 46/1 genotype between *TET2*-mutant versus nonmutant patients (20% vs 16% [*P* = .31] occurrence with an allelic frequency of 20% and 25% for mutant versus nonmutant patients, respectively). A screen for the JAK2V617F mutation in 200 patients from our cohort identified only one patient (MDS/MPN-unclassifiable [U]) carrying the V617F mutation with no coexistent *TET2* mutation, suggesting that in MDS such mutations were not a downstream effect of a *TET2* lesion.

TET gene expression

Expression of *TET1*, *TET2*, and *TET3* was analyzed in a subset of 116 patients [RCMD = 48, RAEB-II = 26, RAEB-I = 13, AML with MDS-related changes (sAML) = 12, MDS/MPN-U = 8, CMML-I = 3, CMML-II = 3, and RCMD with ringed sideroblasts (RS) = 3], of which 18 had *TET2* mutations. There was no

Table 2. *TET2* mutations and clinical details

UPC	WHO	Residue change/ mutation position	RMA (mean)	SD of RMA	<i>JAK2</i> haplotype	Disease progression	IPSS	Outcome
4	RAEB-II	T1726fsX18	39.5	0.7	No	No	3	Dead
55	sAML	P1937S	15.3	1.2	No	Yes	3	Dead
		P1941S	17.7	0.6				
571	CMML-I	A1344E	46.0	7.1	Yes	No	1	Dead
682G	RAEB-II	G1275E [19]	68.0	1.4	No	Yes	3	Dead
		G1282D	68.0	1.4				
		D1844G	38.0	4.2				
750	RCMD-RS	R1216Q	21.5	4.9	No	No	1	Dead
783	RA	Q1529X	25.0	1.0	No	n/a	n/a	n/a
798	sAML	H1380P	47.0	1.0	No	No	2	Dead
919	RAEB-II	A619fsX19	86.7	1.5	Yes	Yes	n/a	Dead
957	RCMD	W564X	32.5	3.5	Yes	No	1	Dead
1074	RCMD-RS	V872fsX44	38.5	0.7	No	No	0	Dead
1160	CMML-I	Splice acceptor 5' exon 9	42.3	0.6	No	No	1	Dead
1161	RCMD	R1216X [9,11]	18.0	2.6	Yes	No	1	Dead
1190	CMML-I	R1216X [9, 11]	31.5	2.1	No	No	1	Dead
		T1726fsX18	61.0	1.4				
1217	CMML-I	H1904Q	45.0	4.2	Yes	Yes	1	Dead
		E1845fsX42	42.0	1.4				
1288	RARS	Q726X	24.0	5.7	No	No	0	Dead
1326	RCMD	C1193W	49.0	3.0	No	Yes	0	Dead
1392	CMML-I	D905fsX16	38.5	3.5	Yes	No	0	Dead
1444	RCMD	L371X	47.5	0.7	No	No	0	Alive
		Y1569fsX1	37.0	2.8				
1490	CMML-I	Q1699X	49.3	1.5	No	Yes	0	Dead
		G1288V	43.3	2.1				
1586	RCMD	R1216X [11]	41.0	1.4	No	No	1	Dead
1619	RAEB-I	Q888X	44.7	0.6	Yes	No	1	Alive
		R1465X [23]	35.0	2.8				
1625	RCMD	S716X	44.7	0.6	Yes	No	0.5	Alive
1672	RCMD-RS	F1377fsX22	84.5	0.7	No	No	1	Dead
1681	RCMD	I1195V	47.0	0.0	No	No	0	Alive
1684	CMML-I	N1743fsX17	45.5	0.7	No	No	0	Alive
1702	CMML-I	V1371fsX76	94.0	1.4	No	No	0	Dead
1740	RCMD	Q1529X	38.0	4.2	Yes	No	0.5	Alive
1757	RAEB-II	Q644X	36.3	3.2	No	No	3	Dead
1857	CMML-I	R550X	45.5	0.7	No	Yes	1	Dead
1869	RCMD	Q526X	41.0	1.4	Yes	No	1	Dead
1906	RAEB-II	Splice acceptor 5' exon 5	79.0	1.4	No	No	3	Dead
1932*	RCMD	V1862fsX24	97.5	0.7	Homozygous	No	0	Alive
1980	CMML-I	A707fsX9	49.5	0.7	No	No	0	Alive
2007	RCMD	P1962L [11]	40.0	2.1	Yes	No	0	Alive
		L1360R	36.0	4.2				
		Q644X	35.5	0.7				
2222	RAEB-II	A1355P	26.0	2.0	No	No	2	Dead
		V160fsX22	19.0	2.6				
2236	CMML-I	Splice donor 3' exon 10	42.0	2.6	No	No	0	Dead
		Y1631fsX28	50.5	0.7				
2296	RCMD-RS	E259X	42.7	3.1	No	n/a	1	Dead
		Q888X	47.5	0.7				
2298	RAEB-I	E576X	43.5	0.7	No	No	1	Alive
2306*	RCMD	T1393P	39.0	2.1	No	No	0	Alive
2392	RCMD	P1962L [11]	50.0	0.5	No	No	0	Dead
2483	CMML-I	R1216X	92.7	3.1	Yes	No	1	Alive
2545	RAEB-II	F1287fsX75	20.3	7.1	Yes	Yes	3	Alive
2582	sAML	R1261G	42.0	2.8	Yes	Yes	3	Dead
2644	RCMD-RS	A1876G	46.7	1.5	Yes	Yes	0	Alive
		H1904R	46.0	1.0				
2705	sAML	P1194A	41.0	1.4	No	No	3	Alive
2719*	MDS/MPN-U	C1273F	70.7	2.1	No	Yes	0	Alive
		R1452X	20.0	3.0				
2789	RAEB-II	R1465Q	40.0	0.0	Yes	Yes	2	Dead
2859	RCMD	I1873N	39.0	1.4	Yes	No	0.5	Dead
		Q1687X	40.5	0.7				
2874	RCMD	P555fsX5 [11]	49.5	0.7	No	No	1	Alive
2984	CMML-II	H1380Y	50.0	0.6	Yes	No	2	Dead
3181	CMML-II	Q888X	43.0	1.4	Yes	No	.	Dead
3198	RAEB-II	P1629fsX31	89.5	0.7	No	Yes	2	Alive
3232	CMML-I	S1303fsX58	44.0	1.4	Yes	No	0	Alive
3266	CMML-I	Q1020fsX16	85.0	6.1	No	Yes	0.5	Alive
3338	RCMD	Q910X	37.0	1.4	No	No	0.5	Alive

TET2 mutations in the 355 MDS patient cohort. Details of all *TET2* mutations and patient context are shown. The average (mean) and standard deviation (SD) for relative mutation abundance (RMA) are shown and are based on a combined sequence coverage of > 500×. Relevant clinical details are also shown for each patient, who are referred to by a unique patient code (UPC).

*Genomic aberrations at the *TET2* locus relating to SNP6 experiments are indicated. All frame-shift mutations (fs) are, by their nature, Indel mutations and are described by the number of amino acids corresponding to the length of the new frame after the Indel event, which is indicated at the end of the mutation description. Mutations previously referenced by the 4 largest studies, to date, are highlighted and referenced. The presence of the *JAK2*V617F 46/1 predisposition genotype (haplotype), indicated by a "yes," corresponds to a heterozygous state.

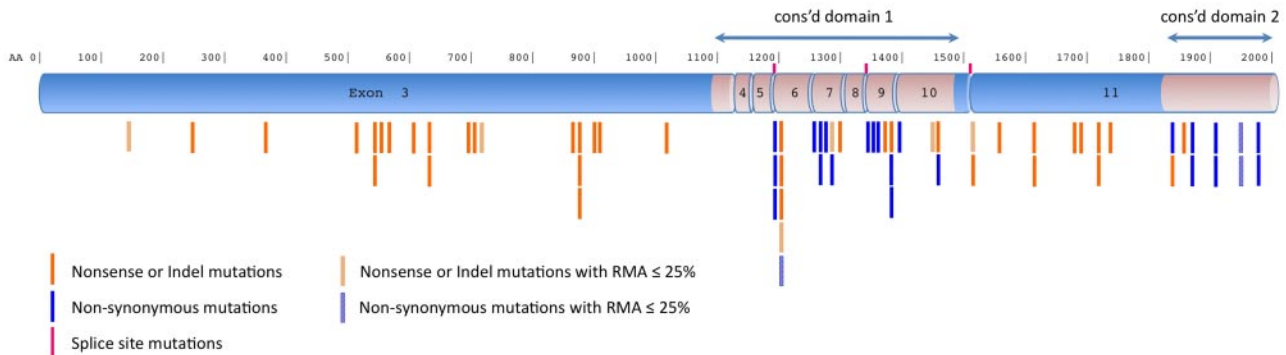


Figure 1. Seventy-one mutations in 55 of 355 persons were identified by NGS and mapped to the *TET2* coding region. Mutations were identified that existed at $\geq 10\%$ relative mutation abundance (RMA). These were mapped against *TET2* translated exons 3-11 (NM.001127208; 2002 amino acids). Classification of mutations is indicated in the figure key and includes nonsense or insertion/deletion mutations (Indels), nonsynonymous amino acid changes, and splice-site mutations, indicated by orange, blue, and pink bars, respectively. Mutation level is also defined as $>$ or $\leq 25\%$ RMA, indicated by the solid or stippled bars, respectively. Regions conserved across the TET protein family, and implicated in the conversion of 5' methylcytosine to 5' methylhydroxycytosine (5-hmC), are shown (cons'd regions 1 and 2) and correspond to amino acids 1104-1478 and 1845-2002, respectively ([11][18]).

significant difference in the mRNA levels of *TET1* ($P = .82$), *TET2* var1 ($P = .26$), *TET2* var2 ($P = .44$), or *TET3* ($P = .67$) between *TET2*-mutant and nonmutant patients (Figure 4A). Analysis of *TET* expression in different subtypes of myeloid neoplasms did not identify any significant differences ($P > .4$; Figure 4B).

Tracing *TET2* mutations through differentiation

To investigate the origins of *TET2* mutations and the selective advantage these lesions confer, we quantified the mutation load in $CD34^+$ and $CD3^+$ cells for 12 patients with *TET2* mutations (Table 3). The RMA was similar between $CD34^+$ cells (median 38.5%) and total bone marrow (median 43%). However, in UPC 1932, we detected an expansion of the *TET2* mutation from 50% to 97% RMA for $CD34^+$ and total bone marrow, respectively. In this patient, SNP array karyotyping identified a UPD at the *TET2* locus

in the total bone marrow DNA (see below). Importantly, in 9 of 11 patients, $CD3^+$ cells had a reduced, but significant, mutation load, compared with total bone marrow and $CD34^+$ cells, maintaining a mutant allele burden of $\geq 14\%$ (median, 23.5% RMA) and indicating a very early event affecting a primitive stem cell. Significantly, relevant levels of mutant clones were not detected in paired skin samples, confirming the acquired nature of these mutations.

TET2 mutations and SNP karyotyping

We analyzed 91 cases with no previous history of hematologic malignancy, using SNP 6.0 arrays. We used a size cut-off of: > 270 kb for deletions, > 500 kb for gains, and > 6.9 Mb for UPD, which were based on detecting deletions (106 kb), gains (207 kb), and UPD (2.2 Mb) at the 95th percentile in normal tissue.

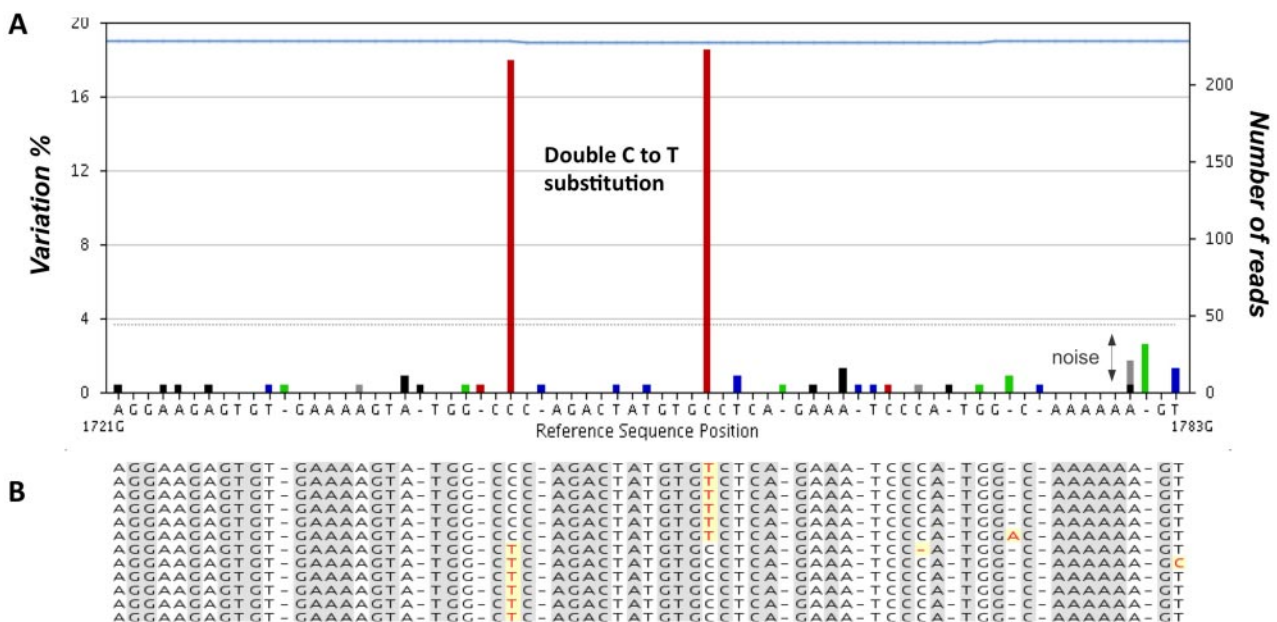


Figure 2. NGS clonal sequencing allows direct detection of independent disease clones. (A) Two C to T substitutions in *TET2* exon 11 are depicted by the dominant red bars and are found in approximately 17% of sequence reads (17% variation [left axis]) from the reference sequence [x-axis]. The sequence coverage (number of reads) across the region is shown on the right axis and is traced by the blue line. Experimental noise is shown underneath the dotted line with reads for bases A (green bars), C (blue bars), G (black bars), T (red bars), and base deletions (gray bars) at less than 4% of reads. (B) A selection of individual sequence reads lined up against each other is shown and demonstrates that the 2 mutations are found in separate molecules being sequenced and thus belong to independent clones or alternate alleles.

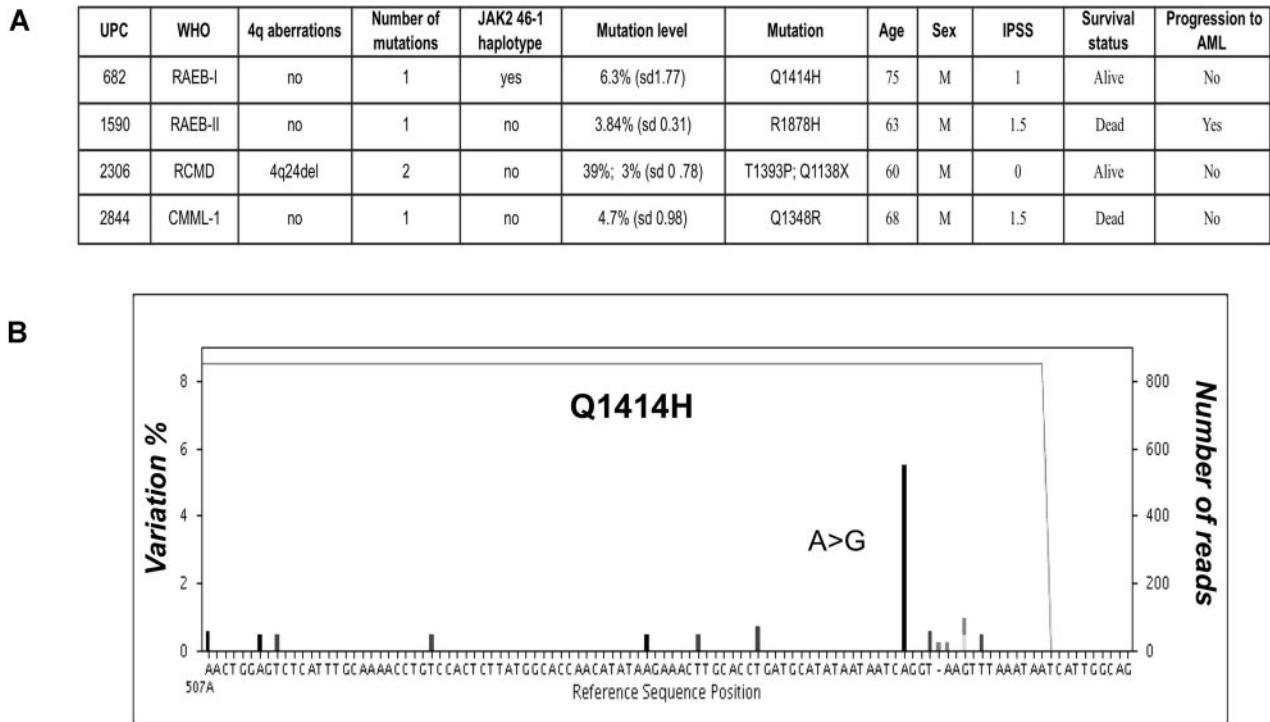


Figure 3. Deep sequencing reveals low-level mutant clones. (A) After deeper sequencing at 450× coverage for 96 patients, an additional 4 mutations were identified, which were present at between 3% and 6.3% RMA within total bone marrow DNA. One of these mutations was found in a patient with an additional mutation, which had previously been identified. Means and standard deviations for RMA corresponding to 3 independent experiments are shown. (B) Representative sequence readout for 1 of these mutations is exemplified and shows an A>G substitution, denoted by the prominent black bar, in 6% of sequence reads (6% variation from the reference sequence, left axis), leading to a Q1414H nonsynonymous amino acid change. The sequence depth is 820× across this region traced by the gray line running across the top of the graph. Experimental noise accounts for ≤ 1% of variation.

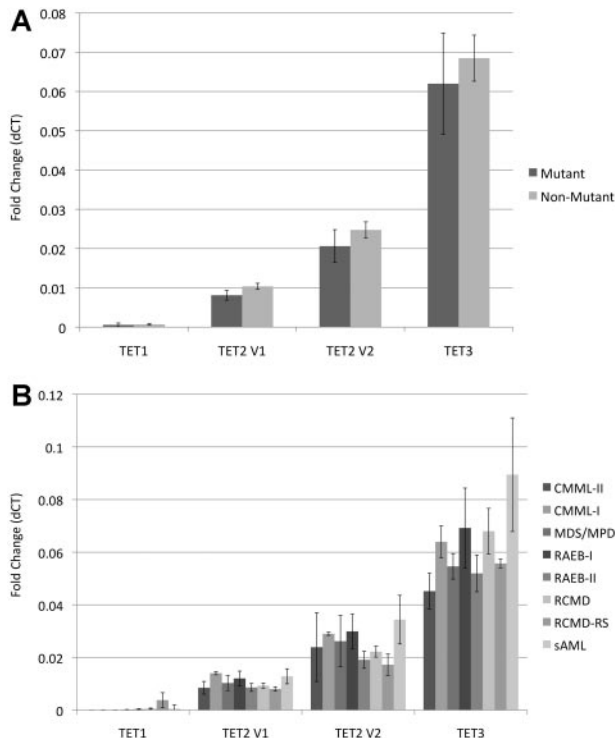


Figure 4. Bar graphs showing *TET1*, *TET2* var1, *TET2* var2, and *TET3* quantitative PCR gene-expression analysis in 116 MDS cases. (A) Gene-expression ratios in mutant (n = 18) and wild-type cases (n = 98) with standard error bars. (B) Gene-expression ratios of *TET* genes in different disease subtypes within our cohort.

DNA from bone marrow (n = 88) and peripheral blood samples (n = 3), as well as paired constitutional DNA from 12 patients, was analyzed. In addition, 155 somatic aberrations were identified in 41 patients (supplemental Figure 1). These consisted of 67 deletions (43%) of median size 2 Mb (range, 0.3-158 mb),

Table 3. Abundance of patient-specific *TET2* mutations across different tissues

Patient	% relative mutation abundance			
	BM	CD34 ⁺	CD3 ⁺	Skin
2984	50	40	22	5
2236	51/42	37/35	36/36	8/4
1932*	98	49	15	6
2859†	39/41	46/42	30/34	2/5
1619	45/35	40/30	26/17	6/2
1684	46	42	4	na
2719	71/20	60/21	54/18	na
1490	49/43	25/27	19/15	5/7
1625	45	45	26	6
1326	49	42	4	0
1444	48/37	43/27	27/25	na
2222	26	32	5	1

TET2 mutations are an early event. Twelve patients who were found to have mutant *TET2* clones within their bone marrow tissue were also screened for mutation status within paired germline (skin), CD34⁺, and CD3⁺ T-cell tissue, where available. Data for % relative mutation abundance (RMA) refers to the same mutation(s) across different tissues for any 1 patient. The median RMA for all patients is 43% for bone marrow, 38.5% CD34⁺ cells, 23.5% for CD3⁺ cells, and 5% for germline (skin) tissue. Data for individual RMA readings is an average based on at least 2 independent experiments. Patients with more than 1 mutation are denoted as n/n RMA.

*Higher mutation abundance in total bone marrow than CD34⁺ cells, consistent with a genomic aberration rendering the *TET2* locus hemizygous after differentiation.
 †High abundance of mutation within the T-cell population.

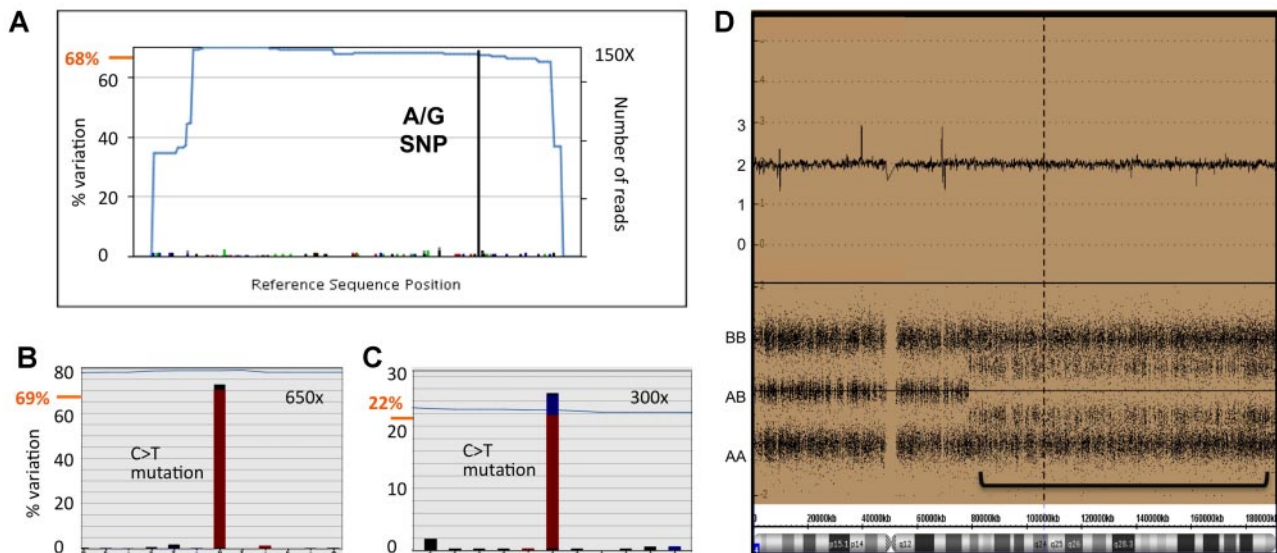


Figure 5. The quantitative nature of NGS provides greater insight into LOH events. For UPC 2719, representative sequence read-out for one SNP (A) and 2 mutations (B-C) are displayed alongside SNP array data (D), showing a UPD over the *TET2* locus. (A) An A to G SNP found in *TET2* exon 11 is depicted by the dominant black bar and is found at 68% RMA (68% variation from the reference sequence indicated on the left axis), indicative of homozygosity and LOH in a population with mixed clonality. (B-C) Likewise, a C to T mutation in *TET2* exon 7 is found at 69% RMA and another in exon 10 at 22% RMA, respectively, again indicative of multiple and distinct disease clones. The sequence coverage (number of reads) across the regions is indicated outside or inside the right axis in panel A or panels B and C, respectively, and traced by the blue lines in all cases. (D) Consistent with the sequence data, SNP6 array data show a region with LOH, highlighted by the green bar. The top panel represents a smooth copy number signal plot and a copy number of 2. The bottom panel shows individual genotypes for the SNP probes and a background of AB heterozygote calls consistent with mixed clonality.

54 gains (35%) of median size 3.7 Mb (range, 0.5-171 Mb), and 34 regions (22%) of UPD of median size 14 Mb (range, 6.9-104 Mb). Aberrations were detected in 15 RA/RCMD, 1 RARS, 12 RAEB-III, 3 CMML-III, 6 sAML, and 4 MDS/MPN-U patients. There was no significant difference observed in detected aberrations between cytogenetic risk groups (good, intermediate, and poor) and IPSS scores (low and high risk).

Of the 41 patients with genomic aberrations, 8 (20%) had *TET2* mutations. Of these, UPCs 1932 and 2306 had deletions spanning the *TET2* locus at 4q24, and UPC 2719 had UPD on 4q21.21-q35.2. UPC 1932 and 2719 had no additional aberrations, whereas UPC 2306 had additional deletions at 4q21.22-q22.1 (7.3 Mb) and 4q34.3 (492 kb). The remaining 5 patients (UPC 1326, UPC 1625, UPC 2844, UPC 2984, and UPC 1740) had no genomic aberrations associated with chromosome 4q24, but had additional aberrations as follows: UPC 1326 had UPD6q14.1-q15 (8.8 Mb), UPC 1625 had del 7q22.1 (736 kb), and UPD had 7q22.1-q36.3 (57 Mb), UPC 2844 had del 12p13.2 (277 kb), UPC 1740 had del 6q12.1-11.2 (2.1 Mb), and UPC 2984 had del 17p132.1-p13.1 (1.6 Mb), gains on 13q34 (518 kb), and 17q25.3 (751 kb) and UPDs on chromosomes 13 q13.3-q14.2 (10.4 Mb), q21.1-q21.33 (13.2 Mb), and q33.1-q34 (10.2 Mb). Nine patients with a *TET2* mutation had no detectable genomic aberrations. Of note, there was no significant difference ($P < .462$) in the total number of genomic aberrations between *TET2*-mutated and nonmutated patients.

Sequencing versus SNP array data

Patients 1932 and 2719 had *TET2* mutations with an RMA of 98% and 71/20% (2 clones), respectively, consistent with loss of heterozygosity (LOH) at the *TET2* locus (see above). Interestingly, patient 2306, with LOH on the *TET2* locus, had a mutation that was at 39% RMA, indicative of the mutation arising after the chromosomal abnormality. Consistent with these findings, patient 2306 was homozygous for SNPs within the *TET2* coding region, according to the sequencing data (not shown). Interestingly, the

sequencing data were also predictive of a further 8 LOH regions at the *TET2* locus where SNP array analysis was not informative, having mutations that were found at greater than 65% RMA and consistent SNP frequencies (Table 1; example shown in Figure 5). Furthermore, the quantitative nature of the sequencing data allowed us to confirm cellular mixes, comprising clones with LOH and ones without. As an example, patient 2719 had an A/G SNP (nucleotide position 5284, rs2454206) that was found at 68% RMA, indicative of an LOH within a clonal mix (Figure 5A). This is consistent with the mutation status of this patient with 2 mutations, one at 69% and one at 22% RMA (shown in Figure 5B-C as a representative experiment and in Table 2 as an average of 3 such experiments [71% and 20%]), suggesting the existence of 2 disease clones and the occurrence of a second mutation event after recombination. Alternatively, this may also indicate multiple mutant clones existing entirely independently, perhaps as products of a single reciprocal recombination event occurring in an early clone containing both mutations. SNP data for this patient showed a UPD at this locus with a relatively high level of background noise, in line with expectations for a mixed population (Figure 5D). This could also be the case for other patients with multiple mutations that differ significantly in frequency from each other (see above). Conversely, patients with no genomic aberrations at the *TET2* locus had a mutation frequency (RMA) consistent with heterozygosity.

Clinical associations

The association between *TET2* mutation status and clinical variables is shown in Table 1. Previous studies have shown a high incidence of *TET2* mutations in CMML patients.¹⁰ Likewise, a significantly higher proportion of CMML cases (46%) in our cohort carried the *TET2* mutation (Table 1), compared with other World Health Organization (WHO) subtypes (12%; $P < .001$). Of note, none of the 8 cases with isolated 5q- abnormalities had *TET2* mutations. Cytogenetic status, sex, WHO subtypes, transformation to AML, and IPSS score did not correlate with *TET2* mutations.

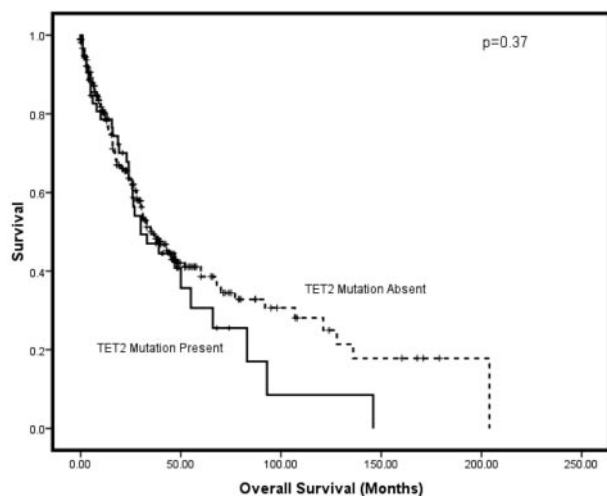


Figure 6. Survival curve depicting the impact of the presence or absence of *TET2* mutations on the overall survival of the cohort.

Similarly, no clinical association was attributed to the size of the mutant clone ($P = .921$) or the presence of single versus multiple *TET2* mutations ($P = .627$). In addition, there was no significant difference in the treatment modalities received by *TET2*-mutated and nonmutated patients. Furthermore, exclusion of patients treated with 5-azacitidine or EPO/G-CSF (erythropoietin/granulocyte colony-stimulating factor) did not alter the significance ($P = .68$). However, increasing patient age was significantly associated with a higher incidence of *TET2* mutations: 17% of patients above 50 years had a *TET2* mutation, compared with 9% of patients aged less than 50 years ($P = .02$).

Effect of *TET2* mutations on survival

At last follow-up, there were 170 deaths in the cohort, with 33 deaths being from the group of 55 patients with *TET2* mutations. The median overall survival (OS) for the entire patient cohort was 35.0 months (95% confidence interval [CI]: 26.8-43.2). There was no significant difference in OS between patients with or without *TET2* mutations (median OS: 30.0 months; 95% CI: 14.5-45.5 vs 36.0 months; 95% CI: 28.0-44.0.7; $P = .37$; Figure 6). When CMML ($n = 35$) and MDS ($n = 320$) patients were analyzed separately, no significant difference in OS was seen ($P = .12$ and $P = .42$, respectively). Among patients with a *TET2* aberration, there was no difference in OS between patients with a high ($> 25\%$ RMA) or low ($\leq 25\%$) level mutation (median OS: 26.0 months; 95% CI: 22.3-29.7 vs 35.0 months; 95% CI: 23.0-47.0; $P = .45$). In addition, homozygosity for *TET2* mutations, defined as $> 65\%$ RMA, did not affect the OS. Inclusion of the 4 low-level mutations identified after sequencing the first 96 patients at greater depth did not alter the survival outcome in that subgroup. Likewise, there was no significant difference in OS between patients based on the presence of single or multiple mutations, even when all the variants initially identified (82 variants) were included in the analysis (65 patients; $P = .15$). Furthermore, when nonmutant patients with *TET2* mRNA expression levels less than the 95th percentile ($n = 15$) were included together with cases having a *TET2* mutation, OS remained nonsignificant ($P = .77$).

Within the cohort, 289 patients with MDS were evaluated based on the IPSS score at the time of diagnosis. Among the 192 patients with low/Int-1 IPSS scores, there was no difference in OS between patients with or without *TET2* mutations (median OS: 52 months; 95% CI: 30.9-69.1 vs 65.0 months; 95% CI: 41.9-90.3; $P = .33$).

Likewise, for the 97 patients with int-2/high-risk disease, the overall survival for *TET2*-mutated and nonmutated patients was equivalent (17.4 months; 95% CI: 3.3-31.4 vs 17.2 months; 95% CI: 11.3-23.0; $P = .20$). In the low/Int-1 subgroup of patients who were untreated ($n = 162$), there was no difference in OS between *TET2*-mutated and nonmutated patients (54.0 months; 95% CI: 40.1-68.5.0 vs 68.0 months; 95% CI: 16.6-119.4; $P = .42$).

Univariate analysis performed on the subgroup of patients with normal cytogenetics demonstrated that the presence of a *TET2* mutation was associated with a significantly worse OS (32 vs 90 months; $P = .03$). However, this association could be confounded by the fact that patients with normal cytogenetics and a *TET2* mutation were older, and more likely to have CMML.

Multivariate analysis showed that the presence of a *TET2* mutation did not influence overall survival (odds ratio [OR]: 1.11; 95% CI: 0.98-1.25; $P = .08$). Advanced patient age (OR: 1.21; 95% CI: 1.05-1.38; $P = .02$), WHO classification (OR: 1.18; 95% CI: 1.07-1.38; $P < .01$), and cytogenetic subgroup (OR: 2.02; 95% CI: 1.66-2.46; $P < .01$) were the only variables significantly affecting OS (supplemental Table 3).

Discussion

TET2 mutations have been identified in a range of myeloid malignancies and are one of the most frequently acquired genetic aberrations associated with MDS.^{11,26} However, the prognostic significance of *TET2* mutations remains unclear, with some studies suggesting favorable prognosis, whereas others show no significant contribution to survival in MDS.^{11,23,27} Previous studies of *TET2* mutations have used conventional Sanger sequencing, which is constrained by a high cost and limitation in sensitivity, making this technology unsuitable for large-scale sequencing projects and inefficient in mutation screening. We have recently shown the utility of NGS technology for the detection of novel mutations across the *TET2* gene, with a mutant-allele detection sensitivity of 1%-2% in total bone marrow DNA.¹² The present study is the first to use NGS and high-resolution SNP array technology in 355 and 91 MDS/CMML patients, respectively, to simultaneously map and quantify mutant *TET2* clones and additional genomic aberrations, and correlate the findings to the clinical status of these patients.

NGS technology has recently been used to sequence the complete genome of a person with AML, primarily confirming aberrations in known disease-associated genes. We used the massively parallel pyrosequencing NGS technique (Roche GS FLX), which, because of its relatively long sequence reads, is particularly suitable for mutation screening through exonic PCR amplicons.¹² Furthermore, the increased sensitivity of NGS allows the identification of low-abundance mutant clones, enabling the study of disease evolution.²⁸ We demonstrated an overall *TET2* mutation rate of 15.5% in a cohort of 355 patients, with a *TET2* mutation rate of 12%, specifically among MDS patients (vs 46% in CMML). As these mutations were predicted to significantly alter protein function, with a majority being nonsense or Indel mutations likely to abrogate *TET2* protein function completely,^{9,11,12,23} the data strongly implicate *TET2* mutations as having a distinct, yet unknown, biologic significance, rather than being a chance event, and is consistent with a tumor-suppressor role for this protein.

A prominent finding in our study was the infrequency of low-abundance *TET2* mutant clones. Identification of such low-abundance mutant clones is important in understanding the relevance of *TET2* protein function in disease and its contribution to

clonal evolution. By increasing the sequence depth and sensitivity by 3-fold for 96 patients, we identified only 4 patients with low-abundance *TET2*-mutant clones (RMA of between 3% and 6.3%), of whom 3 had no previously detectable mutations. Moreover, by doubling sequence depth yet again for 13 of these patients, no additional *TET2* mutations were discovered, suggesting that the moderate sequencing depth achieved in the primary mutation screen was adequate in identifying the majority of mutant cases. Of note is that all the low-abundance mutations seen here were predicted to abrogate protein function and therefore were unlikely to be background or neutral events. Moreover, the existence of low, but significant, numbers of minor clones is indicative of clonal disease evolution.^{29,30} This is highlighted by several multiple-mutant cases here, where we see one dominant and one low-abundance population, suggestive of clonal competition.

Previous reports have highlighted the occurrence of *TET2* mutations in CD34⁺ cells, reinforcing the view that this aberration is an early disease lesion.^{9,11} To extend these studies, we screened CD34⁺ cells, CD3⁺ T cells, and skin tissue for *TET2* mutations that were present in the bone marrow of affected patients. In line with previous findings,¹² matched *TET2* mutations were found in both CD34⁺ and bone marrow cells at similar levels of abundance (median, 38.5% and 43% RMA, respectively). Significantly, *TET2* mutations were also found in CD3⁺ T cells (median, 23.5% RMA), confirming our previous findings.¹² Background mutation levels in skin confirmed the acquired nature of these aberrations. This indicated that *TET2* mutations were ubiquitous across myeloid and lymphoid lineages and suggests that such events occur early in hematopoiesis. Previous reports have suggested that *TET2* mutations may precede the JAK2V617F mutation in MPN.²⁷ However, after screening our samples for the JAK2V617F mutation and associated 46/1 predisposition genotype, no significant difference was observed between *TET2* mutant versus nonmutant patients, showing that the 46/1 genotype was not associated with *TET2* mutations in MDS and CMML.

We have previously identified a common region of UPD on chromosome 4q24 in MDS (8%) using 250-K SNP arrays that were subsequently shown to harbor *TET2* mutations.^{6,12} In the present study, we used high-resolution SNP6 arrays in 91 patients and found that aberrations affecting the 4q24 region were identified in only 3 of 18 patients with *TET2* mutations. These results support previous studies, which suggest that *TET2* mutations and genomic aberrations affecting the 4q24 locus are independent events.⁷ using NGS sequencing to complement SNP6 arrays, we identified 8 additional cases with SNPs and mutations at > 65% RMA, indicative of LOH. Thus, 11 of 55 (20%) of *TET2* mutant cases were associated with LOH at the *TET2* locus. Interestingly, we identified 2 patients with 4q24 UPD and *TET2* mutations at RMA 97% and 39%, respectively. The former mutation is consistent with homozygosity, whereas the latter would have had to have been acquired after the recombination event. Moreover, one patient (UPC 2719) who had UPD at *TET2* also had 2 mutations that differed significantly in abundance from each other in bone marrow (69% vs 22% RMA), a relationship that was maintained in isolated CD34⁺ and T-cell populations, suggesting that LOH and *TET2* mutations are independent events in some MDS patients.

An important feature of MDS is the increased frequency of UPD,^{5,6,8} which is an indirect measure of genomic instability.^{31,32} There was no significant difference between patients with mutant or wild-type *TET2* in the frequency of UPD or copy-number aberrations, suggesting that *TET2* defects do not contribute to general genomic instability. Furthermore, *TET2* mutations are not

linked to the 46/1 JAK2V617F mutation predisposition genotype, the only characterized genetic background associated with myeloid disease.^{24,33,34}

We did not identify any correlation between the presence of *TET2* mutations and overall survival for either MDS or CMML patients. This is in contrast to Kosmider et al,²³ who have, in 2 separate studies, shown *TET2* mutations as an independent favorable prognostic factor in MDS and associated with poor outcome in CMML patients. Any contradiction between the Kosmider MDS study and our own may, in part, be due to a relatively frequent transformation to AML in the non-*TET2*-mutated cases for the aforementioned study. Given that both Kosmider studies were retrospective in nature, it is possible that patient selection bias in either study could also have contributed to the disparity in observations. Our study benefited from the uniformity of data collected from a large cohort of patients from only 2 centers. Although the incidence of *TET2* mutations in this study was slightly lower (15.5% overall; 12% MDS and 46% CMML) than that reported by other centers,^{9,11} this could be accounted for by the observation that advanced age correlates with an increased incidence of *TET2* mutations, and that the median age in our study (64 years) was lower, compared with other studies.

The size of the mutant *TET2* clone, presence of multiple versus single mutations and *TET2* mRNA expression levels did not affect the overall survival. In addition, there is no evidence from our study that cases with homo- or hemizygous *TET2* mutations show reduced survival relative to heterozygous mutations, as might be expected if aberrant *TET2* were driving the disease independently.

The lack of correlation between the presence of *TET2* mutations and prognosis in MDS suggests that several other factors affect the MDS phenotype. Nonetheless, the acquisition of *TET2* mutations is an important event, both in the pathogenesis as well as the transformation of MDS. The latter is supported by 2 studies showing that, in MPN, the acquisition of *TET2* mutations is associated with transformation to AML, and additional mutations in genes, such as *ASXL1* or *IDH1*, are present in either a *TET2* mutant or wild-type background bone marrow culture.^{20,21} Whether mutations in *TET2* create a permissive environment or provide a “mutator phenotype”³⁵ for subsequent mutations, perhaps via alterations in the hydroxymethylome, remains to be seen.¹⁸ Studies that interrogate both *TET2* function and clonal evolution in parallel are likely to clarify the role of this gene in disease.

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Authorship

Contribution: A.E.S. and A.M.M. contributed equally and were involved with all aspects of the study's design, execution, analysis, and manuscript preparation; G.J.M. contributed to design, analysis, and manuscript preparation as well as providing project leadership; J.G. and N.L. contributed to experimental design and analysis;

G.J.M., A.K., Z.L., and N.W. provided clinical details and analysis; N.W. provided Tissue Bank support and contributed to manuscript preparation; S.M. and both C.S. and E.N. contributed to sequencing and SNP array experiments, respectively; C.S., N.G., and U.G. provided clinical samples and details; and B.P. and J.M. were involved with SNP array analysis.

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