

250K Single Nucleotide Polymorphism Array Karyotyping Identifies Acquired Uniparental Disomy and Homozygous Mutations, Including Novel Missense Substitutions of *c-Cbl*, in Myeloid Malignancies

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Abstract

Two types of acquired loss of heterozygosity are possible in cancer: deletions and copy-neutral uniparental disomy (UPD). Conventionally, copy number losses are identified using metaphase cytogenetics, whereas detection of UPD is accomplished by microsatellite and copy number analysis and as such, is not often used clinically. Recently, introduction of single nucleotide polymorphism (SNP) microarrays has allowed for the systematic and sensitive detection of UPD in hematologic malignancies and other cancers. In this study, we have applied 250K SNP array technology to detect previously cryptic chromosomal changes, particularly UPD, in a cohort of 301 patients with myelodysplastic syndromes (MDS), overlap MDS/myeloproliferative disorders (MPD), MPD, and acute myeloid leukemia. We show that UPD is a common chromosomal defect in myeloid malignancies, particularly in chronic myelomonocytic leukemia (CMML; 48%) and MDS/MPD-unclassifiable (38%). Furthermore, we show that mapping minimally overlapping segmental UPD regions can help target the search for both known and unknown pathogenic mutations, including newly identified missense mutations in the proto-oncogene *c-Cbl* in 7 of 12 patients with UPD11q. Acquired mutations of *c-Cbl* E3 ubiquitin ligase may explain the pathogenesis of a clonal process in a subset of MDS/MPD, including CMML. [Cancer Res 2008;68(24):10349–57]

Introduction

Among chromosomal aberrations involved in the pathogenesis of hematologic malignancies, somatic uniparental disomy (UPD) is increasingly recognized as a common molecular defect that results in copy-neutral loss of heterozygosity (LOH). It is likely that this defect is random and occurs either as a result of mitotic recombination or as an attempt to correct loss of chromosomal material (1). Important as a clonal marker, UPD may participate in the malignant pathologic process, particularly if UPD results in duplication of either an activating or loss of function mutation, or even perhaps an aberrant germ line genetic variant. UPD can also lead to increased or decreased gene expression through alteration

of an encoded epigenetic pattern (2). Perhaps the most well-known example of UPD involved in hematologic malignancies is UPD9p (3), which led to the identification of the Janus-activated kinase (JAK)2 V617F mutation in myeloproliferative disorders (MPD; refs. 4–6).

Routine detection of UPD was not easily possible in the past and required systematic, labor-intensive microsatellite and copy-number analysis limited in resolution. Recently, the advent of single nucleotide polymorphism (SNP) array (SNP-A) technology has allowed for the efficient and effective detection of segmental UPD in addition to other, previously undetectable microdeletions and duplications. Previously, we and others have shown that clonal UPD occurs frequently in myelodysplastic syndromes (MDS), secondary acute myeloid leukemia (AML), MPD, and MDS/MPD overlap disease entities (7–9). Other studies have shown that in patients with AML, regions of UPD can correlate with homozygous somatic mutations affecting proteins including FLT3 and CEBPA (10–12). However, systematic analysis of commonly affected areas of UPD using SNP-A technology in a broader cohort of patients with myeloid malignancies has not been performed. In this study, we have applied high-density 250K SNP-A to patients with malignant myeloid disorders to identify segmental UPD, map shared/overlapping lesions, suggest candidate genes that may be involved in disease pathogenesis, and examine relationships between UPD and corresponding clinical phenotypes.

Materials and Methods

Patients. Bone marrow aspirates and/or blood was collected from 301 patients with myeloid malignancies (mean age, 64 y; range, 17–87) seen between 2002 and 2008 at participating institutions. Informed consent for sample collection was obtained according to protocols approved by the Cleveland Clinic and Johns Hopkins University Institutional Review Boards. Samples obtained from 116 healthy individuals at the Cleveland Clinic were used as controls. In addition, a cohort of 61 CEPH (Utah residents with ancestry from northern and western Europe; CEU) HapMap individuals was used for comparison (13); however, it should be noted that the criteria used to assign membership in the CEPH population have not been specified, except that all donors were residents of Utah.⁴

DNA extraction. DNA was extracted from patient specimens using the ArchivePure DNA Blood kit (5Prime) as per the manufacturer's instructions. The concentration of the DNA was determined using a ND-1000 spectrophotometer (NanoDrop), and the quality determined by gel

Note: Supplementary data for this article are available at Cancer Research Online (<http://cancerres.aacrjournals.org/>).

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⁴ International HapMap Consortium. The responsible use and publication of HapMap data. The International HapMap Consortium Project. 2007. <http://www.hapmap.org/citinghapmap.html>.

Table 1. Characteristics of patients

	Diagnosis	Subgroup	N	% of total with abnormal SNP-A	% of total with UPD	% of those with abnormal SNP-A having UPD
Total (n = 301)	MDS (n = 140)	RA/RARS/RCMD/5q	67	63	18	29
		RAEB I/II	28	79	14	18
		Secondary AML	45	67	20	30
	MDS/MPD (n = 81)	MDS/MPDu	29	66	34	53
		CMML	27	74	41	55
		Secondary AML*	25	88	60	68
		N/a	7	100	57	57
	MPD (n = 14)	Secondary AML	7	86	57	67
	pAML (n = 66)	N/a	66	56	24	43

NOTE: Samples from 116 healthy individuals were used as controls.

Abbreviations: N/a, not applicable; pAML, primary AML.

*Includes patients who evolved to AML from both MDS/MPDu and CMML.

electrophoresis. CD3+ lymphocytes were isolated by magnetic bead separation using the RoboSep instrument (StemCell Technologies).

SNP-A analysis. The Gene Chip Mapping 250K Assay kit (Affymetrix) was used for SNP-A analysis per the manufacturer's instructions as previously described (14). Lesions identified by SNP-A were compared with the Cancer Genome Anatomy Project database⁵ and our own internal control series to exclude known copy number (CN) variants. To confirm regions of LOH detected by 250K SNP-A, we repeated samples when possible (n = 95) on ultrahigh density Affymetrix 6.0 arrays and analyzed using Genotyping Console v2.0 (Affymetrix). Signal intensity was analyzed and SNP calls determined using Gene Chip Genotyping Analysis Software Version 4.0 (GTYPE). CN and areas of UPD were investigated using a Hidden Markov Model and CN Analyzer for Affymetrix GeneChip Mapping 250K arrays (CNAG v3.0) as previously described (14, 15).

Mutational screening. Screening for the JAK2 V617F and c-MPL W515L mutations was performed using a DNA tetraprimer ARMS assay as previously described (8, 16, 17). For *FLT3*-ITD detection, capillary electrophoresis was used; PCR amplification primers were prepared as follows: ITD-F, 5'-FAM-GCAATTTAGGTATGATGAAAGCCAGC-3' and ITD-R, 5'-CTTTCAG-CATTTTGACGGCAACC-3'. After PCR amplification, 1 µL product and 0.5 µL of 1:1 diluted 500 HD Rox size standard (Applied Biosystems) were added to 13 µL formamide and analyzed using the ABI Prism 310 Genetic Analyzer (Applied Biosystems) with an extended run time of 30 min. Patients negative for the ITD mutation will have a single 332-bp fragment. For patients positive for an ITD mutation, an additional fragment ranging from 350 to 554 bp will be present.

NRAS mutational screening. Screening for mutations in exons 1 and 2 of *NRAS* was carried out using direct genomic DNA sequencing. The primer sets used were as follows: 1F, 5'-GGCCGATTAATCCGGTGT-3'; 1R, 5'-TGGGTAAGATGATCCGACA-3'; and 2F, 5'-GCAATTTGAGGGGACAAACCA-3'; 2R, 5'-TGTTAACCTCATTCCCCATA-3'. PCR conditions were as follows: 94°C for 4 min, 30 cycles of 94°C for 30 s, 51°C for 30 s, and 72°C for 30 s; 1 cycle at 72°C for 5 min. After PCR amplification, product was prepared for direct sequencing using standard conditions and analyzed using the 3100-Avant Genetic Analyzer (Applied Biosystems).

c-Cbl mutational screening. To screen patients for mutations in *c-Cbl*, direct genomic DNA sequencing was performed. Exons 7, 8, and 9 from genomic DNA were amplified using the following primer sets: 7F, 5'-ACACCAGTTGCCCTTTTAG-3'; 7R, 5'-GTCAATGGGTTCCAATGAAT-3'; 8F, 5'-GGACCCAGACTAGATGCTTTCT-3'; 8R, GAAAATACATTTCTAGAGATCAAAA-3'; 9F, 5'-CTGGCTTTTGGGGTTAGGTT-3'; and 9R, 5'-TCGTTAAGTGTTTTACGGCTTT-3'. PCR reaction conditions were as

follows: 94°C for 4 min, 30 cycles of 94°C for 30 s, 48.5°C for 30 s, and 72°C for 30 s; 1 cycle at 72°C for 5 min. The PCR product was then prepared for sequencing under standard conditions.

Results

Detection of UPD using 250K SNP-A. To define acquired somatic UPD and distinguish this lesion from germ-line, nonclonal regions of homozygosity, we first performed SNP-A analysis on a control cohort consisting of 116 healthy internal controls. Areas of homozygosity were defined as regions spanning >400 consecutive homozygous SNPs as detected by the AsCNAR (Allele-specific Copy Number Analysis using anonymous References) algorithm, which allows for the detection of abnormal clones comprising 20% to 30% of the total cell population (14, 18). The criteria of >400 homozygous SNPs is specific for our 250K array data and was chosen based on the uneven distribution of SNP probes on the array. Certain regions of the genome are not sufficiently covered by the 250K array due to relatively long stretches of LOH definable by only a few SNPs (18). This provides a greater risk of technical artifacts. Thus, the longer the series of homozygous SNPs, the higher the probability the LOH is genuine. Four hundred sequential SNPs was deemed to be of sufficient length to account for these regions. Although such an approach has a greater risk of generating false negative results, it is also more likely to eliminate technical artifacts. However, this criterion is not necessary for the 6.0 array due to the much higher, more uniform density of the SNP probes on the array (see below).

Using this criteria, and through the application of CNAG v3.0 software, segmental regions of homozygosity were identified in 3.5% (4 of 116) of the internal control cohort, comparable with previous reports (19). SNP-A analysis of CD3+ lymphocytes in these controls confirmed the germ-line derivation of these regions. Moreover, we identified a similar percentage (4.9%) of individuals harboring germ line regions of homozygosity in the CEU HapMap cohort (13). The average size of these regions in both control groups was 10.3 Mba (4.2–22.4 Mba) and occurred in small areas on chromosomes 2, 3, 4, 6, 9, 13, and 15 (Supplementary Table S14). In general, regions of homozygosity detected in controls were all short, interstitial fragments that did not regularly overlap with commonly affected regions of UPD seen in patients. Any potential

⁵ <http://cgap.nci.nih.gov>

gene candidates located in regions of homozygosity found in controls were subsequently excluded in patient analyses.

Potentially pathogenic UPDs in patients were chosen based on a rational algorithm of exclusion compared with controls. All UPD regions smaller than 20 Mba (the median size of lesions found in controls + 2× SD) were excluded from analysis unless they extended to the telomere. The rationale for the latter is based on

our analysis of control cases (none had UPD extending to the telomere) and the observation of frequent extension of UPD to the telomere in JAK2 V617F cases with UPD9p (5, 8). Although such stringent criteria may be associated with a higher false-negative rate, in the interest of minimizing false positives, we feel that this method is the best method to remove any nonpathogenic regions in patient samples. This exclusion principle was substantiated by

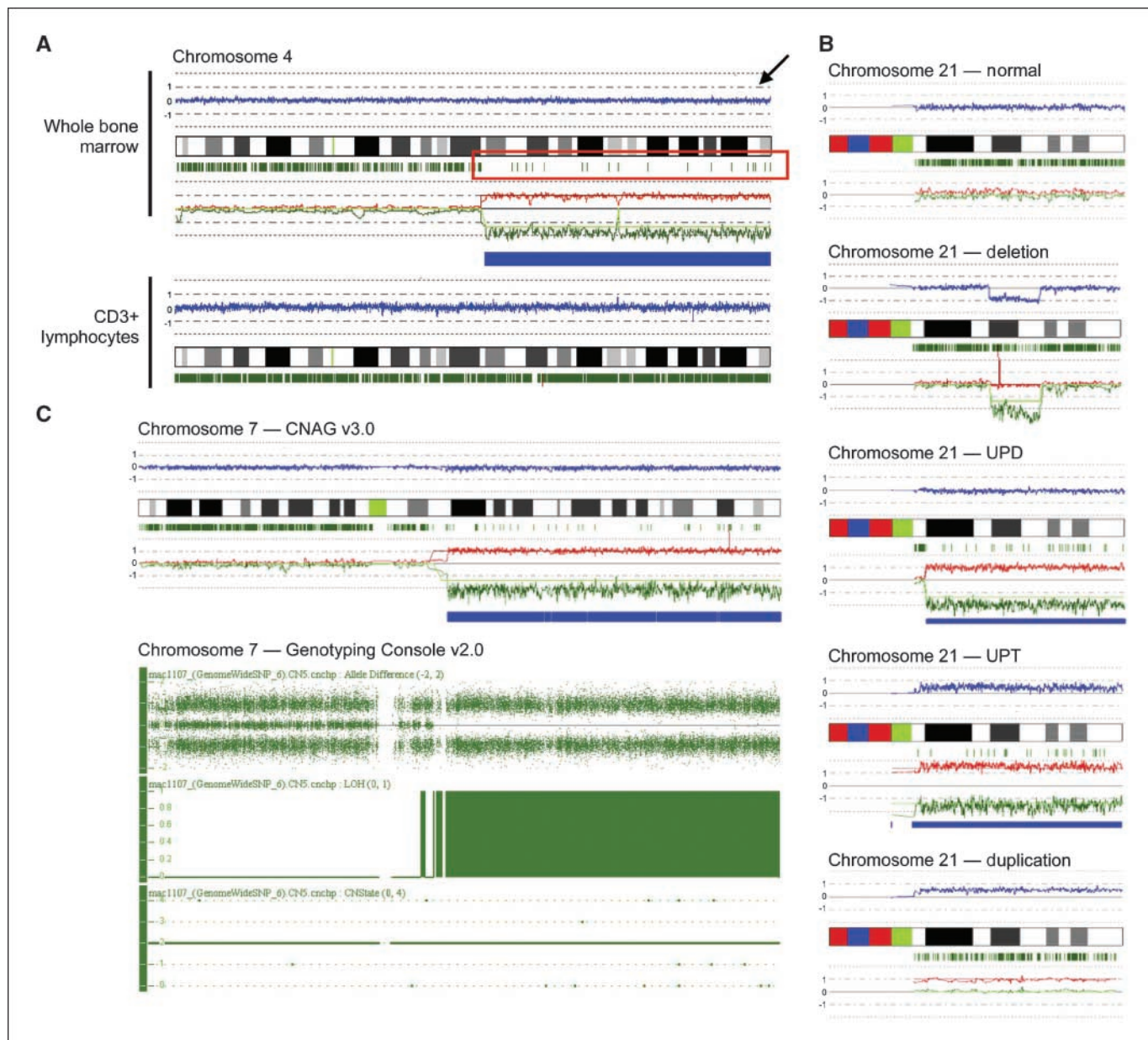


Figure 1. Detecting acquired, segmental UPD using 250K SNP-A technology. **A**, SNP-A “karyograms” of both whole bone marrow cells and CD3+ lymphocytes in one patient show the somatic nature of acquired UPD. In this representative sample, chromosome 4 is displayed. The blue line (*black arrow*) represents average CN signal intensity of SNPs on the array chip. In this instance, there are no CN variations, and thus, the blue line does not deviate from normal diploid CN. The green marks below the ideogram represent heterozygosity at particular DNA loci. In the region of UPD seen in whole bone marrow cells, drastic reduction of heterozygous loci denotes the region of UPD (*red box*). Remaining green marks in the region of UPD delineate the presence of nonclonal cells in the sample. *Green and red lines*, the AsCNAR algorithm showing the relative signal intensity for individual homologues. In CD3+ sorted lymphocytes, a normal chromosome 4 is seen. **B**, SNP-A analysis of chromosome 21 in 1 healthy control and in 4 patients shows the various types of lesions observed as follows: deletions, UPD, duplications, and in a unique patient, uniparental trisomy (UPT). **C**, comparison of Affymetrix 250K and 6.0 arrays in the detection of UPD7q for one patient. CNAG v3.0 analysis (*top*) shows clear UPD of chromosome 7 both by loss of heterozygous loci and allelic imbalance. Repeated testing on the 6.0 array and analysis using Genotyping Console v2.0 software (*bottom*) confirms the 250K SNP-A findings. Note that the Genotyping Console output includes Allele Difference, LOH, and CN variation (CNState) plots. The Allele Difference graph represents the genotypes for each individual SNP. Dots with a value of 1 represent SNPs with an “AA” genotype, whereas those with a value of -1 represent SNPs with a “BB” genotype. Dots at 0 represent heterozygous SNPs (“AB”). Complete loss of all AB SNPs indicates copy-neutral LOH. This is further shown by both the LOH and CNState graphs, which show no loss in CN but clear LOH.

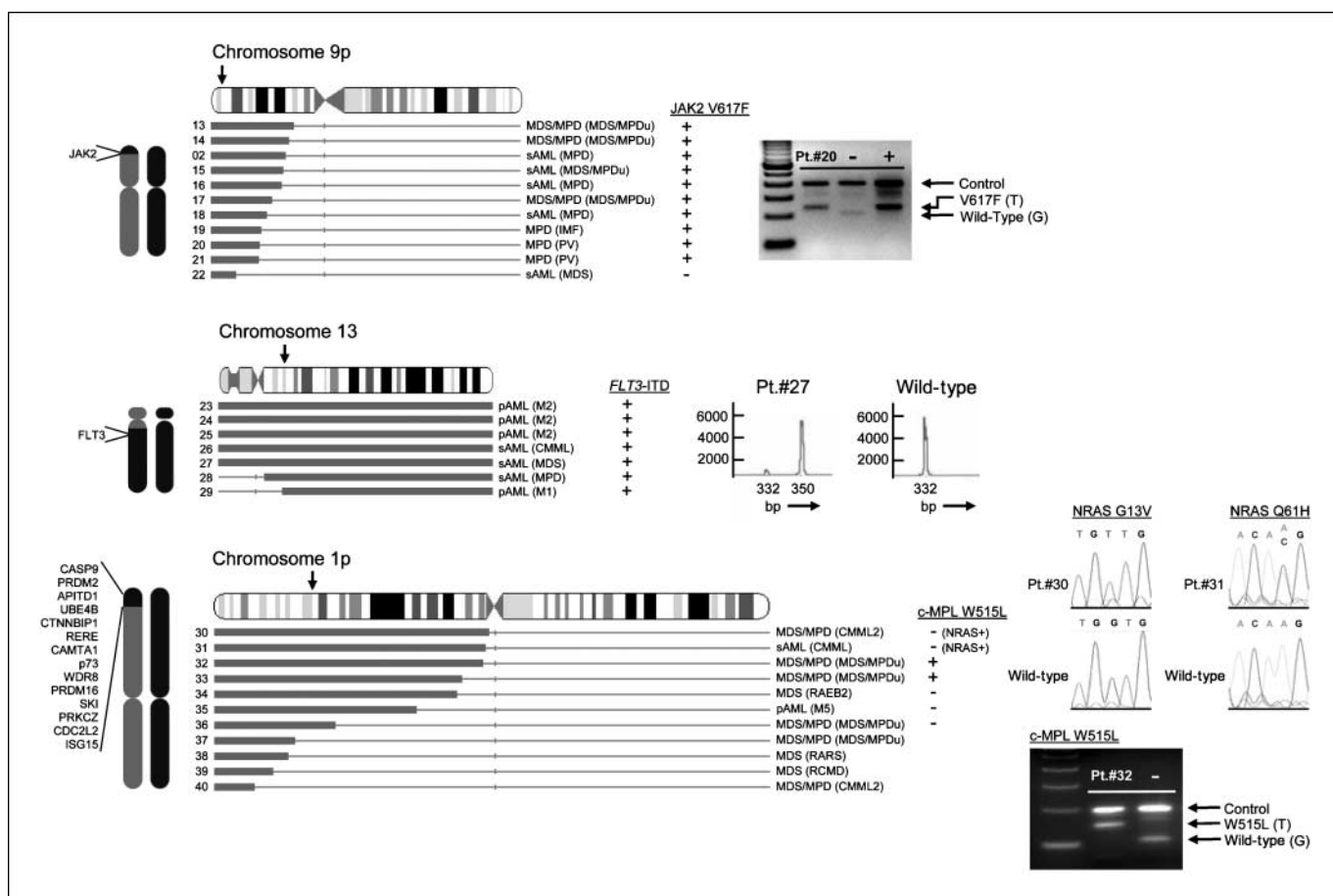


Figure 2. Acquired UPD detected by 250K SNP-A may serve as a molecular marker for mutated genes in patients. Topographical maps show regions of UPD in individual patients on chromosomes 9, 13, and 1 (left). Grey bars below the ideogram, regions affected for each patient. Individual diagnoses are listed to the right of the depicted lesion (disease subtypes are indicated in parentheses; for those patients who transformed to secondary AML, the initial presentation is given in parentheses). Arrows above the ideogram, the physical location of the *JAK2*, *FLT3*, and *c-MPL* genes, respectively. Right, mutational screening of *JAK2*, *FLT3*, *NRAS*, and *c-MPL* in patients with segmental UPD in the region of each gene confirms their homozygous status (note that sequencing analysis of *NRAS* exon 2 in patient 31 seems heterozygous due to the presence of nonclonal cells in the sample). Cartoons to the left of the maps represent the diploid chromosome set and identify the minimally overlapping region on each chromosome along with representative candidate genes in the region, regardless if the segment overlaps with known gene mutations or not. Gel images have been cropped and enhanced.

running CD3+ lymphocyte fractions from patients showing regions of UPD similar to those seen as controls as well as those with longer, telomeric UPD ($n = 18$). In agreement with findings from the control cohort described above, small interstitial regions of UPD identified in patients using CD3+ comparison were found to be germ line, whereas longer, telomeric regions of UPD were all clonal. Comparison of 250K results of both blood and bone marrow samples from the same patients showed identical results each time ($n = 6$; ref. 9).

Of the 301 total patients in our cohort, 44 had chronic myelomonocytic leukemia (CMML) or AML evolving from CMML, 140 had MDS or AML evolving from MDS, 37 had MDS/MPDu or AML evolving from MDS/MPDu, 14 had MPD or AML evolving from MPD, and 66 had primary AML (Table 1). Across diseases tested, UPD is a frequent type of chromosomal lesion, occurring in 18% of MDS cases (25 of 140), 37% of MPD cases (8 of 14, particularly UPD9p in those positive for the *JAK2* V617F mutation), and 24% (16 of 66) of all primary AML cases; however, there seems to be a slightly higher frequency of segmental UPD in patients with MDS/MPDu and CMML, occurring in 38% (14 of 37) and 48% (21 of 44) of patients, respectively. The average size of UPD in patients was 52.4 Mba. Copy number changes and UPD were identified

using CNAG v3.0 software (Fig. 1A). In contrast to deletions, which show clear loss of CN signal intensity, UPD is characterized by drastic loss of heterozygous loci but normal, diploid CN (Fig. 1B). In 1 patient with AML, a distinct area of LOH was accompanied by hyperploid CN consistent with the presence of uniparental trisomy 21 (Fig. 1B).

In many patients ($n = 95$), we have confirmed the presence of segmental LOH detected on 250K arrays, including UPD, by repeated analysis using new, ultrahigh density Affymetrix 6.0 arrays and Genotyping Console v2.0 software (Fig. 1C). Of 95 patients tested, 21 had previously identified UPD. There was remarkable concordance between the 6.0 array and the 250K arrays. All UPD regions detected by 6.0 arrays were detected by the 250K; however, it should be noted that because of the greater resolution of the 6.0 array, which allowed for the identification of smaller regions of homozygosity in controls, the average size of UPD detected in controls was reduced. Therefore, if our same "cutoff" criteria method was applied, the minimal size of pathogenic lesions would be decreased from 20 to ~10 Mba for 6.0 arrays.

Acquired, segmental UPD serves as a molecular marker for known gene mutations. Previous studies in AML have shown that regions of acquired UPD detected by SNP-A may correlate with

areas that harbor genes implicated in disease pathogenesis (12). One could hypothesize that acquired UPD leading to homozygosity of a preexisting pathogenic mutation provides an already mutant clone with an additional growth advantage. Examples of recurrent UPD regions identified in our cohort that include genes previously associated with homozygosity are presented in Fig. 2. For example, in 10 of 11 patients, including 4 of 37 patients with MDS/MPDu (11%) and 6 of 14 patients with MPD or AML evolved from MPD (43%), we found UPD9p coinciding with homozygous *JAK2* V617F mutations (Fig. 2). In addition, we also identified 7 of 143 patients (5%) with either primary or secondary AML harboring segmental UPD in the region of *FLT3*. All seven of these patients were positive for a biallelic *FLT3*-ITD mutation (Fig. 2).

In addition to *JAK2* and *FLT3*, we tested for a homozygous *c-KIT* mutation in patient 45 who had UPD in the region of 4q12 (Fig. 3), and c-MPL mutations in 7 patients with UPD1p. Although patient 45 had wild-type *c-KIT*, we did identify homozygous c-MPL W515L mutations (1p34.2) in 2 of the 7 patients with UPD1p in this region (Fig. 2). In patients 30 and 31, who were both negative for a c-MPL mutation, we hypothesized that given the localization of their lesions and their history of CMML, they may instead harbor homozygous *NRAS* (1p13.2) mutations (20). Direct genomic DNA sequencing of *NRAS* exons 1 and 2 in these 2 patients revealed missense mutations at codon 13 (G13V) in patient 30, and at codon 61 (Q61H) in patient 31.

Other previously reported gene mutations (not investigated in this study) occurring in biallelic configuration as a result of acquired UPD have included *WT1*, *CEBPA*, *RUNX1*, and *NFI* (12, 21). Thus, segmental UPD detected by SNP-A may serve as a clear molecular marker for potentially pathogenic homozygous gene mutations.

Subsequently, we hypothesized that the presence of biallelic mutations may have prognostic significance. Previous studies have shown that hemizygosity for the *FLT3*-ITD mutation due to

deletion of the remaining wild-type allele has a negative effect on overall survival (22). Therefore, as an example, we tested whether duplication of a *FLT3*-ITD mutation due to acquired UPD would also have prognostic effect. We screened all primary and secondary AML patients for *FLT3*-ITD mutations on whom DNA was available. Twenty-three of 127 patients (18%) were positive for a *FLT3*-ITD mutation, 7 of which occurred in homozygous configuration due to acquired UPD. When survival outcomes of AML patients heterozygous for a *FLT3*-ITD mutation ($n = 16$) were compared with those with a homozygous *FLT3*-ITD mutation ($n = 7$), Kaplan-Meier analysis revealed a significant detrimental effect on overall survival ($P = 0.0011$; Supplementary Fig. S1A).

Overlapping segmental UPD in patients reveals other regions that may harbor pathogenic gene mutations. In our cohort of 301, we have identified several regions of the genome commonly affected by UPD that may also harbor genes implicated in disease pathogenesis. Minimally overlapping regions of UPD found in >2 patients on any chromosome are defined in Figs. 2 and 3. Chromosomes frequently affected by UPD include 1p ($n = 11$), 4q ($n = 6$), 7q ($n = 8$), 9p ($n = 11$), 13 ($n = 7$), 17 ($n = 8$), and 21 ($n = 6$). Eighty-three percent of these lesions occurred in patients with an otherwise normal SNP-A karyotype (data not shown).

The chromosome arm most often affected was 11q, occurring in 12 of 301 patients, 6 of which had MDS/MPDu, CMML, or related disorders by clinical phenotype (Fig. 4A). Certain lesions were associated with specific disease subtypes. For instance, all patients with UPD17p had advanced stages of MDS or AML evolving with MDS, perhaps consistent with a *p53* mutation (23). Likewise, all patients with UPD17q had MDS/MPD or abnormal monocytic features, including CMML and M5 primary AML (Fig. 3). A similar scenario occurs in patients with UPD4q. Perhaps not surprisingly, all but one patient with UPD21 have either primary or secondary AML, consistent with a potential *RUNX1* mutation (20). However,

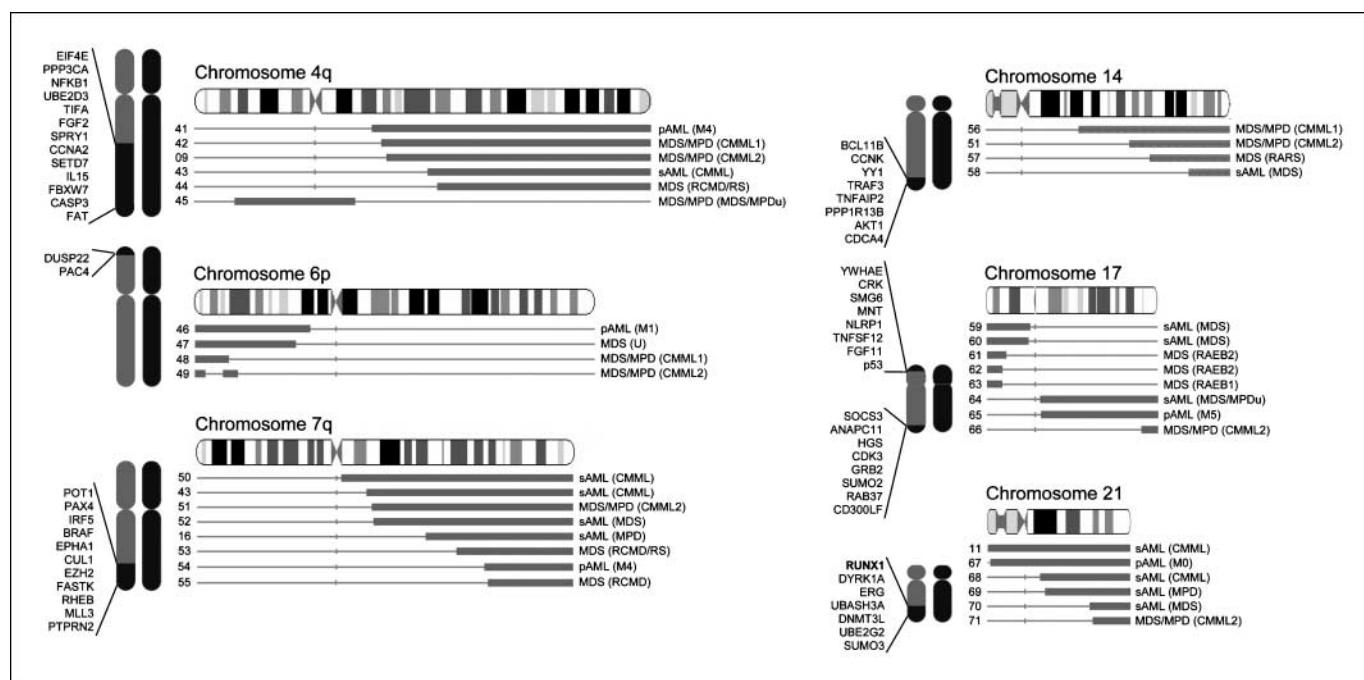


Figure 3. Genome-wide scan of overlapping UPD in patients. Topographical maps show all overlapping regions of UPD occurring in more than two patients. Gray bars, regions affected for each patient. Individual diagnoses are listed to the right of the depicted lesion. Cartoons to the left of the maps represent the diploid chromosome set and identify the minimally overlapping region on the chromosome along with representative candidate genes in each region.

greater variation exists on chromosomes 7q and 14q, where a broader range of patients harbor UPD in these regions. This is consistent with the clinical spectrum of chromosome 7q deletions in MDS, AML, and MPD.

Identification of point mutations in *c-Cbl* (11q23.3) using SNP-A. Given the prevalence of UPD on chromosome 11q, we screened for candidate genes located in this region in an attempt to identify novel mutations. Candidate genes were chosen using a simple algorithm based on physical location and function of the encoded protein. Genes were first considered according to their location within minimally affected regions of UPD. Next, genes involved with leukemia, cell signaling, cell cycle regulation, or apoptosis were selected (particularly receptor tyrosine kinases or those linked to tyrosine kinase activity). Finally, genes already known to be involved with disease processes were isolated. In this manner, potential gene candidates were minimized from a list of many to a small few.

Among several candidate genes on 11q, including *ATM*, *MLL*, and *STSI*, was *c-Cbl*, a gene that encodes an E3 ubiquitin ligase involved in the ubiquitylation and degradation of active protein tyrosine kinase receptors (24). Oncogenic mutations of *c-Cbl* occurring in a highly conserved α -helix linker joining an SH2 tyrosine kinase receptor binding domain with a RING domain have been previously identified in cell lines and in a small number of

patients with primary AML (5 of 162 in all; refs. 25–28); however, to our knowledge, no mutations occurring in other *c-Cbl* protein domains and/or in other hematologic malignancies have been found thus far. Therefore, *c-Cbl* seemed a logical target for sequence analysis.

Among our UPD11q cohort, all 12 patients harbored UPD11q in the region of the *c-Cbl* gene (Fig. 4A). Direct genomic sequencing of *c-Cbl* in these patients revealed the presence of three unique missense mutations, all occurring within or directly adjacent to the RING finger domain. In total, 7 of 12 patients were affected (Table 2). One mutation, occurring in 2 of 7 patients, resulted in the substitution of an arginine residue with either glutamine or proline at position 420 (R420Q/P). R420Q, lying just outside the RING domain, has been previously identified in 1 of 150 AML patients (28); however, we also found two additional, newly identified missense mutations, both affecting the cysteine residues of the RING finger in the remaining 5 patients (Fig. 4B and C). The cysteine residues, which bind zinc ions that form the zinc-chelating loops responsible for accommodating contacts with UbcH7, are critical for maintaining proper RING domain structure (29). In 2 of 5 patients, residue 384 was altered by substitution of a tyrosine. In the other 3 patients, residue 404 was altered by substitution of either a tyrosine (in 1 patient) or serine (in 2 patients; Fig. 4D). The presence of each somatic mutation was confirmed by

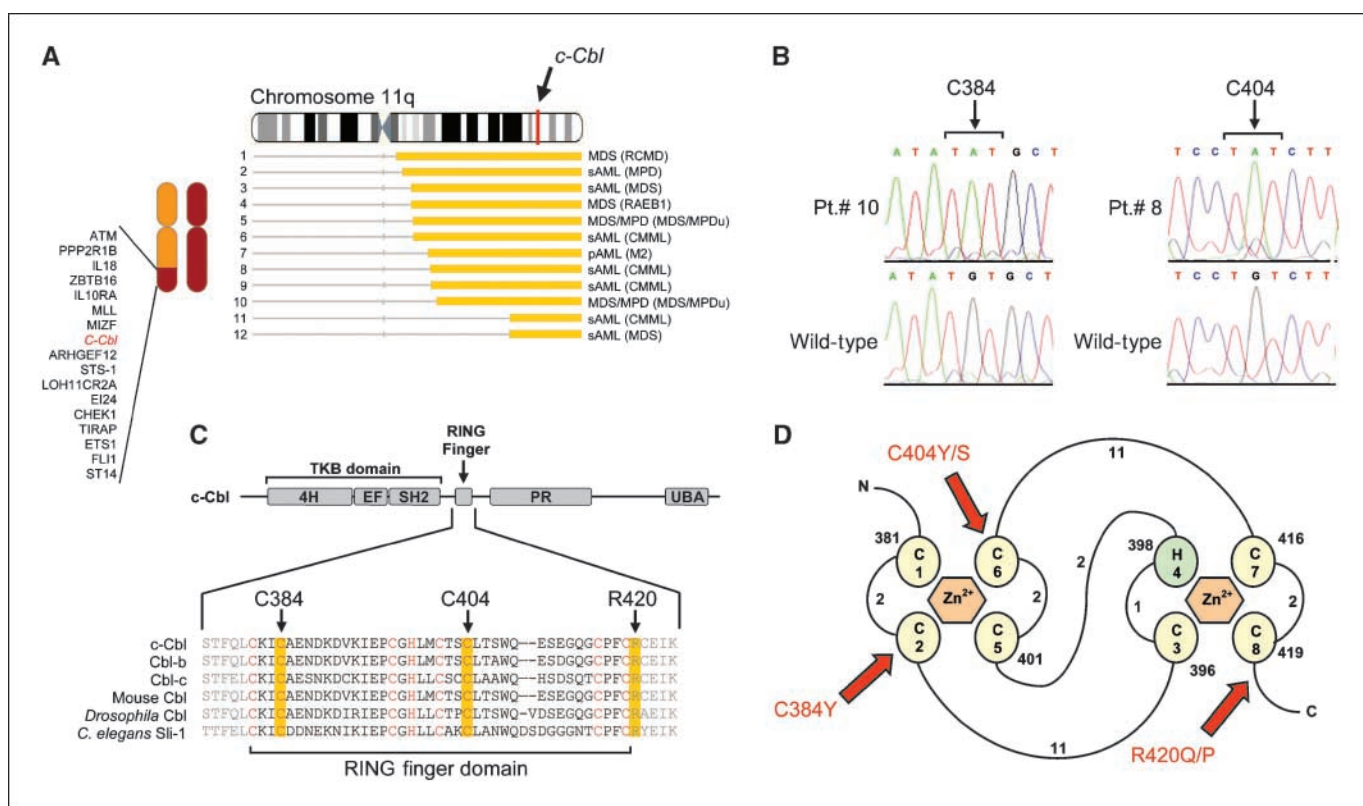


Figure 4. Identification of unique missense mutations in *c-Cbl* (11q23.3). **A**, map of individual UPD lesions on chromosome 11q and the location of the *c-Cbl* gene. **B**, direct genomic DNA sequencing of exon 8 in *c-Cbl* reveals the presence of missense mutations. Each bp change occurs in a homozygous state due to the copy-neutral LOH and results in the substitution of cysteine residues for tyrosine or serine residues at positions 384 (in patients 9 and 10) and 404 (in patients 6, 8, and 12). **C**, schematic representation showing the major domains of mammalian *c-Cbl*, primarily the tyrosine kinase binding (*TKB*) domain, RING finger, proline-rich (*PR*) region, and ubiquitin-associated domain (*UBA*). Also shown is the amino acid sequence spanning the RING finger domain of mammalian *c-Cbl* along with homologues *Cbl-b*, *Cbl-c*, and those found in *Mus musculus*, *Drosophila*, and *C. elegans* (*Sli-1*). **Red**, critical cysteine/histidine residues that make up the RING finger domain. Residues affected by mutations are highlighted in yellow. **D**, schematic of the prototypical RING finger domain found in *c-Cbl* (adapted from ref. 37; Fig. 1) **C**, cysteine residues; **H**, histidine residues. These are numbered in the order with which they occur in the domain. **Curved lines**, amino acid chains which are accompanied by numbers that denote their lengths. **Red arrows**, affected residues in patients positive for the mutations, accompanied by the resulting amino acid substitutions.

Table 2. Characteristics of patients with *c-Cbl* mutations

Pt.	Age (y)	Sex	Initial presentation	Current dx.	Current BM blast %	Metaphase karyotype	SNP-A karyotype			c-Cbl mutation
							Gain	Loss	UPD	
2	61	F	MPD (IMF)	sAML	10	46,XX,del(13)(q12q22)[15]/46,XX[5]	N	13q13.2q31.1	9p13.3pter 11q12.3qter	R420Q
4	49	F	MDS-RAEB1	N/c	7	47,XX,+21[20]	21	N	6p21.3p22.1 11q13.1qter	R420P
6	78	M	CMML2 (MP)*	sAML	60	46,XY[20]	8	N	2p22.3p23.3 11q13.2qter	C404S
8	54	F	CMML1 (MP)*	sAML	60	45,X,-X[17]/46,XX[3]	N	N	11q13.4qter	C404Y
9	69	M	CMML2 (MD)*	sAML	50	46,XY[20]	21q11.1	7q21.1	4q21.21qter 11q13.4qter	C384Y
10	82	M	MDS/MPDu	N/c	0	46,XY[20]	N	3p14.2, 7p21.2	11q13.5qter	C384Y
12	44	F	MDS-RAEB2	sAML	14	46,XX[20]	15q13.3	N	11q22.3qter	C404S

NOTE: Patient numbers correspond to numbers/lesions presented in Fig. 4A.

Abbreviations: Dx, diagnosis; MP, myeloproliferative phenotype; MD, myelodysplastic phenotype; sAML, secondary AML; N/c: no change.

*Refers to FAB CMML classification based on WBC counts (ref. 36).

bidirectional DNA sequencing of multiple isolates and comparison against CD3+ sorted lymphocytes when available.

Given the preponderance of *c-Cbl* mutations in patients who exhibit MDS/MPD features, we tested for the presence of monoallelic mutations in all patients with a diagnosis of MDS/MPD, including those negative for UPD11q; however, in 40 such patients screened, no *c-Cbl* mutations were identified.

Discussion

SNP-A technology, with its improved resolution and ability to detect UPD, increases the overall detection rate of chromosomal abnormalities and complements metaphase cytogenetics (MC) in the delineation of chromosomal lesions associated with hematologic malignancies. Recently, we and others have shown that in MDS, MDS/MPD, and MDS-derived AML, previously unrecognized chromosomal defects detected by SNP-A have similar effect on prognostic variables including overall survival as those lesions detected by MC (7, 9). In addition to its potential clinical implications, SNP-A karyotyping is also an excellent investigative tool, facilitating delineation of invariant chromosomal defects and, in turn, corresponding genes that may be implicated in the pathogenesis of hematologic malignancies.

Using 250K SNP-A and CNAG v3.0 software, we have identified in a large cohort of patients with MDS, MDS/MPD, MPD, and AML, a significant proportion of UPD, particularly in MDS/MPD. Often times, overlapping regions of UPD correlated with the presence of homozygous gene mutations, including *FLT3*-ITD in 7 of 7 cases with UPD13q, accounting for 30% of all cases positive for a *FLT3*-ITD mutation ($n = 23$), and *JAK2* V617F mutations in all MDS/MPD and MPD cases characterized by UPD9p. In addition, we identified *c-MPL* or *NRAS* point mutations in four patients with UPD1p. *c-MPL* mutations were associated with MDS/MPD, and we believe that this is the first report of biallelic *c-MPL* or *NRAS* mutations occurring as a result of acquired UPD. Although not formally tested here, one could also hypothesize that, in accordance with previous studies (11, 12), there are homozygous gene mutations affecting either *RUNX1* or *CEBPA* in those patients with acquired UPD on chromosomes 21 ($n = 6$) or 19 ($n = 2$), respectively.

Systematic mapping of copy-neutral LOH has allowed for the identification of other frequently shared areas of UPD that are

currently not attributable to any known gene mutations in myeloid malignancies. Identification of novel mutations in these regions may explain the pathophysiology of those individual cases affected. Examples of these commonly affected areas include segments on chromosomes 4q, 6p, 7q, 14, 17, and 11q. Various genes potentially involved in the pathogenesis of leukemia are located in these regions including *EIF4E*, *CCNA2*, and *FGF2* on 4q; *DUSP22* and *PAC4* on 6p; *BRAF*, *CUL1*, and *EZH2* on 7q; *TNFAIP2* and *AKT1* on 14; and *p53* and *GRB2* on 17.

When we examined patients with UPD11q, we noted several common clinical phenotypic trends, including history of MDS/MPD, the presence of monocytic blasts, or increased numbers of differentiated monocytes, reticulin fibrosis, and propensity to AML transformation. Monoallelic mutations in the α -helix linker domain of *c-Cbl* disrupt binding of c-Cbl to E2 ubiquitin-conjugating enzymes and, thus, deregulate proper ubiquitylation and degradation of phosphorylated protein tyrosine kinase receptors in animal models (30) and in a small number of patients with AML (25–28). We hypothesized that mutated, dysfunctional c-Cbl could lead to deregulated activity along proliferative transduction pathways consistent with the phenotype of MDS/MPD. For example, other TK receptors including CSF1R, platelet-derived growth factor receptor β and epidermal growth factor receptor have all been described to be inactivated by members of the Cbl ubiquitin ligase family either directly or via binding to growth factor receptor binding protein 2 (31–35).

In addition to 2 patients harboring biallelic R420Q/P mutations previously reported, we identified new, biallelic mutations of *c-Cbl* in 5 additional patients, which cause single amino acid substitutions affecting the highly conserved cysteine residues at positions 384 and 404 of the RING domain. Analysis of nonclonal T-cells from these cases confirmed the acquired nature of these mutations. However, unlike previous reports, sequencing targeted to identify RING and linker domain monoallelic mutations in heterozygous configuration did not identify patients with monoallelic *c-Cbl*. Based on our observation of only biallelic mutations in our MDS/MPD patients, we hypothesize that the pathogenic role of *c-Cbl* mutations may differ in AML versus MDS/MPD. Alternatively or additionally, the discrepancy between our findings and those previously reported in AML may be complicated by technical problems that arise when distinguishing between true monoallelic

mutations in one clone and a mixture of abnormal clonal cells harboring biallelic lesions (e.g., due to UPD) and nonclonal, wild-type cell contributions. We have shown that patients with *c-Cbl* mutations all have corresponding UPD11q; perhaps SNP-A applied to the previously described AML patients would also reveal UPD11q.

Patients with *c-Cbl* mutations showed a great deal of diversity with regard to clinical phenotype (Table 2). Although *c-Cbl* mutations have been previously linked to primary AML only (27, 28), all patients positive for *c-Cbl* mutations in this study had developed from a previous hematologic disease. In addition, some patients with *c-Cbl* mutations have additional chromosomal aberrations and/or genetic mutations that could modify the resultant clinical phenotype. For example, one patient (patient 2; Table 2) was JAK2 V617F+. Another two patients (patients 6 and 12; Table 2) were NPM1+. In addition, related genes *Cbl-b* (3q13.11) and *Cbl-c* (19q13.31) may harbor mutations with pathogenetic significance, which could explain other cases of myeloid malignancies at the molecular level. These results serve to illustrate that advanced myeloid malignancies likely have many cooperating genetic and epigenetic lesions, and that high-resolution SNP arrays provide insight into sorting out affected molecular pathways.

Systematic analysis of UPD also provides many important clues as to the pathogenesis that may operate in MDS, MDS/MPD, or AML. We have noted that certain chromosomes are as frequently affected by UPD as deletion (e.g., 7 and 21). However, UPD may suggest a different mechanism and type of gene affected (e.g., duplication of an activating mutation) compared with deletion, which can lead to haploinsufficiency or "unmasking" of a pre-existing inactivating mutation in a tumor suppressor gene. This may explain why some chromosomes are more frequently affected by deletion than UPD. For example, although we have identified 35 cases with deletions involving the minimally affected region on chromosome 5q, only 1 example of UPD was found in the corresponding area, suggesting that affected genes in this region do not provide an additional selection advantage if duplicated.

It should also be noted, however, that not all UPD is likely to be pathogenic but may simply serve as a marker of clonality and/or underlying previous DNA damage (1). To enable analysis of as many samples as possible and determine clinical applicability of SNP-A, most patient samples were run using whole bone marrow and/or blood. Most often the admixture of nonclonal with clonal cells is not an issue in samples with advanced MDS, AML, and CMML as the percentages of nonclonal cells are low. However, in low-grade MDS, it is possible that the admixture of clonal cells with nonclonal cells may be much lower, and thus, some UPD may have been undetected by our analysis.

In sum, our investigations provide a proof of principle that delineation of shared areas of UPD identified by SNP-A may facilitate the identification of genes affected by pathogenic mutations and potentially explain clinical phenotype. Segmental UPD seems to be a common clonal lesion in myeloid malignancies. As more sensitive techniques and study of additional myeloid malignancy cases accrue, comparison of individual UPD lesions with current clinical, molecular, genetic, and cytogenetic prognostic schemes in larger cohorts of patients will likely provide additional important diagnostic, prognostic, and translational biomarkers.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

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