Abstract 7022: Mutational landscape and clinical characterization of over 17000 patient samples with myeloid malignancies using real world data

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Background

- Myeloid neoplasms represent a broad spectrum of hematological disorders.
- Somatic mutation status in key driver genes is important for diagnosis, prognosis and treatment.
- We summarize findings from 17181 clinical samples from 16133 patients analyzed by a next generation sequencing (NGS) laboratory developed test targeting 50 myeloid associated genes.
- Samples were analyzed comprehensively and as part of individual cohorts specific to acute myeloid leukemia (AML), myelodysplastic syndrome (MDS), and myeloproliferative neoplasms (MPN)

Methods:

- Whole blood or bone marrow samples from patients with cause-for-testing for hematological symptoms were submitted for analysis by a referring clinician.
- DNA was extracted and assayed by a targeted, NGS panel to detect and report single nucleotide variants and small indels within 50 genes associated with myeloid malignancies.
- Sequenced on an Illumina MiSeq or NextSeq (Illumina, San Diego, CA).
- Multiple somatic variant classes were called including single nucleotide variants, insertions, and deletions. Copy number variants in the gene *KMT2A* were also reported.
- Results were reviewed, orthogonally confirmed unless previously validated, and reported by clinical laboratory directors.
- Disease status or symptoms used in this study were taken from test requisitions for each patient.

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Figures

Figure 1. Patient demographics and clinical mutations detected

Characteristics	Results					
Total patient samples, n	17181					
Total patients	16133					
Age, mean (SD)	67.17 (15.67)					
Gender, n (%)						
Male	9525 (55.4)					
Female	7656 (44.6)					
Number of clinical indications, mean (SD)	1.26 (0.51)					
Select reported clinical indications ^a , n (%)						
AML	2225 (13.0)					
MDS	4176 (24.3)					
MPN	2018 (11.6)					
MDS/MPN	590 (3.43)					
Symptoms only (including anemia, thrombocytopenia and leukopenia)	8159 (47.5)					
Patients with positive NGS Result, n	12102 (70.4)					
Patients with negative NGS Result, n	5079 (29.6)					
Tier I-III mutations detected, n	34581					
a. Patients with multiple clinical indications are counted for all indications on test requisition form.						



A) Patient demographics for all samples included in this analysis. The number (n) and proportion (%) of patients for each factor are listed. All clinical indications were included as noted within the test requisition form.

B) Distribution of the number of clinically relevant mutations detected per clinical sample. The mean number of mutations per test was 1.98 (95% CI: 1.95 to 2.01) with a range of 0-14.

C) Boxplot showing number of mutations by detected by age of the patient at testing. The number of clinically relevant (Tiers I-III) mutations identified for each of the 17181 patient samples was compared to the age of the patient at testing and were found to be positively correlated (Spearman's rank correlation coefficient, p=0.30, p<0.0001). Text in blue is the mean number of mutations per patient for that particular age group



Associations are shown for the entire sample cohort regardless of indication. Associations are colored by odds ratio showing a gradient from orange (co-mutation) to white (no association) to purple (mutual exclusivity) as shown on the legend on the left. Circular labels are sized by the p-value calculated from the Fisher Exact test with the largest size corresponding to the lowest probability. Associations that are significant after correction for multiple hypotheses are indicated by *. Pair-wise mutation analyses found 21 mutually exclusive pairs including between genes associated with RNA splicing (SF3B1, SRSF2 and U2AF1; OR<0.29, p < 0.001) suggesting possible candidates for targeted therapy. For gene pairs co-mutated, the largest significant effect size was between the AML driver mutations FLT3 and NPM1 (OR = 23.4, p < 0.001).







During the evaluation period, 1021 patients were tested at multiple time-points. These patients provided an average of 2.24 samples (range 2-7) which were taken an average of 233 days (range 0 – 1141 days) after the initial sample. 498 (48.8%) patients showed loss or gain of a mutation between sample dates, potentially the result of tumor evolution and/or therapeutic intervention. The following four individual case studies show specific examples of possible changes in tumor evolution and clonal architecture. For each case study, the variant allele frequency (VAF) of the detected mutations is shown on the y-axis and is plotted against time (days) on the x-axis. Each unique variant measured is colored according to the key and is described according to the gene and nucleotide change observed. Dots represent the test dates. Dots with time = 0 days represent the first test date. VAF = 0 means that a variant is not detected on the particular test date. (A) Patient with 4 unique variants detected over 4 test dates spanning 200 days. Changes in tumor fraction may be represented by changes in VAF for mutations in ASXL1 (c.2077C>T), IDH2 (c.419G>A), and SRSF2 (c.284C>A). These 3 mutations are absent by test date four (200 days). (B) Patient with 10 unique variants detected over 7 test dates spanning 465 days. Mutations in ASXL1 (c.1900_1922del23), DNMT3A (c.2645G>A), and TP53 (c.830G>A) are present in all 7 test dates. Over the course of testing, the patient shows gains and losses of mutations in ASXL1 (c.1934dupG), IKZF1 (c.503_505delinsAT), NRAS (c.35G>A) and TP53 (c.358A>G, c.524G>A, c.733G>A, and c.743G>A) which may correspond to the evolution of individual sub-clones. (C) Patient with 6 unique variants detected over 5 test dates spanning 398 days. Patient shows the loss of a mutation in CEBPA (c.912_913insTTG) after 27 days along with three other mutations in CEBPA (c.191delinsGG), KIT (c.2447A>T), and TET2 (c.4054delG). The same CEBPA (c.912_913insTTG) mutation reemerges at 398 days alongside a mutation in CSF3R (c.1853C>T) potentially representing an expansion of a new sub-clone. (D) Patient with 10 unique variants detected over 7 test dates spanning 878 days. Four mutations are ever-present over the course of testing: ASXL1 (c.1934dupG), RUNX1 (c.964_965dupTC), SRSF2 (c.284C>A), and STAG2 (c.3097C>T). Six other mutations are either gained or lost on subsequent dates possibly reflecting gains or losses of individual sub-clones. The differing VAFs observed for mutations detected on the same test date suggest a complex clonal architecture for this patient.

Mutation 1	Mutation 2	ELN Prognosis	Samples with both	Mutation 1 only	Mutation 2 only	Samples with neither	Odds Ratio	Adjusted p value
FLT3-ITD ^{low}		Intermediate						
FLT3-ITD ^{high}		Adverse						
FLT3-ITD ^{low}	NPM1	Favorable	50	77	171	1927	7.3	1.17 x 10 ^{-19**}
FLT3-ITD ^{high}	NPM1	Intermediate	6	11	215	1993	5.1	0.013**
FLT3-ITD	NPM1		56	88	165	1916	7.4	5.24 x 10 ^{-22**}





Samples were categorized into cohorts for Acute Myeloid Leukemia (AML), Myelodysplastic Syndrome (MDS) and myeloproliferative neoplasms (MPN) based upon the clinical indication(s) isted on the test requisition form. A) % of patients with a mutation in each gene for AML cohort. 2225 (13.0%) samples had an indication for AML Mutations that are diagnostic for AMI were observed as follows: FLT3 in 12.0% (267 of 2225), NPM1 in 10.0% (221 of 2225), and RUNX1 in 14.5% (322 of 2225) of AML samples. FDA approved targeted therapies associated with mutated genes in the panel include midostaurin (FLT3), ivosidenib (IDH1), and enasidenib (IDH2). IDH1 was observed in 12.3% (273 of 2225) and IDH2 in 8.0% (180 of 2225) of AML patients.

B) % of patients with a mutation in each gene for MDS cohort. 4176 (24.3%) samples had an indication for MDS. SF3B1 mutations which are diagnostic for MDS with ring sideroblasts were found in 12.4% (516 of 4176) of MDS patients. C) % of patients with a mutation in each gene for MPN cohort. 2018 (11.6%) of samples had an indication for MPN. The canonical MPN mutations were observed as follows: JAK2 (606 of 2018, 30.0%), CALR (154 of 2018, 7.6%), and MPL 2.5% (50 of 2018, 2.5%).

Table 1. NPM1 and FLT3 internal tandem duplicate mutation status amongst patients with an indication of AML

The European Leukemia Net (ELN) assigns a favorable, intermediate, or adverse risk based on a patient's FLT3 internal tandem duplicate (ITD) and NPM1 mutation status The clinically favorable co-mutation of NPM1 with FLT3 internal tandem duplicate was significantly enriched in the AML population. FLT3-ITD^{high} denotes allele ratio > 0.5 while FLT3-ITD^{low} denotes allele ratio <= 0.5.

** denotes statistical significance.

Table 2. NPM1 and ASXL1/RUNX1 mutation status amongst patients with an indication of AML

ation 1	Mutation 2	ELN Prognosis	Samples with both	Mutation 1 only	Mutation 2 only	Samples with neither	Odds Ratio	Adjusted p value
XL1		Adverse						
NX1		Adverse						
XL1	NPM1	Favorable	18	428	203	1576	0.33	Mutual exclusive: 3.8x10 ^{-4**}
NX1	NPM1	Favorable	8	314	213	1690	0.20	Mutual exclusive: 3.9x10 ^{-5**}
XL1	RUNX1		138	308	184	1595	3.88	Co-mutation: 1.5x10 ^{-19**}

The European Leukemia Net (ELN) assigns an adverse risk based on ASXL1 or RUNX1 mutation. Co-mutation of ASXL1 or RUNX1 with NPM1 mutation status gives a favorable outcome however this comutation was significantly less common than expected. In contrast, co-mutation of ASXL1 and RUNX1 was enriched in AML (OR = 3.9, p < 0.001). Further investigation is required to determine whether this co-

mutation is clinically significant.

** denotes statistical significance.

Summary

- 17181 patient samples assessed for somatic mutations in 50 genes.
- Disease status used in this study were taken from the test requisitions for each patient.
- 47.3% of patient samples had 2 or more somatic mutations. Pair-wise mutation analyses found 21 mutually exclusive
- pairs including between genes suggesting possible
- candidates for targeted therapy.

- The clinically favorable co-mutation of *NPM1* with *FLT3* internal tandem duplicate was significantly enriched in the AML population.
- The clinically favorable co-mutation of *NPM1* with *ASXL1* or *RUNX1* was significantly less common than expected in the AML population.
- Co-mutation of *ASXL1* with *RUNX1* significantly enriched in the AML population.
- Individual case studies of patients tested at multiple timepoints show evidence of tumor evolution and/or therapeutic intervention.

Conclusion

Parallel testing of multiple genes in addition to the canonical driver mutations encompasses the mutations contributing to the etiology of myeloid neoplasms. Consistent patterns of mutations are routinely observed that can help the clinician tailor the treatment and chart the progression of myeloid disease for each patient.