**FLT3-ITD Allelic Burden and Acute Promyelocytic Leukemia Risk Stratification**

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**Simple Summary:** Around 12–38% of acute promyelocytic leukemia (APL) patients carry the FLT3-ITD mutation, which has been associated with several poor-prognosis indicators such as high white blood cell counts, M3v variant morphology, and the bcr3 isoform. We aimed to retrospectively study the impact of FLT3-ITD mutations in APL patients in regard to clinical features, treatment courses, and outcomes. We demonstrate that Sanz high-risk status APL correlates with high FLT3-ITD allelic burdens, with every 1% increase in allelic burden correlating with a 0.6 × 10^9/L increase in white blood cell count (WBC). The presence of FLT3-ITD was associated with decreased remission rates and higher 5-year mortality from the time of diagnosis. These findings provide novel revelations regarding the features of FLT3-ITD APL, particularly in regard to allelic burden, that warrant further study.

**Abstract:** The significance of FLT3-ITD in acute promyelocytic leukemia (APL) is not well-established. We performed a bi-center retrospective study of 138 APL patients, 59 (42.8%) of whom had FLT3-ITD. APL patients with FLT3-ITD had higher baseline white blood cell counts (WBCs) (p < 0.001), higher hemoglobin, (p = 0.03), higher aspartate aminotransferase (p = 0.001), lower platelets (p = 0.004), lower fibrinogen (p = 0.003), and higher incidences of disseminated intravascular coagulation (p = 0.005), M3v variant morphology (p < 0.001), and the bcr3 isoform (p < 0.001). FLT3-ITD was associated with inferior post-consolidation complete remission (CR) (p = 0.02) and 5-year overall survival (OS) of 79.7%, compared to 94.4% for FLT3-WT (wild-type) (p = 0.02). FLT3-ITD was strongly associated with baseline WBCs ≥ 25 × 10^9/L (odds ratio (OR): 54.4; 95% CI: 10.4–286.1; p < 0.001). High FLT3-ITD allelic burdens correlated with high-risk (HR) Sanz scores and high WBCs, with every 1% increase in allelic burden corresponding to a 0.6 × 10^9/L increase in WBC. HR APL was associated with a 38.5% increase in allelic burden compared with low-risk (LR) APL (95% CI: 19.8–57.2; p < 0.001). Our results provide additional evidence that FLT3-ITD APL is a distinct subtype of APL that warrants further study to delineate potential differences in therapeutic approach.

**Keywords:** APL; leukemia; FLT3-ITD
1. Introduction

First described by Norwegian hematologist Leif Hillestad in 1957 through a series of three cases, acute promyelocytic leukemia (APL) was aptly named for its predominance of promyelocytes and deadly coagulopathy characterized by “a very rapid fatal course of only a few weeks’ duration” [1]. Today with the advent of all-trans retinoic acid (ATRA) and arsenic trioxide (ATO), APL is highly curable if treatment is initiated promptly. APL is classified as the French–American–British (FAB) subtype M3 of acute myeloid leukemia (AML), classically arising from a balanced reciprocal translocation between chromosomes 15 and 17. The fusion between the promyelocytic leukemia (PML) and retinoic acid receptor alpha (RARα) genes results in the PML–RARα rearrangement t(15;17)(q24;q21), leading to the disruption of the RARα-regulated maturation of myeloid progenitors at the promyelocytic stage [2]. APL comprises less than 10% of AML, with an estimated 0.01/100,000 incidence in Western countries, affecting men and women equally [3].

FMS-like tyrosine kinase 3 (FLT3) is a proto-oncogene implicated in leukemogenesis. The FLT3 ligand binds to the extracellular FLT3 receptor, inducing homodimerization that potentiates a downstream signaling cascade involved in the regulation of the proliferation, differentiation, and apoptosis of early myeloid and lymphoid progenitor cells. The most common FLT3 aberration is a 3-to-400-base-pair in-frame internal tandem duplication (ITD) mutation within exon 14 that leads to the constitutive activation of the FLT3 receptor and ligand-independent autophosphorylation [4–9]. FLT3-ITD mutations are found in 20–30% of young adults with AML and are a poor prognostic indicator [9]. Targeted FLT3 inhibitors have been developed such as giltertinib, which was associated with significantly longer overall survival (OS) and higher complete remission (CR) rates compared to those with salvage chemotherapy in relapsed or refractory FLT3-ITD AML [10].

Whereas the importance of FLT3-ITD in AML from prognosis to treatment is well known, its significance in APL is less established. Present in 12–38% of acute promyelocytic leukemia [5,6], FLT3-ITD has been associated with high white blood cell counts (WBCs), the short PML-RARα breakpoint cluster region 3 (bcr3) isoform, and microgranular variant M3 (M3v) APL [6–9,11]. The Sanz grouping classifies APL patients as high risk (HR) (WBC > 10 × 10⁹/L), intermediate risk (IR) (WBC ≤ 10 × 10⁹/L, platelets ≤ 40 × 10⁹/L), and low risk (LR) (WBC ≤ 10 × 10⁹/L, platelets > 40 × 10⁹/L) for relapse-free survival (RFS) [12]. Sanz HR APL is associated with FLT3-ITD [8,12]. The clinical outcomes in FLT3-ITD APL remain controversial, and little is known about the impact of ITD insertion length and allelic burden in APL. In this retrospective study, we investigated the significance of FLT3-ITD, insertion length, and allelic burden in APL.

2. Materials and Methods

Patients 18 years or older treated at the University of Maryland Medical Center and Johns Hopkins Hospital from January 2000 to May 2020 were included. APL was defined by the cytogenetic and/or molecular confirmation of PML-RARα. FLT3-ITD mutations were targeted at the juxtamembrane region of the FLT3 gene (exons 14–15) via primers, amplified, and identified by fluorescent PCR. The FLT3-ITD allelic burden was estimated as a percent ratio of the area under the curve (AUC) of the variant peak divided by the AUC of the wild-type (WT) peak. The ITD insertion length was determined by subtracting the 328-base-pair PCR product of the FLT3 gene from the base pair size of the variant peak. De-identified patient ages, genders, ethnicities, body mass indices (BMIs), laboratory measurements, Sanz and FLT3-ITD statuses, ITD allelic burdens and insertion lengths, induction chemotherapy regimens, and outcomes were inputted into Microsoft Excel. These data were then transferred to Stata, Version 16.1 (StataCorp, College Station, TX, USA). All the statistical analyses were performed and graphs were made using Stata.

The Chi-square test of independence was run for two categorical variables (e.g., Sanz and FLT3-ITD status). Mean differences between two groups (e.g., FLT3-ITD status) were tested using independent-sample t-tests, and mean differences between three or more groups (e.g., Sanz) were tested using one-way analysis of variance (ANOVA), followed by
Scheffe tests. Pearson’s and Spearman’s correlations were used to study the associations of two continuous variables (e.g., allelic burden and WBC). For multiple regression analyses, logistic regression models were used for the associations of independent binary outcomes (e.g., \( \text{FLT3-ITD} \) status) with independent variables of interest (e.g., Sanz), before and after adjustment for other covariates (e.g., age, gender, and ethnicity). Similarly, linear regression models were used for the association of continuous outcomes (e.g., allelic burden) with independent exposures of interest (e.g., Sanz) before and after adjustment for other covariates. Overall survival (OS) was compared between all groups (\( \text{FLT3-ITD} \), \( \text{FLT3-WT} \), and Sanz HR/IR/LR) using the log-rank test for the equality of survivor functions and graphed using the Kaplan–Meier method. Cox regression models were used to compare survival between groups after adjustment for covariates of interest.

3. Results

We identified 138 patients (47 (34.1%) HR, 54 (39.1%) IR, and 37 (26.8%) LR) (Table 1). \( \text{FLT3-ITD} \) was detected in 59/138 (42.8%) of the APL patients. There were no significant differences in demographics between any of the groups (age, gender, ethnicity, and BMI). The \( \text{FLT3-ITD} \) patients were more likely to possess higher WBC (\( p < 0.001 \)), M3v (\( p < 0.001 \)), and bcr3 (\( p < 0.001 \)) characteristics, reported to be associated with worse outcomes in APL [6–9,11]. Higher hemoglobin (\( p = 0.03 \)) and aspartate aminotransferase (AST) (\( p = 0.001 \)), lower platelets (\( p = 0.004 \)) and fibrinogen (\( p = 0.003 \)), and a higher incidence of disseminated intravascular coagulation (DIC) (\( p = 0.005 \)) were also noted in our \( \text{FLT3-ITD} \) cohort, albeit without differences in all-cause bleeding (\( p = 0.39 \)), intracranial hemorrhage (\( p = 0.80 \)), or thrombosis (\( p = 0.33 \)). There were no significant differences noted in the incidences of differentiation syndrome when the cohort was stratified by the presence of \( \text{FLT3-ITD} \) (\( p = 0.29 \)), Sanz risk status (\( p = 0.20 \)), and both \( \text{FLT3-ITD} \) and Sanz risk statuses (\( p = 0.22 \)).

Sanz HR patients were more likely to have \( \text{FLT3-ITD} \) compared to IR and LR patients (Chi-square test of independence, \( X^2 (2, N = 138) = 37.8, p < 0.001 \)). However, the association between HR status and \( \text{FLT3-ITD} \) (odds ratio (OR): 13.4; 95% CI: 4.7–38.3; \( p < 0.001 \)) disappeared when adjusting for WBCs and platelets (OR: 0.7; 95% CI: 0.1–5.5; \( p = 0.80 \)). Sanz LR and IR patients with WBCs < 10 \( \times 10^9 \)/L comprised roughly 66% of the cohort, leaving the remaining 34% of HR patients divided evenly into two groups: WBCs of 10–25 \( \times 10^9 \)/L and >25 \( \times 10^9 \)/L. \( \text{FLT3-ITD} \) was strongly associated with WBCs \( \geq 25 \times 10^9 \)/L (OR: 54.4; 95% CI: 10.4–286.1; \( p < 0.001 \)), similar to the previously reported WBCs \( \geq 20 \times 10^9 \)/L [7]. WBCs of 10–25 \( \times 10^9 \)/L were, to a lesser degree, also associated with \( \text{FLT3-ITD} \) (OR: 8.65; 95% CI: 2.71–27.5; \( p < 0.001 \)). No differences in post-induction complete remission (CR) (\( p = 0.42 \)), post-consolidation CR (\( p = 0.61 \)), induction deaths (\( p = 0.80 \)), and OS (\( p = 0.33 \)) were noted between the Sanz risk groups.

HR APL was associated with an ITD insertion length decrease of 20 base pairs compared to LR APL (95% CI: \(-40.0 \) to \(-0.23 \); \( p = 0.05 \)); however, this finding was lost when adjusting for platelets (\(-13.5 \); 95% CI: \(-38.2–11.2 \); \( p = 0.3 \)). Whereas the ITD insertion length was associated with higher platelet counts (\( r_s = 0.39, p = 0.003 \)), no association was noted between the insertion length and WBC (\( r_s = -0.06, p = 0.65 \)). A longer ITD insertion length and ITD mutant/wildtype ratio greater than 0.5–0.66 have been associated with shorter RFS, and OS in APL [9–14]. No correlation between the insertion length and OS was noted in this study (\( p = 0.38 \)).
Table 1. Baseline demographic and clinical data of 138-patient acute promyelocytic leukemia (APL) cohort differentiated by presence of FLT3-ITD and Sanz risk status.

| Variable | ITD, HR (n = 59) | WT, HR (n = 79) | p-Value |
|----------|------------------|------------------|---------|
| **Age** (Years) | 47 | 50 | 0.27 |
| **Gender** | M (44%) | M (52%) | 0.39 |
| **Ethnicity** | M (49%) | M (48%) | 1.0 |
| **Bleeding** | Yes (49%) | Yes (48%) | 0.07 |
| **DIC** | Yes (53%) | Yes (56%) | 0.20 |
| **Bleeding** | Yes (30%) | Yes (41%) | 0.005 |
| **ICH** | Yes (14%) | Yes (11%) | 0.80 |

| Variable | ITD, HR (n = 59) | WT, HR (n = 79) | p-Value |
|----------|------------------|------------------|---------|
| **BMI** | 32.3 | 31.8 | 0.74 |
| **WBC** | 29.9 | 5.1 | <0.001 |
| **LDH** | 44.5 | 30.2 | 0.001 |
| **ALT** | 44.8 | 33.7 | 0.09 |
| **ALP** | 83.7 | 81.4 | 0.73 |
| **TBili** | 1.0 | 0.9 | 0.42 |
| **Fib** | 968.1 | 1030.5 | 0.92 |
| **Morph** | 168.2 | 225.0 | 0.003 |
| **BCR** | 3 (53%), 2 (22%) | 3 (52%), 2 (22%) | <0.001 |
| **DS** | Yes (46%), No (54%) | Yes (49%), No (51%) | 0.29 |
| **DIC** | Yes (73%), No (27%) | Yes (79%), No (61%) | 0.005 |
| **Bleeding** | Yes (49%), No (51%) | Yes (55%), No (59%) | 0.39 |
| **ICH** | Yes (14%), No (84%) | Yes (19%), No (81%) | 0.80 |

| Variable | ITD, HR (n = 59) | WT, HR (n = 79) | p-Value |
|----------|------------------|------------------|---------|
| **BMI** | 32.3 | 31.8 | 0.74 |
| **WBC** | 29.9 | 5.1 | <0.001 |
| **LDH** | 44.5 | 30.2 | 0.001 |
| **ALT** | 44.8 | 33.7 | 0.09 |
| **ALP** | 83.7 | 81.4 | 0.73 |
| **TBili** | 1.0 | 0.9 | 0.42 |
| **Fib** | 968.1 | 1030.5 | 0.92 |
| **Morph** | 168.2 | 225.0 | 0.003 |
| **BCR** | 3 (53%), 2 (22%) | 3 (52%), 2 (22%) | <0.001 |
| **DS** | Yes (46%), No (54%) | Yes (49%), No (51%) | 0.29 |
| **DIC** | Yes (73%), No (27%) | Yes (79%), No (61%) | 0.005 |
| **Bleeding** | Yes (49%), No (51%) | Yes (55%), No (59%) | 0.39 |
| **ICH** | Yes (14%), No (84%) | Yes (19%), No (81%) | 0.80 |
### Table 1. Cont.

| Variable | ITD (n = 59) | WT (n = 79) | p-Value |
|----------|--------------|-------------|---------|
| Clot     | Yes (19%), No (81%) | Yes (11%), No (89%) | 0.33 |
| CR\textsubscript{induc} | Yes (89%), No (4%), Death (7%) | Yes (93%), No (3%), Death (4%) | 0.70 |
| CR\textsubscript{cons} | Yes (81%), No (4%), Death (15%) | Yes (97%), No (0), Death (3%) | 0.02 |
| 5-Year OS | 79.7%, 94.4% | 0.02 | 87.1% |

| Variable | ITD (n = 47) | IR (n = 54) | LR (n = 37) | p-Value |
|----------|--------------|-------------|-------------|---------|
| ITD, HR  | Yes (22%), No (78%) | Yes (20%), No (80%) | Yes (14%), No (86%) | 0.12 |
| WT, HR   | Yes (100%), No (0%), Death (15%) | Yes (85%), No (0%), Death (5%) | Yes (95%), No (0%), Death (5%) | 0.42 |
| ITD, IR  | Yes (88%), No (6%), Death (6%) | Yes (91%), No (6%), Death (3%) | Yes (89%), No (6%), Death (3%) | 0.42 |
| WT, IR   | Yes (100%), No (0%), Death (15%) | Yes (99%), No (0%), Death (15%) | Yes (99%), No (0), Death (15%) | 0.61 |
| ITD, LR  | Yes (83%), No (7%), Death (10%) | Yes (93%), No (0%), Death (0%) | Yes (83%), No (7%), Death (10%) | 0.61 |
| WT, LR   | Yes (100%), No (0%), Death (14%) | Yes (99%), No (0%), Death (14%) | Yes (100%), No (0%), Death (14%) | 0.61 |

| Variable | ITD (n = 37) | IR (n = 54) | LR (n = 37) | p-Value |
|----------|--------------|-------------|-------------|---------|
| ITD, HR  | Yes (22%), No (78%) | Yes (20%), No (80%) | Yes (14%), No (86%) | 0.12 |
| WT, HR   | Yes (100%), No (0%), Death (15%) | Yes (85%), No (0%), Death (5%) | Yes (95%), No (0%), Death (5%) | 0.42 |
| ITD, IR  | Yes (88%), No (6%), Death (6%) | Yes (91%), No (6%), Death (3%) | Yes (99%), No (0), Death (3%) | 0.42 |
| WT, IR   | Yes (100%), No (0), Death (15%) | Yes (99%), No (0), Death (15%) | Yes (100%), No (0), Death (15%) | 0.61 |
| ITD, LR  | Yes (83%), No (7%), Death (10%) | Yes (93%), No (0%), Death (0%) | Yes (93%), No (0), Death (0%) | 0.61 |
| WT, LR   | Yes (100%), No (0), Death (14%) | Yes (99%), No (0), Death (14%) | Yes (99%), No (0), Death (14%) | 0.61 |

| Variable | ITD (n = 37) | IR (n = 54) | LR (n = 37) | p-Value |
|----------|--------------|-------------|-------------|---------|
| ITD, HR  | Yes (22%), No (78%) | Yes (20%), No (80%) | Yes (14%), No (86%) | 0.12 |
| WT, HR   | Yes (100%), No (0), Death (15%) | Yes (85%), No (0), Death (5%) | Yes (95%), No (0), Death (5%) | 0.42 |
| ITD, IR  | Yes (88%), No (6), Death (6%) | Yes (91%), No (6), Death (3%) | Yes (99%), No (0), Death (3%) | 0.42 |
| WT, IR   | Yes (100), No (0), Death (15%) | Yes (99), No (0), Death (15%) | Yes (100), No (0), Death (15%) | 0.61 |
| ITD, LR  | Yes (83), No (7), Death (10) | Yes (93), No (0), Death (0) | Yes (93), No (0), Death (0) | 0.61 |
| WT, LR   | Yes (100), No (0), Death (14) | Yes (99), No (0), Death (14) | Yes (100), No (0), Death (14) | 0.61 |

ITD = FLT3 internal tandem duplication mutation, WT = FLT3 wild-type, HR = Sanz high risk, IR = Sanz intermediate risk, LR = Sanz low risk, Age = mean age in years, Gender = male (M) or female (F), Ethnicity = white = W, black = B, other = O, BMI = admission body mass index, WBC = admission white blood cell count in $\times 10^9/L$, Hgb = admission hemoglobin in g/dL, Plt = admission platelet count in $\times 10^9/L$, Creat = admission creatinine in mg/dL, AST = admission aspartate aminotransferase in units/L, ALT = admission alanine aminotransferase in units/L, ALP = admission alkaline phosphatase in units/L, TBili = admission total bilirubin in mg/dL, LDH = admission lactate dehydrogenase in units/L, Fib = admission fibrinogen in mg/dL, Morph = classic or variant, BCR = breakpoint cluster region, DS = differentiation syndrome during induction, DIC = disseminated intravascular coagulation during induction, Bleeding = any bleeding event during induction, ICH = intracranial hemorrhage during induction, Clot = any thrombotic event during induction, CR\textsubscript{induc.} = complete remission after induction, CR\textsubscript{cons.} = complete remission after consolidation, OS = overall survival (survivor function %) in 5 years.
The Sanz risk status significantly correlated with allelic burden according to one-way ANOVA ($F(2, 30) = 12.1, p < 0.0001$). According to post hoc Scheffe tests, the allelic burden differed between HR/IR ($p = 0.007$) and HR/LR ($p = 0.001$), but not in IR/LR ($p = 0.32$). These findings persisted when adjusting for center, age, gender, ethnicity, WBC, and platelets via linear regression. HR APL was associated with a 38.5% increased allelic burden compared with LR APL (95% CI: 19.8–57.2; $p < 0.001$). According to Spearman’s Rho, the FLT3-ITD allelic burden was associated with higher WBCs ($r_s = 0.49$, $p = 0.001$). For every 1% increase in allelic burden, the WBC increased by $0.6 \times 10^9$/L. The relationship between the FLT3-ITD allelic burden and WBC can be visualized in Figure 1. Of note, the majority of the reported allelic ratios (33/59 patients) were from those seen after 2012. There was no significant relationship between the FLT3-ITD allelic burden and OS ($p = 0.97$).

**Figure 1.** Distribution of FLT3-ITD allelic burden (%) and associated WBC count ($\times 10^9$/L) at diagnosis per patient in FLT3-ITD APL.

While studies have reported a higher incidence of induction death and inferior CR rates, OS, and RFS in FLT3-ITD APL [6,9–11,15,16], we report no significant differences in the CR duration after induction ($p = 0.70$) or death during induction ($p = 0.13$). We found shorter post-consolidation CR durations ($p = 0.02$) and OS ($p = 0.02$) in FLT3-ITD APL (Figure 2). The 5-year OS for patients with FLT3-ITD was 79.7% compared to 94.4% for FLT3 wild-type patients. FLT3-ITD was associated with a higher mortality risk, with a hazard ratio of 3.25 (95% CI: 1.14–9.25; $p = 0.027$). Other studies reported no association between FLT3-ITD and CR/RFS/OS/early death in APL [7,8,11,17–20]. The variability of the outcomes may be attributed to the small cohort size, differences in inclusion criteria, varying treatment protocols, and adherence to follow-up.

Pre-arsenic trioxide (ATO) studies showed reduced OS and increased relapse rates with FLT3-ITD [9,15,21] in APL, whereas post-ATO studies demonstrated no prognostic significance of FLT3-ITD [17–20], suggesting the effect of FLT3-ITD may be mitigated with ATO. Our cohort consisted of 26 FLT3-ITD patients treated with ATO, compared to 32 without ATO, and 41 FLT3-WT patients treated with ATO, compared to 37 without (two patients with unclear induction regimens). When mutually adjusting for FLT3-ITD...
status, the ATO-containing regimens were not associated with improved OS, with a hazard ratio (95% CI) of 1.13 (0.41–3.16). When stratified by the ATO versus non-ATO regimens, FLT3-ITD was associated with increased mortality in both groups. The hazard ratio (95% CI) was 4.78 (1.01–22.6) for the non-ATO regimens and 3.17 (0.58–17.3) for the ATO-containing regimens (p-value for interaction = 0.46), suggesting no evidence for the impact of ATO-containing regimens on the prognostic significance of FLT3-ITD in this study. However, the study sample size was modest, making the power relatively low for detecting effect modification. The number of patients treated with gemtuzumab ozogamicin was too small (n = 4) to perform statistical analysis on outcomes.

4. Discussion

APL therapy differs in Sanz HR versus IR/LR groups, and risk-stratifying APL is important for optimizing outcomes. Genes impacting differentiation are downregulated and genes involved in cellular adhesion, invasiveness, and metastasis are upregulated in FLT3-ITD APL [16]. Given the associations of FLT3-ITD with poor prognostic features and outcomes, FLT3-ITD APL has been proposed to be a distinct subtype of APL [7,9]. The high WBCs, M3v, and bcr3 reported in this study have been well-established in FLT3-ITD APL, whereas the shorter post-consolidation CR duration and OS found here are less well-established.

We found a strong correlation between FLT3-ITD and leukocytosis in APL, suggesting a WBC cutoff of $\geq 25 \times 10^9$/L as an indicator for considering testing for FLT3-ITD in APL. The well-known association between leukocytosis and FLT3-ITD in APL is further expanded upon in this study by the novel revelation of the WBC’s relationship with the ITD allelic burden, with every 1% increase in allelic burden equating to a $0.6 \times 10^9$/L increase in WBC. Interestingly, the allelic burden was not found to be associated with OS.

Figure 2. Kaplan–Meier survival curves comparing OS between (a) all patients: FLT3-ITD vs. FLT3-WT, (b) FLT3-ITD/Sanz LR vs. FLT3-WT/Sanz LR, (c) FLT3-ITD/Sanz IR vs. FLT3-WT/Sanz IR, and (d) FLT3-ITD/Sanz HR vs. FLT3-WT/Sanz HR.
5. Conclusions

Taken together, these results support the importance of additional study of the significance of the FLT3-ITD mutation and ITD allelic burden in APL. More data are required to determine the utility of incorporating FLT3-ITD into risk-adapted treatment algorithms and molecular monitoring. The absence of routine testing for FLT3-ITD in APL, the lack of international standardized FLT3-ITD assays, and the rarity of the disease pose limitations for studies of FLT3-ITD APL [6,13].

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Informed Consent Statement: Patient consent was waived due to the retrospective nature of the study and inclusion of only de-identified patient information in the study.

Data Availability Statement: The data presented in this study are available on request from the corresponding author. The data are not publicly available due to the protection of the de-identified patient information.

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Abbreviations

ALP alkaline phosphatase
ALT alanine aminotransferase
AML acute myeloid leukemia
ANOVA analysis of variance
APL acute promyelocytic leukemia
AST aspartate aminotransferase
ATO arsenic trioxide
ATRA all-trans retinoic acid
AUC area under the curve
bcr3 breakpoint cluster region 3
BMI body mass index
CR complete remission
DIC disseminated intravascular coagulation
DS differentiation syndrome
FAB French–American–British
Fib fibrinogen
Hgb hemoglobin
HR Sanz high risk
ICH intracranial hemorrhage
FLT3-ITD fms-like tyrosine kinase 3 internal tandem duplication
IR Sanz intermediate risk
LDH lactate dehydrogenase
LR Sanz low risk
M3v microgranular variant M3
Morph morphology
OS overall survival
PCR polymerase chain reaction
Plt platelet
PML-RARA promyelocytic leukemia retinoic receptor alpha
RFS relapse free survival
TBili total bilirubin
WBC white blood cell
WT wild-type

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