Integrated genomic sequencing in myeloid blast crisis chronic myeloid leukemia (MBC-CML), identified potentially important findings in the context of leukemogenesis model

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Chronic myeloid leukemia (CML) is a model of leukemogenesis in which the exact molecular mechanisms underlying blast crisis still remained unexplored. The current study identified multiple common and rare important findings in myeloid blast crisis CML (MBC-CML) using integrated genomic sequencing, covering all classes of genes implicated in the leukemogenesis model. Integrated genomic sequencing via Whole Exome Sequencing (WES), Chromosome-seq and RNA-sequencing were conducted on the peripheral blood samples of three CML patients in the myeloid blast crisis. An in-house filtering pipeline was applied to assess important variants in cancer-related genes. Standard variant interpretation guidelines were used for the interpretation of potentially important findings (PIFs) and potentially actionable findings (PAFs). Single nucleotide variation (SNV) and small InDel analysis by WES detected sixteen PIFs affecting all five known classes of leukemogenic genes in myeloid malignancies including signaling pathway components (ABL1, PIK3CB, PTPN11), transcription factors (GATA2, PHF6, IKZF1, WT1), epigenetic regulators (ASXL1), tumor suppressor and DNA repair genes (BRCA2, ATM, CHEK2) and components of spliceosome (PRPF8). These variants affect genes involved in leukemia stem cell proliferation, self-renewal, and differentiation. Both patients No.1 and No.2 had actionable known missense variants on ABL1 (p.Y272H, p.F359V) and frameshift variants on ASXL1 (p.A627Gfs*8, p.G646Wfs*12). The GATA2-L359S in patient No.1, PTPN11-G503V and IKZF1-R208Q variants in the patient No.3 were also PAFs. RNA-sequencing was used to confirm all of the identified variants. In the patient No. 3, chromosome sequencing revealed multiple pathogenic deletions in the short and long arms of chromosome 7, affecting at least three critical leukemogenic genes (IKZF1, EZH2, and CUX1). The large deletion discovered on the short arm of chromosome 17 in patient No. 2 resulted in the deletion of TP53 gene as well. Integrated genomic sequencing combined with RNA-sequencing can successfully discover and confirm a wide range of...
variants, from SNVs to CNVs. This strategy may be an effective method for identifying actionable findings and understanding the pathophysiological mechanisms underlying MBC-CML, as well as providing further insights into the genetic basis of MBC-CML and its management in the future.

**Abbreviations**

AP  Accelerated phase  
DP  Depth of coverage  
ACMG  American college of medical genetics and genomics  
ASCO  American society of clinical oncology  
AMP  Association for molecular pathology  
B  Benign  
BMA  Bone marrow aspiration  
CML  Chronic myeloid leukemia  
CGI  Cancer genome interpreter  
CAP  College of American pathologists  
CNV  Copy number variation  
Dx  Diagnosis  
Dv  Driver  
ER  Epigenetic regulators  
GATK  Genome analysis tool kit  
kDv  Known driver  
LPC  Leukemia progenitor cell  
LSC  Leukemia stem cell  
LB  Likely benign  
LP  Likely pathogenic  
MDS  Myelodysplastic syndrome  
MBC  Myeloid blast crisis  
MPS  Massively parallel sequencing  
MPNs  Myeloproliferative neoplasms  
NGS  Next generation sequencing  
Npa  Not protein affecting  
Ps  Passenger  
P  Pathogenic  
PB  Peripheral blood  
PIF  Potentially important finding  
PAF  Potentially actionable finding  
RT-PCR  Reverse transcriptase-polymerase chain reaction  
SNV  Single nucleotide variant  
TF  Transcription factor  
TSG  Tumor suppressor gene  
VUS  Variant of uncertain significance  
VUS-B  Variant of uncertain significance leaning benign  
VUS-P  Variant of uncertain significance leaning pathogenic  
VAF  Variant allele fraction  
WES  Whole exome sequencing

Chronic Myeloid Leukemia (CML) is a unique model for the evolution of cancer and is classified as a triphasic myeloproliferative neoplasm based on clinical and pathological characteristics. Premalignant leukemia stem cells (LSCs) are, in fact, generated in the bone marrow nich by unknown mutagenesis processes. In this step, the disease may be undetectable for a decade or more. Further oncogenic processes cause LSCs to develop into leukemia progenitor cells (LPCs). Leukemogenic alterations mainly affect five classes of regulatory proteins: signaling pathway components, transcription factors (TFs), epigenetic regulators (ERs), tumor suppressor genes (TSGs), and components of the spliceosome.

Without therapeutic interventions or in the case of drug resistance, CML will progress to accelerated phase (AP) and then into an acute leukemia phase that is called blastic phase or blast crisis. Even in the advent of Tyrosin Kinase Inhibitor therapy, blastic phase survival is less than 12 months. In general, blast crisis in chronic myeloid leukemia is still a fatal disease.

Over the past decade using Massively Parallel Sequencing (MPS) or Next Generation Sequencing (NGS) the multi-dimensional genomic view with higher resolution of molecular profiling has become possible. It was shown that the combination of exome, genome and RNA Sequencing data, called integrative sequencing, paves the way to better understand the mechanisms and pathogenesis of the cancer and can potentially improve the clinical management and finding new therapeutic targets specially in the advanced phase patients.

In the present study, we conducted integrated genomic sequencing through an in-house filtering algorithm to uncover the leukemogenic important/actionable findings and investigate the myeloid leukemogenesis model in three Iranian myeloid blast crisis CML patients.
Methods

Subjects. Three patients (from 3 unrelated families) with the clinical diagnosis of blastic phase CML were recruited from the Shariati Hospital in Tehran, Iran. Inclusion criteria were based on clinical presentations and hematological findings in the peripheral blood and bone marrow, according to the WHO 2016-guideline.

The study was approved by the Ethical Committee of the Iran University of Medical Sciences (Code: IR.IUMS.REC 1395.95-04-87-30235). All subjects and/or their legal guardian(s) gave their informed consent to participate. All methods, including obtaining informed consent, were carried out in compliance with the above-mentioned ethical standards and guidelines.

Whole Exome Sequencing (WES) along with Chromosome-seq and RNA Sequencing were performed on extracted DNA and RNA from peripheral blood samples in the blast crisis phase.

Whole exome sequencing. Peripheral blood samples of patients were collected in EDTA containing tubes and used to extract genomic DNA by Blood SV-mini kit (GeneAll Biotechnology Co., LTD, South Korea) according to the manufacturer instruction. The extracted genomic DNA samples were subjected to WES using Agilent SureSelect V6-post Capture/Target Enrichment Kit. For the third patient TWIST Human Core Exome Kit (Twist Bioscience, USA) was used according to the manufacturer’s instruction. The enriched libraries were sequenced on Illumina platforms (Illumina Inc., CA, USA). Table S1 shows a summary report of data analysis metrics from the patients’ WES.

WES data analysis (variant filtering). Paired-end sequence reads were mapped to the refseq (NCBI) human reference genome (GRCh38 assembly) by BWA-MEM. Picard software was used to mark duplicate reads. Genome Analysis Tool Kit (GATK) software Mutect2 package was used to call single nucleotide variants (SNVs) and short insertions or deletions (Indels). The intronic variants were removed, however the 100 bp flanking each exon was included in VCF files using interval.bed file.

Several tools were used to annotate and interpret variants including CIGI, Varsome, InterVar, CancerVar, and Franklin. The annotated VCF files were filtered to find Tier1 or Tier2 variants according to AMP classification and pathogenic (P), likely pathogenic (LP), or Leaning pathogenic variants based on ACMG classification.

All Tier 1 or Tier 2 variants in AMP classification and the Tier 3 variants that were pathogenic (P), likely pathogenic (LP), or Leaning pathogenic VUS in ACMG classification were selected as potentially important findings (PIFs). PIFs known to be therapeutic, prognostic or diagnostic biomarkers were annotated as potentially actionable findings (PAFs).

RNA sequencing. RNA from whole blood in PaxGene tube was extracted and processed by the TruSeq Stranded Total RNA with Ribozero Gold Kit (Illumina Inc., CA, USA) using 1 µg of DNAse treated RNA. Then paired-end sequencing was performed on Illumina platforms.

CLC genomics workbench 20 was used for RNA sequencing and fusion gene analyses. RNA sequencing results were used for determining the outlier expression analysis of genes harboring important/actionable variants, analyzing the fusion transcripts as well as confirmation of variants found in WES.

Chromosome-sequencing. Vista™ Chromosome Sequencing-100 K an NGS based Whole Genome Sequencing was performed in BGI Clinical Laboratories to evaluate copy number variations (≥ 100 Kb).

Cross validation of the identified variants. The identified SNVs and small Indels in WES were visually validated in the mapping read-tracks after importing WES BAM files in the CLC genomics workbench 20. Besides, the validations of identified WES variants were performed using RNASeq BAM files by confirming both variant and gene expression.

The detected CNVs in chromosome-seq data were validated by CNV analysis from WES data using CLC genomics workbench 20.

Ethics approval and consent to participate. The study was approved by the Ethical Committee of the Iran University of Medical Sciences (Code: IR.IUMS.REC 1395.95-04-87-30235). Informed consent was obtained from all subjects and/or their legal guardian(s) for participation.

Results

Clinical findings. The patient (I), was a 66-year-old female presented by fever, hepatosplenomegaly, leukocytosis, anemia, thrombocytopenia. The blast level in the peripheral blood was 25%. Bone marrow aspiration and immunophenotyping confirmed the myeloid blastic crisis. The patient had a history of thyroid cancer that had been treated many years prior to the development of CML. She was taken to the hospital at the age of 63 with a primary diagnosis of chronic phase CML in terms of leukocytosis with myeloid precursors in the peripheral blood and a positive Philadelphia chromosome in karyotype.

The patient (II), a 55-year old female presented by leukocytosis, anemia, thrombocytopenia. The diagnosis of chronic phase CML was established 15 years before blastic phase at the age of 40 years. There was 36% blast in the peripheral blood. Bone marrow aspiration and immunophenotyping confirmed the myeloid blastic crisis. At blast crisis, the peripheral blood karyotype revealed a clone that was positive for both the philadelphia chromosome and isochromosome 17q, as well as a clone with paracentric inversion in 15q.
Table 1. Clinical and laboratory findings for the CML patients at diagnosis/chronic phase. Dx diagnosis, NA not available, RT-PCR reverse transcriptase PCR, TKI tyrosine kinase inhibitor.

| Patient no | Patient 1 | Patient 2 | Patient 3 |
|------------|-----------|-----------|-----------|
| Gender     | Female    | Female    | Male      |
| Age at diagnosis (years) | 63        | 40        | 43        |
| WBC per µL | 18,500    | 165,200   | 40,000    |
| Diff (CBC) | Myeloblast: 2%, Granulocyte: 79%, Lymphocyte: 10%; Eosinophil: 3%; MetaMyelocyte: 3%; Band: 3% | Myeloblast: 3%, Granulocyte: 58%, Lymphocyte: 9%; Eosinophil: 1%; MetaMyelocyte: 7%; Band: 11%; Myelocyte: 8%; ProMyelocyte: 3% | Myeloblast: 6% Granulocyte: 38%; Lymph:17%; Eosinophil: 4%; MetaMyelocyte: 15%; Band: 3%; Basophil: 1%; Myelocyte: 14%; Monocyte: 2% |
| Platelet per µL | 1,918,000 | 703,000   | 711,000   |
| Hemoglobin (mg/dL) | 9.6       | 12.5      | 11.6      |
| Spleen Size | Palpable (2 cm) | Palpable (2 cm) | NA        |
| Sokal relative risk score | 3.79 (high) | 0.93 (intermediate) | 1.18 (intermediate) |
| Bone Marrow Pathology | High and compact cellularity; Marked myeloid hyperplasia; Giant and lobulated Megakaryocytes with marked hyperplasia; Plasmocytes:3%; Lymphoid: 4%; Normal Iron storage | The marrow is 100% cellular. The hematopoietic populations are replaced by atypical granulocytic series, composed of prominent segmented neutrophils mixed with immature forms including myelocytes and promyelocytes. Myeloblasts in sheets are about 60% of populations. The marrow framework is fibrotic in occasional local areas | NA |
| BCR-ABL1 RT-PCR | Positive | Positive | Positive |
| ABL1 kinase domain Mutation analysis (by PCR-sanger sequencing) | p.F359V mutation was detected (35 months after diagnosis) | p.G250E mutation was detected (74 months after diagnosis) | No mutation was detected (22 months after diagnosis at MBC phase) |
| Probable Mechanism of TKI Resistance | ABL1 kinase domain p.F359V Mutation causes resistance to imatinib and nilotinib | ABL1 kinase domain p.G250E Mutation causes resistance to imatinib | Additional cytogenetic abnormalities (according to the chromosome-seq data in Table 2) |

Patient (III) was a 45-year old male presented by leukocytosis, anemia and thrombocytopenia. The diagnosis of chronic phase CML was established two years before blastic phase at the age of 43 years. There was 40% blast in the peripheral blood. Bone marrow aspiration and immunophenotyping confirmed the myeloid blastic crisis. BCR-ABL1 fusion transcript (b13-a2) was also found in the RNA-seq fusion gene study of all patients during the blast crisis phase.

Table 1 summarizes the patients' detailed clinical and paraclinical findings at chronic phase CML diagnosis. The clinical and paraclinical findings at myeloid blast crisis are presented in the Table 2.

**Genetic findings.** WES analysis identified sixteen PIFs (Figs. 1, 2 and 3) in which ABL1-Y272H, ASXL1-A627Gfs*8, GATA2-L359S, ABL1-F359V, ASXL1-G646Wfs*12, PTPN11-G503V and IKZF1-R208Q variants were classified as PAFs. The ABL1-I418T, PIK3CB-R231H, CHEK2-c.-4C>T, BRCA2-P329L, ABL1-E459G, PRRF8-G1796R, PHF6-R274Q, WT1-R435X and ATM-C117V variants were potentially important cancer variants which were not actionable (Table 3).

**Genetic findings in the patient no.1.** The patient was a BCR-ABL1 positive CML case in the myeloid blast crisis phase. She had abnormal karyotype with Philadelphia chromosome. In chromosome-seq there were no specific copy number alterations.

In WES she had six PIFs which contributed in all classes of leukemogenic genes (Fig. 1) except splicing components (class V). The p.Y272H variant on ABL1 (class I), p.A627Gfs*8 on ASXL1 (class III) and p.L359S on GATA2 (class II) were PAFs.

**Genetic findings in the patient no.2.** The patient was a BCR-ABL1 positive CML case in the myeloid blast crisis phase. She had abnormal complex karyotype with Philadelphia chromosome and isochromosome 15q (in another stemline (46,XX, t(9;22)(q34;q11.2),i(17)(q10) [17]/46,sl,inv(15)(q21.2;q25) [33])). In chromosome-seq (Fig. S1) there were deletions in 15q and 17p, and duplication in 17q (46,XX,del(15q26.3); del(17p11.2p13.3); dup(17p11.2q25.3)). The TP53 gene is a class II leukemogenic gene (transcription factor) located in 17p13.1 which is deleted in this patient.

In WES she had six PIFs that contribute in all classes of leukemogenic genes (Fig. 2). Considering the TP53 involvement by 17p deletion, this patient has seven PIFs. The p.F359V variant in ABL1 (class I), p.G646Wfs*12 on ASXL1 (class III) and TP53 (class IV) deletion were PAFs.

**Genetic findings in the patient no.3.** The patient was a BCR-ABL1 positive myeloid blast crisis CML. In chromosome-seq (Fig. S1) he had several deletions in 7p and 7q (46,XY,del(7p14.3p21.1); del(7p11.2p14.3); del(7q11.23q21.11)). The IKZF1 (7p12.2), EZH2 (7q35-q36) and CUX1 (7q22) are important leukemogenic
| Patient no | Patient 1 | Patient 2 | Patient 3 |
|------------|-----------|-----------|-----------|
| Age of MBC (years) | 66 | 55 | 45 |
| Age of death | 67 (16 days after MBC Dx) | 55 (6 month after MBC Dx) | 45 (3 month after MBC Dx) |
| WBC per µL | 55,350 | 93,520 | 16,310 |
| Diff (CBC) | Blast: 25%, Granulocyte: 55%, Lymphocyte: 20% | Blast: 36%, Granulocyte: 59%, Lymphocyte: 5% | Blast: 40% Granulocyte: 50%, Lymph:10% |
| Platelet per µL | 53,000 | 41,000 | 11,000 |
| Hemoglobin (mg/dL) | 7.4 | 7.8 | 6.4 |
| Spleen Size | 144*63 mm | 30 mm below the left costal margin | 192*70 mm; huge splenomegaly |
| Karyotype | 46,XX, der(9)(9;22)(q34;q11.2)(9;22) (9;22)(p13;q12);p13)(9;22)(q34;q11.2), der(22)(q23)(q12);p13)(9;22)(q34;q11.2) | 46,XX, t(9;22)(q34;q11.2)(17)(q10)(17) | NA |
| Chromosome seq [GRCh37/hg19] | 46,XX, del(15q26.3);seq(99,294,139–100,356,602) × 1; del(15q11.2p13.3) seq(1–18,903,731) × 1; dup(17p11.2q25.3), seq(18,921,492–81,170,888) × 3 | 46,XX,del(7p14.3p11.1).seq(20,665,612–29,695,336) × 1; del(7p11.2p14.3). seq(33,201,686–57,488,614) × 1; del(7q11.23q21.11).seq(76,745,520–85,788,290) × 1 |
| Quantitative BCR-ABL (PB) | 24.1% | 62.4% | 58% |
| Immunophenotyping | Positive for Myeloid markers: (MPO, CD33), Negative for Lymphoid markers (CD3,CD19,CD20) and CD34 Positive | Positive for Myeloid markers: (MPO, CD33, CD117), Negative for Lymphoid markers (CD3,CD19,CD20) and CD34 Positive | Positive for Myeloid markers: (MPO), Negative for Lymphoid markers (CD10,CD19(1),CD20(1)) and CD34 Positive |
| Specific Therapeutic Management at MBC | Cytarabine; Arsenic (discontinued after two days due to cardiotoxicity); Dasatinib | Hydroxyurea; Cytarabine; Nilotinib; Dasatinib | Hydroxyurea; Imatinib; Cytarabine; Chedarabin; Danarubicin; Mitoxantrone; Nilotinib |
| HSCT | [HSCT has been suggested but not performed] | Incomplete [HLA typing was conducted but appropriate donor was not identified] | Incomplete [HLA typing was conducted but HSCT was not performed due to the rapid disease progression and death] |

Table 2. Clinical and laboratory findings for the CML patients in MBC phase. Dx diagnosis, NA not available, MBC myeloid blast crisis, HSCT hematopoietic stem cell transplantation.

genes which are located in the deletion regions. EZH2 gene is an epigenetic regulator gene (H3K27 methyltransferase; class III) and CUX1 is a transcription factor (class II) regulating TP53 and ATM.

He also had four PIFs in WES (Fig. 3). In terms of the pathogenic deletions in the short and long arms of chromosome 7 affecting EZH2 and CUX1, this patient has six PIFs which involve four classes of leukemogenic genes. **PTPN11**p.G507V (class I) and **IKZF1**p.R65Q variants (class II) were PAFs. The **WT1**p.R435X (class II) and **ATM**p.C117Y (class IV) variants were not PAFs. **IKZF1** variant (p.R65Q) in combination with del(7p11.2p14.3) may affect **IKZF1** gene product as an important transcription factor in myeloid malignancies.

**Discussion**

The ability to analyze and model the leukemogenesis process may aid in the effective management and treatment of hematologic malignancies. According to the proposed “Slot machine” model of leukemogenesis by Murati et al., progression from chronic phase to secondary acute myeloid leukemia (blast crisis) can be modeled by the combination of at least four steps that one classes of leukemogenic genes is affected in each step.

Using integrated genomic sequencing to assess the genes affected by SNVs, Indels and CNVs, in the pre-indices, the ability to analyze and model the leukemogenesis process may aid in the effective management and treatment of hematologic malignancies. According to the proposed “Slot machine” model of leukemogenesis by Murati et al., progression from chronic phase to secondary acute myeloid leukemia (blast crisis) can be modeled by the combination of at least four steps that one classes of leukemogenic genes is affected in each step.

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Figure 1. The paired-end mapping read-track displaying reads and detected variants in the first patient. The potentially important findings from WES data analysis is presented in the CLC genomics workbench software. (A) hg38 sequence track with WES mapped reads track; (B) hg38 sequence track with RNA-Seq mapped reads track; (C) Gene expression track; (D) gene, mRNA and CDS tracks; (E) the variant genomic coordinate on hg38 and the variant name. P1: patient No.1.
Figure 2. The paired-end mapping read-track displaying reads and detected variants in the second patient. The potentially important findings from WES data analysis is presented in the CLC genomics workbench software. (A) hg38 sequence track with WES mapped reads track; (B) hg38 sequence track with RNA-Seq mapped reads track; (C) Gene expression track; (D) gene, mRNA and CDS tracks; (E) the variant genomic coordinate on hg38 and the variant name. P2: patient No.2.
The third patient's PTPN11-G507V variant is located in the Phosphatase (PH) domain. Although PTPN11 mutations have been documented in blastic phase CML, the p.G507V alteration is a novel variant in CML and has only been identified in other myeloid malignancies such as JMML, AML and MDS. (COSM14271).

There were several important variants in different transcription factor genes including GATA2, PHF6, WT1 and IKZF1 in MBC-CML patients. In addition, TP53, IKZF1 and CUX1 genes were affected by chromosomal deletions/CNVs. GATA2-L359S variant in the first patient is located on the zinc finger GATA domain. It has been indicated that GATA2 germline mutations strongly predispose patients to leukemia22. GATA2 mutations were described in MBC-CML; however p.L359S variant has not been previously reported in blastic CML. p.L359S variant was suggested as a predisposing germline likely pathogenic variant in MDS/AML 23.

PHF6-R274Q variant in patient No.2 is located in the PHD-like zinc binding domain. Mutations in PHF6 also have been reported in the blastic phase CML while p.R274Q variant is a novel mutation in CML which only reported in Early T-cell precursor (ETP) acute lymphoblastic leukemia (COSM306061).

Biallelic involvement of IKZF1 was observed in the third patient. The detected missense variant was located in the C2H2 zinc finger domain of the IKZF1 (p.R65Q). Because of the 7p deletion, this variant showed a high variant allele fraction (VAF = 100%) in WES analysis. IKZF1 mutations had been found in both chronic and blastic phase CML24. The patient with IKZF1 mutation also had a mutation in PTPN11. It has been shown that in AML patients, mutations in IKZF1 and PTPN11 are associated with aggressive clinical course and primary resistant to chemotherapy25. CUX1 gene involvement was detected in the third patient in the cytogenetic studies in terms of 7q deletion. CUX1 is a transcription factor regulating TP53 and ATM26.

17p deletion in TP53 gene was established by cytogenetic analysis in the second patient. TP53 is a transcription factor located in 17p13.1. TP53 regulate cell cycle, DNA repair and apoptosis. It was observed that 45% of blast phase MPN (MPN-BP) patients have a TP53-related defect such as TP53 gene mutations or haploinsufficiency27. Involvement of TP53 has been described in both chronic and blastic phases of CML28,29.

Additional chromosomal abnormalities (ACAs) have been discovered in the second and the third patients. Patients with high-risk ACA, such as i(17q) and -7/7q-, are known to have a poor response to TKIs and a greater risk of disease progression. -7/7q- are less common in BC than i(17q), although they have a greater unfavorable influence on prognosis30. The patient number 3 with multiple deletions in the chromosome 7 has shorter time to MBC.

WT1-R435X is another significant nonsense variant discovered in the third patient. WT1 is a tumor suppressor gene which has known function as a transcriptional regulator31. This mutation causes loss of function in WT1 by the activation of NMD (Nonsense mediated decay) mechanism. Mutations in WT1 were reported in the blastic phase CML32, Wilms tumor and desmoplastic small round cell tumor (COSM21401). Pathogenic variants in this gene are correlated with poor prognosis in the patients with myelodysplastic syndromes33. The p.R435X germline mutation in WT1 has been classified as pathogenic which is associated to Wilms tumor in the clinvar (RCV00003671).

Figure 3. The paired-end mapping read-track displaying reads and detected variants in the third patient. The potentially important findings from WES data analysis is presented in the CLC genomics workbench software. (A) hg38 sequence track with WES mapped reads track; (B) hg38 sequence track with RNA-Seq mapped reads track; (C) Gene expression track; (D) gene, mRNA and CDS tracks; (E) the variant genomic coordinate on hg38 and the variant name. P3: patient No.3.
Two frameshift single nucleotide deletions were identified in ASXL1 (A627Gfs*8, G646Wfs*12). ASXL1 mutations are more frequent in secondary AML and may contribute to the development of CML to AML. Moreover, ASXL1 mutation might be associated with an aggressive phenotype in myeloid malignancies.

It has been shown that mutations in ABL1 kinase domain, ASXL1, IKZF1, TP53 are common in CML patients who develop BC >30,34. ASXL1 mutation is the most frequent mutation in poor outcome patients. In addition, individuals with an ASXL1 mutation had a longer median time to develop BC. In the current study, the first and second patients with ASXL1 mutation had longer time to MBC than the third patient.

**Table 3.** Potentially actionable (PAF) and potentially important (PIF) findings in WES analysis of BC-CML patients. P = pathogenic, LP = likely pathogenic, VUS = variant of uncertain significance, LP/P = likely pathogenic, LP/B = likely benign, VUS/P = variant of uncertain significance leaning pathogenic, VUS/B = variant of uncertain significance leaning benign, TSG = tumor suppressor gene, OG = oncogene, Dv = driver, kDv = known driver, P = passenger. Biomarker type: T = Therapeutic; D = Diagnostic; P = Prognostic; DP: Depth of coverage; VAF: Variant Allele Fraction. Leukemogenic Class: I (Signaling); II (Transcription Factor), III (Epigenetic Regulator); IV (TSG); V (Spliceosome). $ Novel variants in CML. Npa: not protein affecting.
**PRPF8** was the only RNA splicing factor gene with a significant variant. It is a core component of U2-type and U12-type spliceosome complexes. **PRPF8-G1796R** variant in the second patient is a missense Tier3 variant. This variant has been classified as driver mutation and has not been previously reported.

The self-renewal, differentiation and proliferation are three main processes which are impaired in malignancies. In the current study, the integrated genomic sequencing detected important variants in twelve different cancer genes. Mutations in signaling genes may induce proliferation of malignant cells while mutations in the epigenetic regulators (ERs) or tumor suppressor genes (TSG) may promote self-renewal in the progenitor cells and transcription factors mutations may affect differentiation. Mutations in the spliceosome component may also involve differentiation and self-renewal of progenitor cells. Therefore, the alterations in **ABL1**, **PTPN11** and **PIK3CB** signaling molecules may potentiate the proliferation process in MBC-CML. Variants in the **CHEK2**, **BRCA2** and **ATM** tumor suppressor genes, as well as the **ASXL1** and **EZH2** Epigenetic Regulators, may support the self-renewal of the proliferating progenitors. Variants in the **GATA2**, **PHF6**, **WT1**, **IKZF1** and **CUX1** transcription factors may operate as differentiation blockers. Finally, the variant in the **PRPF8** as spliceosome components may act as differentiation blocker or may cause self-renewal of the proliferating progenitor.

Finally, a recent integrated model of blast crisis showed that one of the most important mechanism associated to the BC progression is the epigenetic reprogramming related to the abnormal Polycomb Repressive Complex (PRC) function. The components of the PRC complex as an epigenetic complex, repress the expression of certain genes including leukemogenic **HOXA** cluster. On the other hand, specific gene products interact with PRC components to add/remove repressive marks or recruitment of PRC components to distinct loci. Based on recent studies on BC genome and transcriptome, it has been shown that the components of the PRC complex have mutated or altered in expression respectively. These changes overall affect the gene expression signature of BC cells. **ASXL1** gene products interact with PRC2 components including **EZH2**. **IKZF1** also recruits PRC2 to the target gene loci. According to this model, in the first and second patients the frameshift mutations in the **ASXL1** may have a critical role in BC progression. In the third patient the large deletions in the chromosome 7 which affect **IKZF1** (7p12.2) and **EZH2** (7q35-q36) may accelerate BC.

A suggested model of leukemogenesis for the hierarchy of events is presented in the Fig. 5. In this model the initial step is the presence of mildly-deleterious variants (VUS-P) mainly in the tumor suppressor genes (TSG)/DNA damage repair (DDR) genes, transcription factors (TF) and/or components of splicesome. These variants may increase the individual's susceptibility and predisposition to malignancies. The BCR-ABL1 fusion gene and mutations in the epigenetic regulator (ER)/transcription factor (TF) genes, which mainly influence the function of PRC, are the key genetic discoveries in the second and third steps. The final step in the accelerated and blast crisis phase may be started by the mutations in signaling molecules and/or additional chromosomal abnormalities (ACAs). **ABL1** mutations specifically in the kinase domain lead to TKI resistance and delay in TKI switching may affect prognosis and survival by blast crisis. ACAs which involve leukemogenic genes (such as ER, TF or TSG) also may exacerbate the clinical prognosis and enter the disease to the blastic phase.
Although mutations in \textit{ASXL1} or \textit{IKZF1} genes may play a key role in BC, they are more often found at the time of diagnosis in the chronic phase, in individuals with poor prognosis. Therefore, mutations in oncogenes such as \textit{ABL1} and \textit{PTPN11} may act as an accelerator.

In this study, samples related to the different phases of disease progression were not available. Future studies using genomic sequencing in different serial time points may allow detecting the sequence of occurrences related to the disease progression. Furthermore, recent advances in single cell sequencing may help to more accurately clarify the disease mechanisms.

Conclusions
The integrated genomic sequencing in this study effectively identified large spectrum of important and actionable variants from SNVs to CNVs. Mildly-deleterious passenger variants mainly in DNA damage repair (DDR) genes may increase the individual’s predisposition to cancer. \textit{BCR-ABL1} fusion, mutations in signaling and epigenetic regulator genes along with ACAs may act as cancer drivers. Epigenetic reprogramming caused by \textit{ASXL1} or \textit{IKZF1} mutations seems to be one of the most critical mechanisms linked with BC development. \textit{ABL1} domain mutations, mutations in other signaling genes and high risk ACAs may promote CML to BC.

Using this strategy we have identified several actionable findings which may improve targeted therapies. These findings may help better understanding the underlying pathophysiological mechanisms and may provide further insights into MBC-CML genetic basis in future. Further studies will shed light on the clinical usefulness of the integrated genomic analysis.

Data availability
The datasets generated and/or analyzed during the current study are available in the ENA https://www.ebi.ac.uk/ena/browser/view/PRJEB51084.

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References
1. Houshmand, M. \textit{et al.} Bone marrow microenvironment: The guardian of leukemia stem cells. \textit{World J. Stem Cells} \textbf{11}(8), 476–490 (2019).
2. Lebecque, B., Bourgne, C., Vidal, V. & Berger, M. G. DNA methylation and intra-clonal heterogeneity: The chronic myeloid leukemia model. \textit{Cancers (Basel)} \textbf{13}(14), 3587 (2021).
3. Valent, P. Imatinib-resistant chronic myeloid leukemia (CML): Current concepts on pathogenesis and new emerging pharmacologic approaches. \textit{Biologics} \textbf{1}(4), 433–448 (2007).
4. Murati, A. \textit{et al.} Myeloid malignancies: Mutations, models and management. \textit{BMC Cancer} \textbf{12}, 304 (2012).
5. Flis, S. & Chojnacki, T. Chronic myelogenous leukemia, a still unsolved problem: Pitfalls and new therapeutic possibilities. \textit{Drug Des. Dev. Ther.} \textbf{13}, 825–843 (2019).
6. Apperley, J. F. Chronic myeloid leukaemia. \textit{Lancet} \textbf{385}(9976), 1447–1459 (2015).
7. Bonifacio, M., Stagno, F., Scaffidi, L., Krampera, M. & Di Raimondo, F. Management of chronic myeloid leukemia in advanced phase. \textit{Front. Oncol.} \textbf{9}, 1132 (2019).
8. Avila, M. & Meric-Bernstam, F. Next-generation sequencing for the general cancer patient. Clin. Adv. Hematol. Oncol. 17(8), 447–454 (2019).
9. Chudasama, P. et al. Integrative genomic and transcriptomic analysis of leiomyosarcoma. Nat. Commun. 9(1), 144 (2018).
10. Robinson, D. R. et al. Integrative clinical genomics of metastatic cancer. Nature 548(7667), 297–303 (2017).
11. Kumar-Sinha, C. & Chinnaiyan, A. M. Precision oncology in the age of integrative genomics. Nat. Biotechnol. 36(1), 46–60 (2018).
12. Branford, S. et al. Integrative genomic analysis reveals cancer-associated mutations at diagnosis of CML in patients with high-risk disease. Blood 132(9), 948–961 (2018).
13. Beaubier, N. et al. Integrated genomic profiling expands clinical options for patients with cancer. Nat. Biotechnol. 37(11), 1351–1360 (2019).
14. Arber, D. A. et al. The 2016 revision to the World Health Organization classification of myeloid neoplasms and acute leukemia. Blood 127(20), 2391–2405 (2016).
15. Bollu, L. R., Mazumdar, A., Savage, M. L. & Brown, P. H. Molecular pathways: Targeting protein tyrosine phosphatases in cancer. Clin. Cancer Res. 23(9), 2136–2142 (2017).
16. Yang, J. et al. Targeting PI3K in cancer: mechanisms and advances in clinical trials. Mol. Cancer 18(1), 26 (2019).
17. Hantschel, O. Structure, regulation, signaling, and targeting of abl kinases in cancer. Genes Cancer 3(5–6), 436–446 (2012).
18. Jones, D. et al. Uncommon BCR-ABL kinase domain mutations in kinase inhibitor-resistant chronic myelogenous leukemia and Ph+ acute lymphoblastic leukemia show high rates of regression, suggesting weak selective effects. Blood 115(26), 5428–5429 (2010).
19. Soverini, S. et al. Next-generation sequencing for BCR-ABL1 kinase domain mutation testing in patients with chronic myeloid leukemia: A position paper. J. Hematol. Oncol. 12(1), 131 (2019).
20. Hochhaus, A. et al. Chronic myeloid leukaemia: ESMO clinical practice guidelines for diagnosis, treatment and follow-up. Ann. Oncol. 28(suppl_4), iv1–iv51 (2017).
21. Whale, A. D., Colman, L., Lensun, L., Rogers, H. L. & Shuttleworth, S. J. Functional characterization of a novel somatic oncogenic mutation of PIK3CB. Sign. Transduc. Target. Ther. 2, 17063 (2017).
22. Donadieu, J. et al. Natural history of GATA2 deficiency in a survey of 79 French and Belgian patients. Haematologica 103(8), 1278–1287 (2018).
23. Gao, J. et al. Heritable GATA2 mutations associated with familial AML–MDS: A case report and review of literature. J. Hematol. Oncol. 7, 36 (2014).
24. Adnan-Awad, S., Kankainen, M. & Mustjoki, S. Mutational landscape of childhood acute leukemia: More than a single oncogene lesion. Leuk. Lymphoma 62(9), 2064–2078 (2021).
25. Zhang, X. et al. The specific distribution pattern of IKZF1 mutation in acute myeloid leukemia. J. Hematol. Oncol. 13(1), 140 (2020).
26. Vгадиас, C. et al. CUX1 transcription factor is required for optimal ATM/ATR-mediated responses to DNA damage. Nucleic Acids Res. 40(10), 4483–4495 (2012).
27. Higgins, A. & Shah, M. V. Genetic and genomic landscape of secondary and therapy-related acute myeloid leukemia. Genes (Basel) 11(7), 749 (2020).
28. Siefert, H. et al. The prognostic impact of 17p (p53) deletion in 2272 adults with acute myeloid leukemia. Leukemia 23(4), 656–663 (2009).
29. Nakai, H. & Misawa, S. Chromosome 17 abnormalities and inactivation of the p53 gene in chronic myeloid leukemia and their prognostic significance. Leuk. Lymphoma 19(3–4), 213–221 (1995).
30. Hehlmann, R. Chronic myeloid leukemia in 2020. Hemasphere 4(5), e468 (2020).
31. Yang, L., Han, Y., Suarez Saiz, F. & Minden, M. D. A tumor suppressor and oncogene: The WT1 story. Leukemia 21(3), 868–876 (2007).
32. Perrotti, D., Jamieson, C., Goldman, J. & Skorski, T. Chronic myeloid leukemia: Mechanisms of ablastic transformation. J. Clin. Invest. 120(7), 2254–2264 (2010).
33. Hosoya, N., Miyagawa, K., Mitani, K., Yasaki, Y. & Hirai, H. Mutation analysis of the WT1 gene in myelodysplastic syndromes. Int. J. Cancer Res. 89(8), 821–824 (1998).
34. Grossmann, V. et al. A deep-sequencing study of chronic myeloid leukemia patients in blast crisis (BC-CML) detects mutations in 76.9% of cases. Leukemia 25(3), 557–60 (2011).
35. Wolf, A. et al. Single base-pair substitutions at the translation initiation sites of human genes as a cause of inherited disease. Hum. Mutat. 32(10), 1137–1143 (2011).
36. Phan, L. M. & Rezaeian, A. H. ATM: main features, signaling pathways, and its diverse roles in DNA damage response, tumor suppression, and cancer development. Genes (Basel) 12(6), 845 (2021).
37. Takagi, M. et al. ATM-dependent DNA damage-response pathway as a determinant in chronic myelogenous leukemia. DNA Repair (Amst.) 12(7), 500–507 (2013).
38. Prakash, R., Zhang, Y., Feng, W. & Jasim, M. Homologous recombination and human health: The roles of BRCA1, BRCA2, and associated proteins. Cold Spring Harb. Perspect. Biol. 7(4), a06600 (2015).
39. Fica, S. M., Oubridge, C., Wilkinson, M. E., Newman, A. J. & Nagai, K. A human postcataclystic spliceosome structure reveals essential roles of metazoan factors for exon ligation. Science 363(6428), 710–714 (2019).
40. Aponte, P. M. & Caicedo, A. Stemness in cancer: Stem cells, cancer stem cells, and their microenvironment. Stem Cells Int. 2017, 5619472 (2017).
41. Ko, T. K. et al. An integrative model of pathway convergence in genetically heterogeneous blast crisis chronic myeloid leukemia. Blood 135(26), 2337–2353 (2020).

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Author contributions
G.E.K.S. made contribution to the conception and data analysis, drafted the work and wrote the manuscript. M.K made contribution to the conception and data analysis. K.M. have made contribution to the conception, supervised the project and encouraged the investigation. S.T. suggested the main conceptual ideas and proofed outline and critically reviewed and revised the article for important intellectual content. M.V and S.M.R had role in the clinical management of the patient and contributed to the clinical analysis. K.K contributed in the NGS data analysis. A.T provided critical feedback and contributed in the NGS data analysis. Sh.R, M.Y and B.Ch contributed in performing hematological, cytogenetic and immunophenotyping studies and related data analysis. All authors read and approved the final manuscript. Golnaz Ensiieh Kazemi Sefat (G.E.K.S.) and Mohammad
Keramatipour (M.K.) contributed equally as the first authors. Saeed Talebi (S.T) and Kazem Mousavizadeh (K.M) contributed equally as corresponding authors. All authors reviewed the manuscript.

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**Competing interests**
The authors declare no competing interests.

**Additional information**

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