Next-generation sequencing identifies major DNA methylation changes during progression of Ph+ chronic myeloid leukemia

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INTRODUCTION

Chronic myeloid leukemia (CML) is a stem cell-derived malignancy characterized by the expansion and accumulation of myeloid cells in the blood and bone marrow (BM), the chromosomal translocation t(9;22) and the associated oncoprotein BCR-ABL1. The disease can be divided into a chronic phase (CP), accelerated phase (AP) and blast crisis (BC). In untreated and drug-resistant patients, evolution of CP to AP and finally to BC reflects the natural course of disease. The prognosis of patients in AP and BC is poor unless these patients can undergo stem cell transplantation.

In CP-CML, BCR-ABL1 is a key driver of oncogenesis, cellular proliferation and survival. Until now, only few BCR-ABL1-independent genetic lesions (for example, c-Myc copy number gain, RB1 mutations) are known to be involved in CML progression. The potential contribution of epigenetic changes, especially of DNA methylation (referred to as methylation) to CML progression, has not been investigated in detail so far.

Methylation is an epigenetic modification involved in the regulation of many biological processes, including embryogenesis, genomic imprinting or X chromosome inactivation. In the mammalian genome, cytosines within CpG sites are the main targets of methylation. CpG sites are spread throughout the genome but are mainly located in small regions of about 0.5–2 kb in length, called CpG islands. These regions are found in ~60% of human gene promoters and less frequently in gene bodies or in intergenic regions. However, in total CpG sites are under-represented in the mammalian genome. Transcriptional gene silencing is associated with methylation of CpG sites and CpG islands located near the transcription start sites (TSS) of genes. In cancer cells, these regions may be frequently methylated resulting in transcriptional gene silencing of many cancer-associated genes. Although methylation of CpG sites in gene bodies may contribute to cancer-causing somatic and germline mutations, the function of intergenic CpG site methylation is barely understood so far. As methylation is reversible, methylated genes may be re-expressed by the use of DNA methyltransferase inhibitors (DNMTi; for example, 5-aza-2′-deoxycytidine (Aza-dC); 5-azacytidine). More- ever, DNMTi and histone deacetylase inhibitors (for example, trichostatin A (TSA)) act synergistically on gene re-expression.

Methylation is considered biologically important in the pathogenesis of many malignant diseases. Therefore, we were interested to study the methylome in CML patients at diagnosis as well as at disease progression. Using reduced representation bisulfite sequencing (RRBS), we analyzed samples from CP, AP and BC of CML patients and from control individuals for differences in CpG site methylation. Moreover, we performed RNA-sequencing (RNA-seq) to study gene expression in these samples.

Overall, we were able (1) to demonstrate that CpG site methylation increases dramatically during the progression from CP-CML to AP-CML/BC-CML and (2) to identify genes that are transcriptionally regulated by methylation in BC-CML samples. Some of these genes are known to be involved in the pathogenesis of various malignancies; however, the role of many other genes in the pathogenesis and progression of malignant diseases, particularly in CML, remains to be determined.
MATERIALS AND METHODS

CML samples and cell lines
Mononuclear cells were isolated from peripheral blood (PB) or BM of 31 samples (17 CP-CML, 5 AP-CML, 9 BC-CML) from 23 CML patients using Ficoll (Sigma, St Louis, MO, USA). PB and BM samples were collected from all patients at diagnosis and from a subset of four patients during their follow-up. All four patients progressed to AP or BC during therapy with BCR-ABL1 tyrosine kinase inhibitors. Clinicopathological characteristics of the patients are shown in Supplementary Table S1. Mononuclear cells obtained from five individuals were used as controls. All donors gave written informed consent. The study was approved by the ethics committee of the Medical University of Vienna, Austria (no. 1594/2015).

Authenticated cell lines K562, KCL22, NALM-1, TOM-1 and BV-173 were purchased from the Leibniz Institute DSMZ (Braunschweig, Germany). KU812 cells were kindly provided by Dr K Kishi (Niigata University, Niigata, Japan), and K562 R cells (matrinin-resistant) were kindly provided by James D Griffin (Dana Farber Cancer Institute, Boston, MA, USA). Cells were cultured in RPMI1640 medium with 10% fetal calf serum at 37 °C. Mycoplasma contamination was tested using the Venor GeM Classic Mycoplasma Detection Kit (Minerva Biolabs, Berlin, Germany). Aza-dC and TSA treatment was performed as reported.11 Untreated control cells were cultured in parallel.

Reduced representation bisulfite sequencing
Genomic DNA was MspI (New England Biolabs, NEB, Ipswich, MA, USA) digested, end-repaired and A-tailed using Klenow Polynucleotise (NEB). Illumina adapters were ligated using Quick Ligase (NEB) followed by AMPure XP size selection (Beckman Coulter, Fullerton, CA, USA). RRBS libraries were bisulfite treated using the EZ-DNA Methylation Direct Kit (Zymo Research Corp, Orange, CA, USA) followed by quantitative PCR quantification. Enrichment PCR was performed using the PhiTuRoxC Hotstart Kit (Agilent Technologies, Santa Clara, CA, USA) followed by Ampure XP clean up. Quality of final libraries was checked by Experion analysis (Bio-Rad, Hercules, CA, USA). Sequencing was performed using the Illumina HiSeq2000 sequencer (Illumina Inc, San Diego, CA, USA). A detailed description of the RRBS assay is provided as Supplementary Information.

RNA-sequencing
Total RNA was extracted using the RNeasy Kit (Qiagen, Hilden, Germany) and processed for sequencing using the TruSeq RNA Sample Preparation Kit (Illumina Inc) as described.27 miRNAs were purified using poly(T)-oligo-attached magnetic beads, fragmented and applied to first-strand complementary DNA (cDNA) synthesis. Second-strand cDNA synthesis was performed using DNA polymerase I and RNase H. cDNAs were end-repaired, A-tailed, ligated to adapters and amplified to create the final cDNA library for sequencing (HiSeq2000, Illumina Inc).

Methylation-sensitive high-resolution melting (MS-HRM) analyses
Genomic DNA was modified by treatment with sodium bisulfite using the EpiTec Bisulfite Kit (Qiagen) as recommended.24 The following primer sequences for methylation-insensitive amplification of a part of the CYP1B1 5′ region (ENSEMBL database, release 69) were designed using the Methyl Primer Express v1.0 software (Applied Biosystems, Carlsbad, CA, USA): fwd, 5′-GGTTAAAGYGGTTTGTGTF-3′; and rev, 5′-CTCCCACCTCCAAAAATAACAAAA-3′. MS-HRM analyses were performed using the EpiTec HRM PCR Kit in a RotorGene Q cycler (Qiagen).25 Methylation standards were constructed by diluting 100% methylated and unmethylated control DNA (Qiagen) at 100, 75, 50, 25, 10 and 0% ratios. Melting curves were normalized by calculation of two normalization regions before and after the major fluorescence decrease using the RotorGeneQ software. Normalized fluorescence values were plotted against the percentage of methylation for each of the methylation standards to generate a standard curve for the calculation of methylation levels of genes in leukemia cells. Water blanks were used as negative controls.

Real-time reverse transcription-PCR (RT-PCR)
Total RNA was reverse-transcribed using the Omniscript Reverse Transcriptase Kit (Qiagen), and cDNA was used for standard RT-PCR analyses. Detailed information about used RT-PCR assays is provided as Supplementary Information.

Statistical analyses
RRBS reads were processed using the RRBS tool of the Trim Galore! software (http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/).

Bismark software was used to align reads to hg19 and for methylation calling.24 Codes are available from Trim Galore! and bismark documentation. Bam files from RRBS analyses were deposited in ArrayExpress database (E-MTAB-4341). Methylation was quantified using the ‘Bisulfite methylation over features’ pipeline of the SeqMonk v0.27.0 software (Babraham Institute, Cambridge, UK). Differential methylation (increased or decreased) was calculated using a windowed replicate filter over a window of 1000 bp using the following cutoff levels: at least 25% difference in methylation and false discovery rate (FDR) < 0.05. Hierarchical clustering was performed using the hclust function of R 2.15.0. Circos software (Canada’s Michael Smith Genome Sciences Center, Vancouver, BC, Canada) was used to visualize RRBS data.26 The R packages MethylSig and MethylKit were used for quality control, calculation of correlation coefficients to describe similarities of CpG site methylation between samples, for CpG island/Refseq annotation and for transcription factor (TF)-binding site enrichment analyses. Pathway enrichment analyses were performed using the GREAT software (Stanford University, Stanford, CA, USA; default settings; binomial test and hypergeometric test cutoff: FDR < 0.05).

Galaxy platform and TopHat2 algorithm were used to align raw RNA-seq data to hg19.27-30 Bam files from RNA-seq analyses were deposited in ArrayExpress database (E-MTAB-4333). Aligned bam files were processed using RNA-seq quantitation pipeline of the SeqMonk v0.27.0 software (Babraham Institute). An adjusted P < 0.05 was defined as cutoff for differentially expressed genes. Single-nucleotide variants (SNVs) were identified using RNAseqmut (https://github.com/davidlw/rnaseqmut) and verified using IGV.31

R software (https://cran.r-project.org/) and GraphPad Prism v.6.03 (GraphPad Software, Inc., San Diego, CA, USA) were used to calculate Bartlett tests, t-tests and Mann–Whitney U-tests. P-values of < 0.05 (two-sided) were considered as significant.

RESULTS

Comprehensive CpG site methylation analyses in CML and in control samples
We investigated the methylene of 17 CP-CML, 5 AP-CML, 9 BC-CML and 5 control samples using RRBS. The bisulfite conversion rate of mapped RRBS reads was > 99% for all samples analyzed. A mean of 1.4 × 107 cytosines were analyzed per sample and about 20% (2.6 × 105) of these cytosines were located in Cpg sites. Examples of RRBS library quality and RRBS data quality are shown in Supplementary Figures S1 and S2.

When we compared the content of methylated CpG sites in CML samples with that in control samples, we observed a higher content in CML samples (Figure 1). Moreover, a higher content of methylated CpG sites was also seen in BC-CML samples compared
with CP-CML samples \( (P = 0.02, \text{Figure 1}) \). However, no statistically significant difference in the content of methylated CpG sites between AP-CML and CP-CML samples was found.

Hierarchical clustering of CML samples and of control samples based on CpG site methylation clearly separated CP-CML, AP-CML and BC-CML samples from control samples (Figure 2a and Supplementary Figure S3A).

Detailed comparison of CpG site methylation between CML and control samples

Analyses were performed separately for the PB (CP-CML, \( n = 6 \); BC-CML, \( n = 4 \); controls, \( n = 3 \)) and BM (CP-CML, \( n = 11 \); AP-CML, \( n = 4 \); BC-CML, \( n = 5 \); controls, \( n = 2 \)) sample groups. In the PB cohort, 666 CpG sites (0.4% of the CpG sites analyzed) were differentially methylated when we compared CP-CML and control samples. While methylation of 292 CpG sites was increased, methylation of 374 CpG sites was decreased (Figure 2b). Six thousand nine hundred and twelve CpG sites (2.3% of the CpG sites analyzed) were differentially methylated when we compared BC-CML and control samples. While methylation of 6081 CpG sites was increased, methylation of 831 CpG sites was decreased (Figure 2b). All these differences were statistically significant.

Similar results were observed in the BM cohort. Two hundred and seventy-six CpG sites (0.2% of the CpG sites analyzed) were differentially methylated between CP-CML and control samples. Increased methylation was observed in 237 CpG sites and decreased methylation in 39 CpG sites. Six thousand one hundred and one CpG sites (1.8% of the CpG sites analyzed) were differentially methylated between BC-CML and control samples and increased methylation was observed in 5729 CpG sites, whereas decreased methylation was found in 372 CpG sites (Figure 2b). Moreover, different CpG site methylation was observed in AP-CML samples compared with control samples. Four thousand one hundred and thirty CpG sites (0.9% of the CpG sites analyzed) were differentially methylated and increased methylation was observed in 3575 CpG sites, whereas decreased methylation was found in 555 CpG sites (Supplementary Figure S3B). Again, all these differences were statistically significant.

From the CpG sites found to be differentially methylated in AP-CML/BC-CML samples, we also determined their location within, around and between CpG islands. The majority of CpG sites with increased methylation (88%; range 87–89%) were located in or close to CpG islands and only 12% were located in between CpG islands \( (P < 0.0001) \). In contrast, about half of the CpG sites with decreased methylation (49%; range 45–53%) were located in between CpG islands mainly in introns and intergenic regions.

Additionally, we assigned CpG sites with increased methylation in AP-CML/BC-CML samples to genomic regions located around TSS of genes (±2000 bp), and we identified 348 methylated genes in BC-CML samples from the PB cohort and 141 and 292 methylated genes in AP-CML and BC-CML samples from the BM cohort, respectively (Figure 3). A detailed description of these genes is listed in Supplementary Tables S2–S4.

CpG site methylation, gene expression and SNV analyses of individual CML patients

Comparison of the methylome of CML patients at different stages of disease. To determine differences between CP-CML and disease progression, we compared CpG site methylation in samples of four patients who were initially diagnosed with CP-CML and who progressed several months/years later (Supplementary Table S1, Figure 4a). From patient 1, PB samples from three different time points were available (CP-CML, CP-CML ~ 5 years later and AP-CML). Overall, we identified 20 172 CpG sites that were differentially methylated in the AP-CML sample compared with the CP-CML samples of this patient. In 72% of these CpG sites, methylation was increased. Interestingly, only 1966 differentially methylated CpG sites were identified when we compared the 2 CP-CML samples of this patient. This finding suggests that CpG site methylation does not differ significantly between various time points during CP-CML even when they are several years apart.

Patients 2–4 were initially diagnosed with CP-CML and developed BC-CML after several months or years. PB samples from patients 2 and 3 and BM samples from patient 4 from two different time points were available for analyses, respectively (Figure 4a). We identified 29 280, 58 691 and 28 268 differentially methylated CpG sites in patients 2–4 in the BC-CML samples compared with the CP-CML samples, respectively. In 96, 89 and 67% of these CpG sites, methylation was increased.

In patients 1–3, increased CpG site methylation was accompanied with increased BCR-ABL1 expression determined by RT-PCR analyses.
using GUSB as well as ABL1 as reference genes (Figure 4a). In patient 4, material for BCR-ABL1 testing by RT-PCR analyses using GUSB as reference gene was used up; however, BCR-ABL1 expression in both samples of this patient was observed in previous RT-PCR analyses using ABL1 as reference gene.

Genomic annotation of differentially methylated CpG sites. We additionally determined the location of differentially methylated CpG sites within, around and between CpG islands in the samples from these four patients. The majority of CpG sites with increased methylation (87%; range 82–92%) were located in or close to CpG islands. Only 13% of them were located in between CpG islands (P < 0.0001; Supplementary Figure S4A) and were evenly distributed in promoter, gene body and intergenic regions. In contrast, about half of the CpG sites with decreased methylation (54%; range 45–63%) were located outside of CpG islands mainly in introns and intergenic regions (Supplementary Figure S4B).

Moreover, we were interested to identify TFs whose activity might be affected by methylation of their binding sites. By TF-binding site enrichment analyses in the samples of

![Figure 3](image-url)

**Figure 3.** Heatmaps summarizing methylation values of genes with increased methylation in AP-CML and BC-CML samples compared with control samples. (a) 348 genes with increased methylation in BC-CML samples compared with control samples (PB cohort), (b) 141 genes with increased methylation in AP-CML samples compared with control samples (BM cohort) and (c) 292 genes with increased methylation in BC-CML samples compared with control samples (BM cohort) are shown. The colors range from blue (low methylation) to red (high methylation). Mean methylation values are shown. Ctrl, controls.

![Figure 4](image-url)

**Figure 4.** CpG site methylation in four CML patients at diagnosis and at disease progression. (a) The time from first diagnosis of CP-CML to disease progression, BCR-ABL1 count (Ph+), blast count and therapies are shown. (b) The circos plots demonstrate the mean percentage of CpG site methylation of genomic regions located ± 2000 bp from TSS of genes. Normal peripheral blood (nPB) and normal bone marrow (nBM) samples from control individuals were used as references. na, not available.
these four patients, we observed an overrepresentation of CpG sites with increased methylation in several TF-binding sites. These TF-binding sites include C-myc (*P* = 5 × 10^{-16}), nuclear factor-κB (*P* = 1 × 10^{-16}), C-fos (*P* = 5 × 10^{-13}) and YY1 (*P* = 2 × 10^{-10}). Detailed results are shown in Supplementary Table S5.

To investigate whether CpG sites with increased methylation in the samples of these four patients are associated with specific biological processes or molecular functions, we performed Gene Ontology (GO) and molecular signature (MSigDB) enrichment analyses. Highly enriched GO/MSigDB terms were mainly associated with cell differentiation, developmental processes, transcriptional gene regulation, targets of polycomb proteins (EED, SUZ12, PRC2), H3K27 trimethylation and gene hypermethylation in cells of different malignant diseases (Supplementary Table S6).

In addition, we assigned CpG sites with increased methylation in AP-CML/BC-CML samples to genomic regions located around TSS of genes (±2000 bp), and we identified 344 (patient 1), 381 (patient 2), 897 (patient 3) and 597 (patient 4) methylated genes in these samples, respectively (Figure 4b). A detailed description of the genes is listed in Supplementary Tables S7–S10. Three hundred and sixty-three of these genes were methylated in samples of at least two of the patients (Figure 5).

Transcriptional gene expression of methylated genes

From patients 3 and 4 also total RNA from CP-CML and BC-CML samples was available and was used together with total RNA samples from four control individuals for RNA-sequencing analyses to investigate gene expression. Overall, the median expression of methylated genes was lower in BC-CML samples compared with both CP-CML and control samples (Figure 6a). Expression of 193 out of 897 (patient 3) and of 141 out of 597 (patient 4) methylated genes in BC-CML samples was downregulated less than twofold in BC-CML samples compared with CP-CML samples of these patients, respectively (Figure 6a). Compared with control samples, expression of a large majority of them was downregulated in the BC-CML samples. Some of these genes are known tumor-suppressor genes (EPP41L3, PRDX2), putative tumor-suppressor genes (PLCL1, TUSC1), regulators of cell proliferation (BCL11B, NDRG2, PID1) or regulators of drug metabolism (CYP1B1).21,32–35 A description of the downregulated genes is shown in Supplementary Tables S11 and S12.

We confirmed CYP1B1, EPB41L3 and PID1 expression data obtained by RNA-seq using RT-PCR in CP-CML and BC-CML samples of patient 3 and in control samples (Figure 6b and Supplementary Figure S5). In addition, we determined both CYP1B1 expression and methylation by MS-HRM in seven leukemia cell lines. In five of them, CYP1B1 was methylated and was not transcribed (Figure 6c). To investigate whether transcription of CYP1B1 may be re-expressed by epigenetically active drugs, we treated K562 and K562 R cells with Aza-dC and TSA and performed RT-PCR. CYP1B1 expression was found to be upregulated 218-fold in K562 cells and 9647-fold in K562 R cells (Figure 6d).

![Figure 5](image-url)

Figure 5. Overlap of genes found to be methylated in BC-CML samples from patients 2–4 who were initially diagnosed with CP-CML and who progressed to BC-CML. The circos plot represents methylation values in CP-CML (outer ring) and BC-CML (inner ring) determined by RRBS. Patients 2 (blue), 3 (green) and 4 (pink) are shown. Blue lines indicate genes that are methylated in two of the patients and red lines indicate genes that are methylated in all three patients.
SNV analyses. When we compared the mutational landscapes of CP-CML and BC-CML samples from patients 3 and 4, we found 73 (patient 3) and 248 (patient 4) SNVs to be present only in BC-CML samples, respectively (Supplementary Tables S13 and S14). Examples for non-synonymous SNVs are ABL1 (p.L248V), NR3C1 (p.G239X), PLK4 (p.S200T) and VRK3 (p.V290A) (Supplementary Figures S6 and S7). Interestingly, BC-CML-specific SNVs of certain epigenetic modifiers (DNMT1, DNMT3A, TET2, TET3, EZH1, EZH2, RUNX1, IKZF1, IDH1 and IDH2) and of certain oncogenes (MYC, RUNX1, MDM2, BCL2, NFKB1, NFKB2, KRAS, NRAS, HRAS, RAF1, MEK/MAP2K1, ERK/MAPK1) were not found. Moreover, no BC-CML-specific karyotypic changes were observed in patients 1, 3 and 4.

**DISCUSSION**

Although changes in methylation patterns were widely analyzed in various malignant diseases, data generated on the impact of methylation in CML are limited and mainly restricted to analyses of single genes or small numbers of genes.\(^5,6,16,36,38\) However, as recently developed techniques for detecting methylation are much more advanced, we were able to perform extensive methylation analyses in CML samples using the next-generation sequencing approach RRBS. The main advantages of this method compared with microarray-based techniques are the high sensitivity, the high CpG coverage, the single-nucleotide resolution and the low input DNA requirement.\(^39-41\)

On average, we analyzed \(2.6 \times 10^6\) CpG sites in each CML and each control sample for methylation. Overall, the frequency of methylated CpG sites was significantly higher in CML samples compared with control samples, indicating that CpG site methylation is involved in CML. Interestingly, we observed differences in CpG site methylation between CP-, AP- and BC-CML samples. In CP-CML samples, the frequency of CpG site methylation was not significantly different from control samples, suggesting that CpG site methylation does not have a central role in CP-CML. These findings support the assumption that CP-CML is mainly driven by the BCR-ABL1 translocation.\(^42\) However, we observed a high increase of methylated CpG sites in AP and BC, indicating that CpG site methylation may be important for disease progression.

To investigate differences in CpG site methylation between CP-CML and CML progression in detail, we compared the methylome of four CML patients at the time of diagnosis (CP-CML) and at the time of disease progression (AP-CML or BC-CML). Although we found similar frequencies of differentially methylated CpG sites between both CP-CML samples of patient 1, we observed a significant increase of differentially methylated CpG sites in AP-CML samples of this patient. However, the highest increase of differentially methylated CpG sites (up to \(58\,691\)) was found in BC-CML samples. In the majority of CpG sites, methylation was increased, suggesting that increased but not decreased CpG site methylation has an important role in the progression of this disease.

Genomic annotation of CpG sites with increased methylation in AP-CML/BC-CML revealed that they are mainly located in or around CpG islands. Our results suggest that these CpG sites may be involved in regulation of transcriptional gene expression as CpG island methylation is strongly associated with gene silencing.\(^13\) Moreover, many methylated CpG sites were found in the binding sites of certain TFs. This finding is of interest as it was reported that methylation of TF-binding sites may deregulate transcriptional gene expression by changing the structure of DNA or by altering the affinity of TFs to DNA.\(^43-45\) In addition, GO/MSigDB enrichment analyses revealed that increased CpG site methylation frequently
happens at genomic regions prone for epigenetic modifications associated with gene silencing. Similar findings were reported by Viré et al. who demonstrated that polycomb group proteins, H3K27 methylation and CpG site methylation are closely linked in epigenetic gene silencing.

Further assignment of CpG sites with increased methylation in BC-CML samples to genomic regions ± 2000 bp from TSS of genes resulted in the identification of up to 897 genes. Many of them were methylated in the BC-CML samples of at least two of the three patients, suggesting that changes in methylation during disease progression are not random events. Although methylation of some of these genes (for example, TAF2PA, EBFB2) was previously reported in CML, methylation of the majority of them was unknown so far. Interestingly, some of them were described to be methylated in other hematological and/or solid malignancies, for example, BNIP3 (multiple myeloma, gastric cancer), HIC1 (myelodysplastic syndrome), CADMI1 and EPB41L3 (lung and breast cancers), PDLIM4 and CIDEB (neoplastic mast cells) and CDH1 (myelodysplastic syndrome).

We also performed genome-wide gene expression analyses in CP-CML and BC-CML samples of two patients who were initially diagnosed with CP-CML and who progressed to BC-CML. These samples were also analyzed by RRB. Expression of many genes identified to be methylated in BC-CML samples was found to be downregulated in these samples, suggesting that expression of them is affected by methylation. Examples are the known/putative tumor-suppressor genes CYP1B1, EPB41L3 and PDI1, which downregulated expression observed by RNA-seq was confirmed by RT-qPCR. CYP1B1 belongs to the cytochrome P450 family, is involved in drug metabolism and steroid synthesis and was found to be associated with a shorter survival of acute lymphocytic leukemia patients. EPB41L3 is involved in cell adhesion, regulation of cell motility, cell proliferation and apoptosis and is frequently inactivated by methylation in various cancer types. PDI1 is a tumor cell growth suppressor and a low PDI1 expression was found to be associated with shorter overall survival of patients with brain tumors. Additional experiments to define the role of these genes during CML progression will be performed.

Because we demonstrated that treatment of CML cells (both sensitive and resistant to imatinib) with Aza-daC lead to gene re-expression and that CpG site methylation is significantly increased in AP-CML/BC-CML samples, we hypothesize that besides BCR-ABL1 tyrosine kinase inhibitor also DNMTi (for example, decitabine) might be useful for the treatment of advanced-stage CML patients. Indeed, Issa et al. reported a complete and partial hematological response rate in 34% and 20% of imatinib-resistant patients treated with low-dose decitabine, respectively. Another study described a complete hematological response rate in 32% and a partial hematological response rate in 4% of AP-CML/BC-CML patients treated with decitabine and imatinib. The efficacy of the combination decitabine and the BCR-ABL1 tyrosine kinase inhibitor dasatinib is currently being investigated in AP-CML/BC-CML patients in a clinical trial (ClinicalTrials.gov, NCT01498445).

It was reported that SNVs may occur in epigenetic modifiers in CML patients. While Schmidt et al. found SNVs in the epigenetic modifiers DNMT3A, EZH2, RUNX1 and TET2 in CP-CML samples, Makishima et al. observed BC-CML-specific SNVs in TET2, ASXL1 and IDH1 family genes when they compared BC-CML and CP-CML samples. However, in our sample collection we neither detected BC-CML-specific SNVs of any of these genes nor of any other epigenetic modifiers investigated which suggests that their role in CML, and particularly in disease progression, is still unclear. Additional studies are necessary to clarify the occurrence of molecular changes of epigenetic modifiers and to investigate a potential role of them in methylation changes in CML patients.

In conclusion, using a novel next-generation sequencing approach, we were able to demonstrate that CpG site methylation is heavily increased in AP-CML/BC-CML compared with CP-CML. Moreover, a large number of genes transcriptionally regulated by methylation were identified in BC-CML samples. From the majority of these genes, methylation in CML was unknown so far. Finally, our findings suggest that investigating the efficacy of DNMTi in patients with progressed CML should be considered.

**CONFLICT OF INTEREST**

PV received research grants from Novartis and Ariad. The other authors declare no conflict of interest.

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