Whole-Genome DNA Methylation Sequencing Reveals Epigenetic Changes in Myelodysplastic Syndromes

Jing-dong Zhou1,2,3, Zi-jun Xu2,3,4, Ye Jin1,2,3, Xin-long Zhang5, Yu Gu1,2,3, Ji-chun Ma2,3,4, Xiang-mei Wen2,3,4, Jiang Lin2,3,4, Ting-juan Zhang5,6* and Jun Qian1,2,3*

1 Department of Hematology, Affiliated People’s Hospital of Jiangsu University, Zhenjiang, China, 2 Zhenjiang Clinical Research Center of Hematology, Zhenjiang, China, 3 The Key Lab of Precision Diagnosis and Treatment in Hematologic Malignancies of Zhenjiang City, Zhenjiang, China, 4 Laboratory Center, Affiliated People’s Hospital of Jiangsu University, Zhenjiang, China, 5 Department of Hematology, The People’s Hospital of Danyang, Zhenjiang, China, 6 Department of Oncology, Affiliated People’s Hospital of Jiangsu University, Zhenjiang, China

Epigenetic dysregulation of cancer-associated genes has been identified to contribute to the pathogenesis of myelodysplastic syndromes (MDS). However, few studies have elucidated the whole-genome DNA methylation in the initiation pathogenesis of MDS. Reduced representation bisulfite sequencing was performed in five de novo MDS patients and four controls to investigate epigenetic alterations in MDS pathogenesis. The mean global methylation in five MDS patients showed no significant difference compared with the four controls. In depth, a total of 1,459 differentially methylated fragments, including 759 hypermethylated and 700 hypomethylated fragments, were identified between MDS patients and controls. Targeted bisulfite sequencing further identified that hypermethylation of DLEU7, FOXR1, LEP, and PANX2 were frequent events in an additional cohort of MDS patients. Subsequently, LEP hypermethylation was confirmed by real-time quantitative methylation-specific PCR in an expanded cohort of larger MDS patients. In clinics, LEP hypermethylation tended to be associated with lower bone marrow blasts and was significantly correlated with U2AF1 mutation. Survival analysis indicated that LEP hypermethylation was associated with a markedly longer survival time but was not an independent prognostic biomarker in MDS patients. Functional studies revealed pro-proliferative and anti-apoptotic effects of leptin in the MDS cell line SKM-1, and it was significantly associated with cell growth and death as well as the Toll-like receptor and NF-kappa B signaling pathways. Collectively, our findings demonstrated that whole-genome DNA methylation analysis identified novel epigenetic alterations such as DLEU7, FOXR1, LEP, and PANX2 methylations as frequent events in MDS. Moreover, LEP might play a role in MDS pathogenesis, and LEP hypermethylation was associated with longer survival but not as an independent prognostic biomarker in MDS.

Keywords: methylation, LEP, myelodysplastic syndromes, genome, epigenetics
INTRODUCTION

Myelodysplastic syndromes (MDSs) represent a group of diverse clonal hematopoietic disorders characterized by peripheral blood cytopenia, ineffective production of blood cells, and high risk of transformation to acute myeloid leukemia (AML) (1). Cytogenetic abnormalities and genetic mutations play crucial roles in the pathogenesis of MDS, and have been proved to be clearly associated with MDS diagnosis and prognosis (2). Moreover, epigenetic dysregulation of cancer-associated genes has been identified as contributing to the pathogenesis of MDS (3). In particular, hypermethylation of CpG islands (CGIs) located at the promoter region of tumor suppressor genes (TSGs) and their consequent gene silencing have been revealed in MDS (4). Recently, epigenetic alterations as new biological makers have also been widely used for predicting prognosis and risk of AML in patients with MDS (5–7). Importantly, epigenetic therapies with demethylating agents such as azacytidine and decitabine have demonstrated clinical effectiveness and have been approved by the Food and Drug Administration (FDA) as antitumor agents for the treatment of MDS (8, 9).

Previously, we determined the genome-wide DNA methylation alterations during MDS progression by reduced representation bisulfite sequencing (RRBS) and revealed that genome-wide DNA hypermethylation changes were a common phenomenon during MDS progression (10). Moreover, hypermethylation of ZNF300, DLX5, SOX30, ID4, and GPX3 genes was associated with the prognosis and disease progression of MDS (10–14). However, only a few studies have elucidated the whole-genome DNA methylation in the initiation pathway of MDS. Based on our previous study, we further re-analyzed the RRBS data of bone marrow (BM) samples from five de novo MDS patients and four controls to investigate epigenetic alterations in MDS pathogenesis.

METHODS

Patients and Samples

In this study, three independent cohorts of MDS patients and controls were included after informed consent was obtained. Firstly, a total of five de novo MDS patients and four healthy donors from the Affiliated People’s Hospital of Jiangsu University and the First Affiliated Hospital of Soochow University were included in the RRBS. Next, another cohort of 36 de novo MDS and 25 healthy donors treated at the Affiliated People’s Hospital of Jiangsu University was used in the targeted bisulfite sequencing. Lastly, the third cohort of 105 de novo MDS and 46 healthy donors treated at the Affiliated People’s Hospital of Jiangsu University was included in the real-time quantitative methylation-specific PCR (RQ-MSP) analysis. BM was collected from all MDS patients at the time of diagnosis with controls. BM mononuclear cells (BMMNCs) were separated by density-gradient centrifugation using Lymphocyte Separation Medium (Solarbio, Beijing, China). Subsequently, DNA extraction was carried out based on the instructions of the manufacturer (10). This study was approved by the Ethics Committee of Affiliated People’s Hospital of Jiangsu University.

RRBS and Targeted Bisulfite Sequencing

RRBS and targeted bisulfite sequencing (MethylTarget) were performed by Genesky Biotechnologies Inc. (Shanghai, China). A detailed description of the RRBS and MethylTarget assay was described in our previous report (10). The primers of the selected genes used in MethylTarget are shown in Table S1.

Bisulfite Modification and RQ-MSP

In our previous literature (10), bisulfite conversion of genomic DNA was reported. RQ-MSP was further performed to quickly detect the methylation level of LEP by using AceQ qPCR SYBR Green Master Mix (Vazyme Biotech Co., Piscataway, NJ) as reported (15). Detailed information regarding the PCR can be referred to in our previous literature (15).

Cell Lines, Cell Culture, and Reagents

The MDS cell line SKM-1 was cultured in RPMI 1640 medium containing 10% fetal calf serum (ExCell Bio, Shanghai, China) and grown at 37°C in a 5% CO2 humidified atmosphere (10). Human recombinant human leptin (R&D Systems, Minneapolis, MN) was dissolved in the medium to a working concentration of 100 ng/ml.

Cell Growth Assays

The tested cells were seeded in 96-well plates (at a density of 5 × 10^3 cells/well) in triplicate and cultured for 0, 24, 48, and 72 h, respectively. A Cell Counting Kit-8 (Dojindo, Kumamoto, Japan) was added to each well and incubated for 2 h, and was measured using a microplate reader at the absorbance at 450 nm. The rate of cell growth was calculated by the OD value.

Cell Apoptosis Assays

The tested cells were cultured with serum-free RPMI 1640 medium for 48 h in 6-well plates (at a density of 5 × 10^5 cells/well) in triplicate. An Annexin V PE Apop Dtec Kit (BD Pharmingen, San Diego, CA) was used to analyze the apoptosis rate by flow cytometry according to the protocols of the manufacturer.

RNA Sequencing

Next Generation Sequencing (NGS) RNA-Seq was performed to analyze the transcriptomes of the tested cells. Total RNA was isolated using the QIAaamp RNA Blood Mini Kit (QIAGEN, Düsseldorf, Germany) according to the instructions of the manufacturer. RNA samples were analyzed by the BGISEQ-
Public Datasets and Bioinformatics Analysis

A cohort of 159 MDS patients and 17 healthy controls downloaded from the Gene Expression Omnibus (GEO) database (GSE58831) (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE58831) was used to identify the mRNA expression of LEP in MDS.

Statistics

Statistical analysis was accomplished using the SPSS 22.0 and GraphPad Prism 5.0 software packages. Comparisons of continuous variables were done by the Independent T/Mann–Whitney’s U test, whereas comparisons of categorical variables were analyzed using Pearson Chi-square/Fisher exact tests. A Spearman correlation test was performed to analyze the correlation between the results obtained using RQ-MSP and targeted bisulfite sequencing in the detection of LEP methylation. Kaplan–Meier analysis (Log-rank test) and Cox regression analysis (univariate and multivariate proportional hazard regression) were performed to evaluate the prognostic impact of LEP methylation on leukemia-free survival (LFS) and overall survival (OS) of MDS patients. The statistical results were considered significantly different if two-sided P-values were less than 0.05.

RESULTS

Genome-Wide Methylation Analysis in MDS Patients

To identify epigenetic alterations occurring in MDS, we performed RRBS in five newly diagnosed MDS patients and four healthy donors. The sequencing data of four newly diagnosed MDS patients and four healthy donors were submitted to the NCBI SRA databases (PRJNA670308) previously. Sequencing data of one remaining MDS patients are available from the corresponding author on reasonable request. The details of the sequencing data were described in our previous study (10). However, the mean global methylation in five de novo MDS patients (51.2, 38.9, 43.3, 42.3, and 44.8%) showed no significant difference compared with four controls (51.0, 49.3, 49.4, and 45.4%) (P = 0.106, Figure S1).

Next, we used MspI fragments (40–220 bp) rather than individual CpG sites or a tiled window approach as the basic analysis units as in our previous report (10). The fragments that were statistically significant (P <0.05, Q <0.05, and also had >25% mean methylation difference) were considered differentially methylated fragments (DMFs). A total of 1,459 DMFs, including 759 hypermethylated and 700 hypomethylated fragments, were identified between MDS patients and controls (Figure 1A and Supplementary Table S2). Moreover, both CpG islands (CGI), promoter (± 2,000 bp from transcription start site), and gene body were also used as the units of analysis, respectively. A total of 922 differentially methylated gene bodies (146 hypermethylated and 776 hypomethylated), 78 differentially methylated promoters (48 hypermethylated and 30 hypomethylated), and 87 differentially methylated CGI (65 hypermethylated and 22 hypomethylated) were identified between MDS patients and controls (Figures 1B–D and Supplementary Table S2).

Identification and Validation of Candidate DMFs in MDS Patients

As is well known, the promoter CpG site hypermethylation is associated with gene silencing, and plays a crucial role in cancer development. Here, to further identify the candidate genes involved in MDS, we first annotated 1,459 DMFs as differentially methylated genes (DMGs), and then selected the promoter-associated DMGs, and finally selected the candidate genes for validation (Figure 3A). Following the procedure, we obtained 184 DMGs (128 hypermethylated genes and 56 hypomethylated genes), which may play a crucial role in MDS pathogenesis (Figure 3B).

The targeted bisulfite sequencing methodology MethylTarget (GENESKY, Shanghai, China) was performed in an additional cohort of 36 MDS and 25 controls to further validate the six candidate genes (DLEU7, FOXR1, LEP, PANX2, RARRES2, and REC8), which may have potential biological functions in cancers predicted by Coremine analysis (http://www.coremine.com/medical/#search). Besides RARRES2 and REC8, the methylation level of DLEU7, FOXR1, LEP, and PANX2 in MDS patients was markedly increased compared with controls (Figure 3).

Further Confirmation of LEP Methylation in a Larger Cohort of MDS Patients

Hypermethylation of LEP was further confirmed in a larger cohort of 105 MDS patients and 46 controls by RQ-MSP developed previously (15). The results obtained by RQ-MSP and targeted bisulfite sequencing in the detection of LEP methylation among MDS patients were highly correlated with each other (R = 0.533, P = 0.001, Figure 4A). Moreover, the LEP methylation level in MDS patients was markedly higher than that in controls, as expected (P = 0.044, Figure 4B). Because of the limited mRNA samples in our MDS cohort, we used the public GEO data to identify the expression of LEP in MDS patients. As shown in Figure 4C, LEP mRNA expression was significantly reduced in MDS patients compared with normal controls (P <0.001).

To analyze the clinical relevance of LEP methylation in MDS, we divided the patients into two groups (LEP hypermethylated and non-hypomethylated) based on the cutoff value of 0.569 (set
as “mean + 2SD” in controls). As shown in Table 1, there were no marked differences when comparing the two groups with regard to sex, age, white blood cells, hemoglobin, platelets, and WHO and IPSS classifications (all $P >0.05$, Table 1). However, LEP hypermethylation tended to be associated with lower BM blasts ($P = 0.052$, Table 1) and was significantly correlated with U2AF1 mutation ($P = 0.016$, Table 1).

**Prognostic Effect of LEP Methylation in MDS Patients**

The prognostic effect of LEP methylation on OS and LFS was further analyzed in MDS patients. Interestingly, Kaplan–Meier analysis demonstrated that patients with LEP hypermethylation exhibited markedly longer OS and LFS times than patients with LEP non-hypermethylation ($P = 0.037$ and 0.038, respectively, Figures 4D, E). However, LEP hypermethylation was not a prognostic biomarker independently affecting OS and LFS in MDS patients ($P = 0.540$, Table 2) by Cox regression analysis.

**Biological Functions of LEP in MDS Cells**

To determine the potential role of LEP during MDS pathogenesis, we carried out gain-of-function experiments in the MDS cell line SKM-1 in vitro. Interestingly, SKM-1 cells treated with human recombinant leptin exhibited a higher growth rate and a lower apoptosis rate than those treated without human recombinant leptin (Figures 5A, B). The biological functions of leptin seemed to be contrary to the hypermethylation pattern in MDS.

We next performed mRNA-sequencing of SKM-1 cells before and after treating with human recombinant leptin to get a better understanding of the biological network in MDS affected by aberrant leptin expression. A total of 31 differential expressed genes (DEGs) including 20 upregulated and 11 downregulated were identified between two groups ([log2 FC] >1, FDR <0.05 and $P <0.05$) (Figures 5C, D). Moreover, the analysis of Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment revealed that these genes were involved in...
cell growth and death (Figures 5E, F) as well as Toll-like receptor and NF-kappa B signaling pathways (Figures 5G, H).

DISCUSSION

Epigenetic alterations, especially aberrant DNA methylations, play a vital role in MDS progression. Previously, Jiang et al. reported that aberrant DNA methylation was a dominant mechanism in MDS progression to AML (17). Similarly, our previous study also confirmed that whole-genome DNA hypermethylation was a frequent event during MDS progression (10). Interestingly, our study did not observe genome-wide DNA hypermethylation in the MDS stage compared with controls, while the whole-genome methylation level in MDS seems to be lower than in controls. Deep analysis of differentially methylated CpG sites revealed that the gene body showed more hypomethylated changes than hypermethylation alterations, whereas CpG islands and promoters showed the opposite phenomena. As is well known, promoter CpG island hypermethylation is associated with TSG silencing during oncogenesis. Gene body hypomethylation harbors extensive intragenic transcriptional activity and is prone to cancer-associated dysregulation (18).

In this study, we further identified several DNA methylation-associated genes as novel epigenetic alterations in MDS. Although aberrant methylation of these candidate genes, including FOXO3 and TERT, has previously been reported in MDS (19, 20), most others remain to be elucidated. Interestingly, some of these genes, such as LEP, ALOX12, RARRES2, DLEU7, and FOXR1, have been reported to be associated with other hematologic malignancies, including AML, chronic lymphocytic leukemia (CLL), and B-cell lymphoma (15, 21–24). Moreover, methylations in RARRES2, REC8, WNK2, and PANX2 have been demonstrated with potential roles in solid cancers (25–28). Therefore, we focused on these genes which were clearly associated with other hematologic malignancies or solid tumors for further validation and revealed that methylations of DLEU7, FOXR1, LEP, and PANX2 were frequent events in MDS. However, due to the limited samples detected in targeted bisulfite sequencing, further investigations are needed to confirm our results.

Since our research group has determined the obesity-related gene LEP methylation in AML, the LEP methylation pattern in MDS patients has come to our attention. Based on our previous study, LEP hypermethylation was not present during MDS progression (10). Moreover, hypermethylation of LEP was a frequent event in primary AML and MDS-derived AML, but they showed no significant difference between primary AML and MDS-derived AML (15). In this study, we further confirmed the LEP hypermethylation phenomenon in a larger cohort of MDS patients, but it was not an independent prognostic biomarker.
affecting LFS in MDS. These results indicated that LEP methylation may only play a vital role in the initiation pathogenesis of MDS rather than disease progression.

As is well known, promoter CpG island hypermethylation is associated with gene silencing. It is a pity that we did not evaluate LEP methylation with LEP expression in MDS, which was mainly caused by the limited samples. Nevertheless, our previous study has proved the significantly negative association between LEP methylation and LEP expression in AML (15). Moreover, we further identified the reduced expression of LEP in MDS patients by using public GEO datasets. Although methylation of LEP and its direct role of leptin protein were less reported in MDS, leptin expression in MDS has been increasingly investigated. Dalamaga et al. revealed that low leptin concentrations were observed in low-risk MDS patients with normal or good prognostic karyotype (29). Subsequently, the same research group further reported that free leptin was associated with a higher risk of MDS, particularly among overweight/obese individuals (30). In our study, although the significant association of LEP hypermethylation with IPSS risks was not observed, we showed that LEP hypermethylation, as a frequent event in MDS, was associated with longer OS and LFS. Similarly, Kraakman et al. demonstrated that leptin-deficient obesity prolongs survival in a murine model of MDS (31). The functional studies showed pro-proliferative and anti-apoptotic effects of leptin in MDS in accordance with a previous study (32). These “conflicting”

**FIGURE 3** | Identification and validation of differentially methylated genes in MDS. (A) The flowchart of the differentially methylated genes screening. The fragments that passed statistical significance (P < 0.05, Q < 0.05 and also had > 25% mean methylation difference) were considered as differentially methylated fragments/gene. (B) Heatmaps summarizing differentially methylated fragments/gene in MDS. (C) The methylation level of the candidate genes in additional samples of de novo MDS (n = 36) and controls (n = 25) analyzed by targeted bisulfite sequencing. P-values were calculated using the Mann–Whitney U-test. NS, no significance; *P < 0.05; **P < 0.01; ***P < 0.001.
Further confirmation of LEP methylation in MDS patients together with its prognostic value. (A) The correlation of LEP methylation results between the targeted bisulfite sequencing and RQ-MSP. The correlation was analyzed by Spearman correlation test. (B) The methylation level of the LEP in larger samples of controls (n = 46) and de novo MDS (n = 105) and AML patients (n = 170) analyzed by RQ-MSP. P-values were calculated using the Mann-Whitney U-test. (C) Leptin expression in MDS patients from public GEO datasets. (D, E) The impact of LEP methylation on overall survival and leukemia-free survival of MDS patients. Survival was analyzed through Kaplan-Meier analysis using Log-rank test.

| Patient’s features | Non-hypermethylated (n = 68) | Hypermethylated (n = 37) | P-value |
|--------------------|-------------------------------|--------------------------|---------|
| Sex (male/female)  | 39/29                         | 19/18                    | 0.682   |
| Median age, years (range) | 62.5 (14–86)                 | 52 (20–86)               | 0.123   |
| Median WBC, ×10^9/L (range) | 2.7 (0.7–26.6)             | 2.85 (0.6–82.4)          | 0.706   |
| Median hemoglobin, g/L (range) | 65.5 (29–140)               | 59 (26–118)              | 0.149   |
| Median platelets, ×10^12/L (range) | 65 (1–1,176)               | 43.5 (0–505)             | 0.110   |
| Median BM blasts, % (range) | 6 (0–18)                    | 2 (0–19)                 | 0.052   |
| WHO classifications | RCUD/RARS 9                  | 5                        | 0.788   |
|                   | RCM/RCMD-RS 24               | 16                       |         |
|                   | RAEB-1 11                   | 8                        |         |
|                   | RAEB-2 21                   | 7                        |         |
|                   | MDS with isolated del(5q) 2 | 1                        |         |
|                   | MDS-U 1                     | 0                        |         |
| IPSS scores       | Low 7                       | 6                        | 0.294   |
|                   | Int-1 33                    | 21                       |         |
|                   | Int-2 13                    | 8                        |         |
|                   | High 8                      | 1                        |         |
|                   | No data 7                   | 1                        |         |
| Gene mutations    | CEBPA (+/-) 1/61            | 2/30                     | 0.266   |
|                   | IDH1/2 (+/-) 3/59           | 1/31                     | 1.000   |
|                   | DNMT3A (+/-) 2/60           | 1/31                     | 1.000   |
|                   | U2AF1 (+/-) 1/61            | 5/27                     | 0.016   |
|                   | SRSF2 (+/-) 2/60            | 1/31                     | 1.000   |
|                   | SF3B1 (+/-) 3/59            | 3/29                     | 0.406   |
|                   | SETBP1 (+/-) 0/62           | 1/31                     | 0.340   |

MDS, myelodysplastic syndromes; WBC, white blood cells; BM, bone marrow; WHO, World Health Organization; IPSS, International Prognostic Scoring System.
results might suggest that LEP hypermethylation-mediated leptin expression was an early event in MDS initiation but did not act as a tumor suppressor, and that they played a protective role during MDS progression, leading to favorable prognosis. These findings provided a theoretical basis and opened new insights for developing leptin-related targeted therapy in MDS.

Collectively, our findings indicated that whole-genome DNA methylation analysis identified novel epigenetic alterations such as DLEU7, FOXR1, LEP, and PANX2 methylations as frequent events in MDS. Moreover, LEP might play a role in MDS pathogenesis, and LEP hypermethylation was associated with longer survival but not as an independent prognostic biomarker in MDS.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and...
accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/bioproject/PRJNA670308.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Ethics Committee of the Affiliated People’s Hospital of Jiangsu University. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

J-dZ and JQ conceived and designed the experiments. T-jZ and J-dZ performed the experiments. Z-jX analyzed the data and provided bioinformatics analysis. YG and X-IZ collected the clinical data. J-cM, X-mW and JL provided the technical and financial supports. J-dZ wrote the paper. All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

FUNDING

The work was supported by the National Natural Science Foundation of China (81900166, 81900163, 81970118), the Zhenjiang Clinical Research Center of Hematology (SS2018009), the Social Development Foundation of Zhenjiang (SH2020055, SH2021052), the Medical Field of Zhenjiang “Jin Shan Ying Cai” Project, Medical Education Collaborative Innovation Fund of Jiangsu University (JDY2022011), and the Scientific Research Foundation of Affiliated People’s Hospital of Jiangsu University for PhD (KFB202002, KFB202202).

ACKNOWLEDGMENTS

We are very grateful for the clinical samples provided by Professor Su-ning Chen from The First Affiliated Hospital of Soochow University and also grateful for the technical support by Genesky Biotechnologies Inc., Shanghai.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2022.897898/full#supplementary-material

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