Abnormal DNA methylation patterns in patients with infection-caused leukocytopenia based on methylation microarrays

CHAO WU*, XIRENNAYI MUHATAER*, WENYI WANG, MINGQIN DENG, RONG JIN, ZHICHUANG LIAN, DAN LUO, YAFANG LI and XIAOHONG YANG

Department of Respiratory and Critical Care Medicine, People's Hospital of Xinjiang Uygur Autonomous Region, Urumqi, Xinjiang Uygur Autonomous Region 830001, P.R. China

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Abstract. The present study aimed to investigate the association between gene methylation and leukocytopenia from the perspective of gene regulation. A total of 30 patients confirmed as having post-infection leukocytopenia at People's Hospital of Xinjiang Uygur Autonomous Region between January 2016 and June 2017 were successively recruited as the leukocytopenia group; 30 patients with post-infection leukocytosis were enrolled as the leukocytosis group. In addition, 30 healthy volunteers who received a health examination at the hospital during the same period were included as the normal control group. In each group, four individuals were randomly selected for whole genome methylation screening. After selection of key methylation sites, the remaining samples in each group were used for verification using matrix-assisted laser desorption/ionization-time of flight mass spectrometry. The levels of serum complement factors C3 and C5 in the leukocytopenia group were significantly lower than those in the other two groups (P<0.05). According to whole-genome DNA methylation detection, 66 and 27 methylation loci may be associated with leukocytopenia and leukocytosis, respectively. Most of these abnormal loci are located on chromosomes 2, 6, 7, 1, 17 and 11. The rates of WW domain containing E3 ubiquitin protein ligase 2 gene methylation at cytosine-phosphate-guanine (CpG)_1, CpG_5/6 and CpG_7 in the leukocytopenia group were higher than in the other two groups (P<0.05); the rate of AKT2 CpG_1 methylation was higher in the leukocytopenia group than in the other two groups (P<0.05); the rate of calcium-binding atopy-related autoantigen 1 gene CpG_2 methylation was higher in the leukocytosis group than in the normal control group (P<0.05); and the rate of NADPH oxidase 5 gene CpG_3 methylation was higher in the leukocytosis group than in the normal control group (P<0.05). Chemotactic factor secretion and cell migration abnormalities, ubiquitination modification disorders and reduced oxidative burst may participate in infection-complicated leukocytopenia. The results of this study shed new light on the molecular biological mechanisms of infection-complicated leukocytopenia and provide novel avenues for diagnosis and treatment.

Introduction

Although most patients with infection exhibit leukocytosis, which contributes to rapid inherent immunological defense, a certain portion of these patients experience leukocytopenia (1). In patients with leukocytopenia, pathogens are more likely to escape from immunocytes, increasing the severity of infection symptoms and complicating infection control, thereby prolonging the course of anti-infective treatment (2). Leukocytopenia can be caused by a variety of pathogens, such as bacteria and viruses; however, the detailed mechanisms underlying leukocytopenia post-infection remain unclear (3,4). Numerous studies have demonstrated that gene methylation plays an important role in a variety of pathophysiological processes, and exploration of pathomechanisms from the perspective of gene regulation may provide novel approaches for the diagnosis and treatment of diseases (5,6).

DNA methylation refers to the process via which methyltransferases modify 5’ terminal promoter region bases and the pentacoordinate carbon of cytosine-phosphate-guanines (CpGs) in the first exon to form 5-methylcytosine (7). A CpG island refers to a region with a CpG sequence density that is 10-20 times higher than the average density of the sequence, with >50% G+C content in a region >200 bases; ~50% of human genes contain CpG islands, for ~29,000 total CpG islands, which are normally located in the promoter region of the gene (8). In normal cells, CpG islands exist in a non-methylated state; however, high levels of methylation at CpG islands in housekeeping genes and tissue-specific genes are closely associated with inactivation of protein function and the
development of cancers and genetic diseases (9). Furthermore, methylation of CpG islands in some non-promoter regions, including intragenic, intergenic and downstream regions, may also cause pathological changes at the molecular level (10).

Previous studies on methylation have primarily focused on tumors (5,9); to the best of our knowledge, the role of DNA methylation in leukocytopenia after infection has not been reported. Overall, studying the methylation of single genes is laborious and inefficient, and it fails to reflect the epigenetic variations in patients with post-infection leukocytopenia. Furthermore, a large amount of work would be required to verify individual candidate methylation-regulated genes to determine whether they cause molecular changes in patients with post-infection leukocytopenia. Gene chip technology, also known as microarray technology, is a systematic, complete analytical method specific for DNA methylation (11). Advantages of this method include easy operation and high throughput (11). Indeed, detection of methylation patterns based on this technique has been reported in a number of studies (11,12).

The present study aimed to apply gene chip technology to investigate the association of DNA methylation with post-infection leukocytopenia. To this end, baseline data, inflammation indices and chemotactic factors of patients with post-infection leukocytopenia were analyzed, and the methylation status of the entire genome was analyzed using 450K methylation chips to establish gene methylation patterns associated with these patients. Key methylated gene sites were then screened and verified.

Materials and methods

Study subjects. A total of 30 patients who were diagnosed with infection complicated by leukocytopenia (<3.5x10^9/l leukocytes; mean age, 52.34±12.74 years; age range, 39-66 years) who underwent treatment at People's Hospital of Xinjiang Uygur Autonomous Region (Urumqi, China), were successively recruited between January 2016 and June 2017. Patients with leukocytosis (>3.5x10^9/l leukocytes; n=30; mean age, 53.62±13.08 years; age range, 39-66 years) were also enrolled. The control group was comprised of 30 healthy volunteers (mean age, 55.76±11.84 years; age range, 43-67 years) who underwent a health examination at the same hospital during the same period. The clinical data of the participants were analyzed. From each group, four individuals were randomly selected for gene-related examination, and the four with leukocytopenia (designated as A1-A4; n=4), four with leukocytosis (C1-C4; n=4) and four healthy volunteers (D1-D4; n=4) constituted the leukocytopenia, leukocytosis and normal control groups, respectively. Patients with infection due to unknown pathogens or with non-respiratory infection were excluded from the leukocytopenia group; those with leukocytopenia due to chemoradiation, hematological disease or sulfonamide and analgesic medication were excluded from the leukocytopenia group; and those with leukocytopenia due to chronic bronchitis and bronchial asthma) were excluded from the leukocytopenia group; those with leukocytopenia due to chemoradiation, hematological disease or sulfonamide and analgesic medication were excluded from the leukocytopenia group; and those with diseases that influence C3 and C5 levels (such as rheumatoid arthritis, idiopathic interstitial pneumonia, chronic bronchitis and bronchial asthma) were excluded from all groups. The 30 patients with leukocytopenia were further divided into slight (<1.5x10^9/l; n=9), moderate (2.5-1.5x10^9/l; n=15) and severe (1.5x10^9/l; n=6) subgroups, according to the degree of decrease in white blood cell count (13).

This study was conducted in accordance with the Declaration of Helsinki. Approval was obtained from the Ethics Committee of People's Hospital of Xinjiang Uygur Autonomous Region (policy no. 2016056). Written informed consent was obtained from each participant.

Data collection. General data, including sex, age and infection site, were collected for each patient. Clinical data, laboratory examination outcomes with regard to blood routine examination, C3 and C5 complement levels, procalcitonin (PCT) and interleukin (IL)-6 were also collected. In total, 4 ml of peripheral venous blood was extracted from each participant after fasting and combined with 0.109 mol/l sodium citrate anticoagulant agent. The samples were immediately stored at -196°C after collection.

C3 and C5 detection. C3 and C5 serum levels were detected using ELISA. Kits for human serum C3 and C5 detection were purchased from Shanghai Fusheng Industrial Co., Ltd. (cat. no. YM-YM10444), the procedures were performed in strict accordance with the instructions of the manufacturer.

Locations of CpG sites with differential methylation. Sites with differential methylation were classified into transcription start sites (TSS)200, 5'untranslated region (UTR), body, TSS1500, 3' UTR and 1st exon categories according to the physical locations of the detected sites, as well as the related annotation information were provided by the University of California Santa Cruz database (14).

Whole-genome methylation detection

DNA preparation and quality examination. Genomic DNA was extracted from blood samples using a Wizard® Genomic DNA Purification kit (Promega Corporation), quantified by spectrophotometry and adjusted to the standard concentration of 50 ng/µl (20 µl in total). The sample was subjected to 0.8% agarose gel electrophoresis. Methylation experiments were performed using samples with a distinct major electrophoresis band with a length no shorter than 10 kb and no noticeable degradation, with a total sample quantity of >5 µg.

Sulfite transformation. Sulfite transformation was performed using an EZ DNA methylation kit (Zymo Research Corp.) according to the manufacturer's protocols.

NaOH denaturation and whole-genome amplification. NaOH (0.1%) was used to dissociate double-stranded DNA into single-stranded DNA. Whole-genome amplification was performed using a PCR amplification kit (cat. no. KG203; Beijing Hongyue Biotech Co., Ltd.). The amplification conditions were as follows: 94°C for 12 min; 10 cycles of 93°C for 50 sec, 62°C for 48 sec (-0.5°C/cycle) and 72°C for 1 min; 35 cycles of 94°C for 45 sec, 56°C for 48 sec and 72°C for 1 min; 72°C for 3 min; and hold at 4°C. After amplification, 2% agarose gel electrophoresis was performed for product verification.

DNA segmentation, precipitation and resuspension. PCR products were fragmented using the random endonuclease method. Isopropanol was applied for centrifugal precipitation of the fragments at 4°C (2x10^4 x g for 15 min). The precipitated
sample was placed at room temperature, dried and dissolved in hybridization buffer.

**DNA/chip hybridization.** Prepared chips were used for DNA hybridization. The capture probe with 50-mer-long beads (7 fmol/µl) was combined with the digested genomic DNA (gDNA) products; annealing was carried out in a hybridization oven for 8-10 h.

**Chip washing and single-base extension and staining.** The wash rack was submerged in the wash dish containing 200 ml PB1 (Shanghai Tichem Chemical Co., Ltd.). The Hyb Chamber inserts were removed from the Hyb Chambers. BeadChips (CapitalBio Corporation) were removed from the Hyb Chamber inserts one at a time, and the cover seal was removed from each BeadChip. Then, the BeadChip was slide into the wash rack, one at a time, ensuring that the BeadChip was complete. These procedures were repeated until all BeadChips (a maximum of 8) were transferred to the wash rack submerged in PB1. Once all BeadChips were in the wash rack, the wash rack was moved up and down for 1 min, the surface of the PB1 was broken with gentle, slow agitation. The wash rack was transferred to the other wash dish containing clean PB1 (ensuring all BeadChips were complete) and the wash rack was moved up and down for 1 min. The surface of the PB1 was broken with gentle, slow agitation. The DNA sequence detected by the probe was used as the template, and nucleotide substrates were labeled with dinitrophenol and biotin using the random primer method (a/T and C/G, respectively) (15). Unqualified DNA was removed, followed by single base extension (the condition for base extension was that gDNA hybridization did not occur). Detectable fluorophores (Fluorescein-5-maleimide) were applied to the chip, and a/T and C/G were used for labeling different fluorocent dyes to distinguish different methylation states. The chip was coated with Xc4 reagent to prolong its lifetime and the extension time of the signals. The chip was then placed in a vacuum dehydration unit for 1 h.

**Chip scanning and data reading.** The fluorophores were excited with a laser to obtain fluorescence. Corresponding ‘manifest’ files were downloaded (https://support.illumina.com/downloads/infinium-methylationepic-v1-0-product-files.html) to generate raw data. The data were then analyzed using GenomeStudio software (version 1.1.0; Illumina, Inc.) to obtain the methylation P-value of each gene and each site of the gene (16,17).

**Key site screening.** To obtain the genes associated with leukocytopenia, the scope of key gene screening was narrowed by intersection analysis (18). Based on P-values and the functions of related genes according to GenBank (https://www.ncbi.nlm.nih.gov; accessed on February 18, 2018), AKT2, NADPH oxidase 5 (NOX5), WW domain containing E3 ubiquitin protein ligase2 (WWP2) and calcium-binding atopy-related autoantigen 1 (CBARA1) were ultimately selected for verification.

**Key gene verification**

**DNA preparation.** DNA was extracted from blood samples using a Wizard Genomic DNA Purification kit (Promega Corporation). Quantitation using spectrophotometry and 1% agarose gel electrophoresis was performed. In general, the length of gDNA obtained was no shorter than 20 kb. DNA was adjusted to a concentration of 50 ng/µl and transferred to a 384-well plate for storage at -20°C.

**Primer design.** Primers were designed by EpiDesigner (Sequenom; http://www.epidesigner.com) to correspond to the methylation sites of AKT2, NOX5, WWP2 and CBARA1. The normal lengths of the amplified fragments ranged from 200-600 bp. To the 5’ terminal of the forward primer, 10-mer tag was applied to balance PCR conditions, and a T7 promoter sequence was applied to the 5’ terminal of the reverse primer for subsequent transcription in vitro. Syntheses were performed by CapitalBio Technology Co., Ltd. The upstream and downstream primers of AKT2 were 5’-TTT ATGTAAAGAGGAAATATGTGGGAAA-3’ and 5’-CCA AAA AACAACAAATTCTCACC-3’, respectively. After modification, the upstream and downstream primers were 5’-aggaggagagTTTATGTAAAGAGGAAATATGTGGGAAA-3’ and 5’-cagtaatacgactcatactagggagaagctCCA-3’ respectively. The upstream, downstream, modified upstream and modified downstream primers of NOX5 were 5’-TGGGGATTTTATTTTAAGTTGTTGGGT-3’, 5’-AATACTACACTCCACTGACCTCCACT-5’, 5’-aggaggagagTTTGGGATTTTTTTTATTGAGTTG-3’ and 5’-cagtaatacgactcatactagggagaagctAATCATT-3’, respectively. The upstream, downstream, modified upstream and modified downstream primers of WWP2 were 5’-TGGGGGATTATTTTTAAGTTGTTGGGT-3’, 5’-AAAACTGAAAACACATTACCACCCCAAT-3’, 5’-aggaggagagTTTGGGATTTTTTTTATTGAGTTG-3’ and 5’-cagtaatacgactcatactagggagaagctAATCATT-3’, respectively and those of CBARA1 were 5’-GGTATGTGTTTTTTTTGGAGAGTT-3’, 5’-ATCACAAGAAATACACCAACAAAT-3’, 5’-aggaggagagGGTAGTGTGTATTGATTGGAGATTTG-3’ and 5’-cagtaatacgactcatactagggagaagctATCACAAGAAATACACCAACAAAT-3’, respectively.

**PCR amplification.** Amplification was performed using a PCR amplification kit (Sequenom). The amplification conditions consisted of 94°C for 12 min, 10 cycles at 93°C for 50 sec, 62°C for 48 sec (-0.5°C/cycle) and 72°C for 1 min, 35 cycles at 94°C for 45 sec, 56°C for 48 sec and 72°C for 1 min, 72°C for 3 min and a final hold at 4°C. Agarose gel (2%) electrophoresis was performed for PCR product verification, with ethidium bromide as the fluorescence dye. RT fluorescent quantitation was performed for PCR products.

**Shrimp alkaline phosphatase (SAP).** SAP treatment was performed in accordance with the instructions of the MassCLEAVE™ reagent kit (Sequenom).

**Chip sample application and mass spectrometric detection.** Purified products were applied to a 384-formatted SpectroCHIP with the MassARRAY(Nanodispenser RS1000 arrayer (Sequenom). The prepared chip was detected using a MassARRAY compact system (Sequenom). Matrix-assisted laser desorption/ionization-time of flight mass spectrometry was performed in accordance with the protocols of the
Infinium HD Assay Methylation Protocol Guide 2011 (19), and data were analyzed using epiTYPER v1.05 (Sequenom).

**Statistical analysis**

**Genomic methylation data.** Raw data were processed by GenomeStudio software to obtain the original signal value and detection P-value of each site (15,16). Site quality control and individual quality control were performed. Color bias adjustment and quantile normalization were performed for the quality-controlled data using Lumi 2.22.1 included in the R software 4.0.0 package (20,21). For probe-type biases, the methylation levels ($\beta$ values) obtained after the preceding adjustment step were further adjusted with BMiQ v1.3 ($\beta$-mixture quantile normalization). The adjusted $\beta$ values were subjected to differential methylation analysis using iMa 3.1.2 of the R software package (22), and the analysis method was based on the limma algorithm (23) of empirical Bayes Statistics. For multiple hypothesis testing, the P-value adjusted by the false discovery rate (FDR) was calculated using the Benjamini-Hochberg method (24). The selection criterion for differential sites was an adjusted P-value (FDR value) $\leq 0.05$ (if the adjusted P-values of all sites were $>0.05$ or when the adjusted P-values of few sites were $\leq 0.05$, a P-value before adjustment of $\leq 0.05$ was used as the criterion). Cluster analysis was performed for the samples using gplots 2.13.0 of the R software package (22), and the analysis method was based on the limma algorithm (23) of Empirical Bayes Statistics. For multiple hypothesis testing, the P-value adjusted by the false discovery rate (FDR) was calculated using the Benjamini-Hochberg method (24).

**Other data.** Experimental data were processed using SPSS 22.0 software (IBM Corp.). For categorical data, $\chi^2$ test was used for group comparisons. For data regarding methylation levels and serum testing, one-way ANOVA was used for comparison among groups. When measurement data were abnormally distributed or exhibited heterogeneity of variance [PCT, C-reactive protein (CRP), acute physiology and chronic health evaluation II (APACHE II) score, C3], the nonparametric Kruskal-Wallis one-way ANOVA was applied for comparison of methylation levels among groups. The significance level was adjusted using the Bonferroni method, and an adjusted P$<0.05$ was considered significant (all stated P-values are adjusted). GraphPad Prism 6 (GraphPad Software, Inc.) was used for plotting.

**Results**

**General data**

**Clinical data.** Clinical data for patients with post-infection leukocytopenia or post-infection leukocytosis, and for normal controls were analyzed (Table 1). The confirmation of infection was based on etiological outcomes or an increase in PCT levels. The three groups did not show significant differences in age ($F=1.026$, P=0.349) or sex ratio ($\chi^2=5.217$, P=0.084). Similarly, the leukocytopenia and leukocytosis groups did not show significant differences in infection sites ($\chi^2=4.692$, P=0.074) or infectious pathogen type ($\chi^2=7.631$, P=0.062).

**Diagnostic values of inflammation indices for post-infection leukocytopenia and leukocytosis.** The leukocytopenia group had significantly higher levels of PCT and IL-6 than the leukocytosis group (P=0.015 and P=0.034, respectively); however,
no significant difference in CRP levels was observed between these two groups (P>0.05; Table II).

**Association of inflammation indices with the total white blood cell count in leukocytopenia.** The 30 patients with leukocytopenia were further divided according to the degree of decrease. The highest serum PCT level was observed in the severe group, followed by the moderate group and the slight group (P<0.05). No significant differences in IL-6 and CRP were observed among the three groups. The highest APACHE II score was observed in the severe group, followed by the moderate group and the slight group (P<0.05). The results are summarized in Table III.

**Changes in serum C3 and C5 levels.** C3 and C5 levels in the leukocytopenia, leukocytosis and normal control groups were detected by ELISA. The leukocytopenia group showed significantly lower levels of C3 and C5 than the leukocytosis and the normal control groups (all P<0.05; Table IV).

**Methylation levels**

Cpg sites and genes with differential methylation. Methylation levels were detected at 473,270 sites across the leukocytopenia, leukocytosis and normal control groups. Pairwise comparisons of methylation rates were performed among the three groups, and the results are shown in Fig. 1. The comparison between the leukocytopenia group and the leukocytosis group demonstrated significantly more hyper- methylation sites compared with hypo-methylation sites, and the comparison between the leukocytosis group and the normal control group found less hyper- methylation sites compared with hypo-methylation sites. However, there were no significant difference between the number of hyper- methylation sites and that of hypo-methylation sites when the leukocytopenia group was compared with the normal control group.

Locations of Cpg sites with differential methylation. Methylation of the TSS, 5’ UTR and 1st exon inhibits gene expression, whereas that of the body and 3’ UTR promotes expression. The results are shown in Fig. 2. A total of 24,230 differential methylation sites were detected between the leukocytopenia group and the leukocytosis group: 11,796 sites were related to the genes whose methylation inhibits gene expression and 9,416 were related to the genes whose methylation promotes gene expression, with the rest undetected in genes. There were 26,892 differential methylation sites detected in total between the leukocytosis group and
the normal control group: 13,390 sites were related to the genes whose methylation inhibits gene expression and 10,899 were related to the genes whose methylation promotes gene expression, with the rest undetected in genes. Moreover,
comparison between the leukocytopenia group and the normal control group identified 50,117 differential methylation sites: 22,459 sites were related to the genes whose methylation inhibits gene expression and 20,210 were related to the genes whose methylation promotes gene expression, with the rest undetected in genes.

Chromosome distribution of sites with differential methylation. The chromosome distribution of sites with differential CpG methylation was also analyzed. Sites with differential methylation between the leukocytopenia and leukocytosis groups were primarily located on chromosomes 1, 2, 6, and 7. In addition, sites with differential methylation between the leukocytosis and normal control groups were primarily located on chromosomes 2, 6, 7 and 17, and those between the leukocytopenia and normal control groups were primarily located on chromosomes 1, 2, 6 and 11 (Table V).

GO enrichment analysis. GO enrichment categories include biochemical processes, molecular function and cellular composition. The comparison between the leukocytopenia group and the leukocytosis group demonstrated that the three biochemical processes with the P-values with the greatest significance are nervous system development, system development and multicellular organismal development; the three molecular functions with the P-values with the greatest significance are protein binding, binding and organic cyclic compound binding; and the three cellular compositions with the P-values with the greatest significance are cell projections, neuron projections and cell junctions (Fig. 3A). Comparing the leukocytosis and normal control groups, the three biochemical processes with the P-values with the greatest significance are cellular component organization or biogenesis, cellular component organization and positive regulation of biological process; the three molecular functions with the P-values with the greatest significance are protein binding, binding and organic cyclic compound binding; and the three cellular compositions with the P-values with the greatest significance are intracellular, intracellular part and cytoplasm (Fig. 3B). The comparison between the leukocytopenia group and the normal control group showed that the three biochemical processes with the P-values with the greatest significance are system development, multicellular organismal development and single-organism developmental process; the three molecular functions with the P-values with the greatest significance are cation channel activity, ion channel activity and passive transmembrane transporter activity; and the three cellular compositions with the P-values with the greatest significance are cell projections, plasma membrane part and extracellular matrix (Fig. 3C).

Pathway analysis
Leukocytopenia vs. leukocytosis. Signaling pathways with the most significant P-values are associated with Rap1 signaling pathway, cholinergic synapse, circadian entrainment and morphine addiction (Fig. 4). Further disease enrichment with the R software identified that diseases with the most significant P-values were cancer, depression, behavior disease and intestinal disease (Fig. S1).

| Chromosome no. | Leukocytopenia vs. leukocytosis (n, %) | Leukocytosis vs. control (n, %) | Leukocytopenia vs. control (n, %) |
|---------------|--------------------------------------|-------------------------------|-----------------------------------|
| 1             | 1,561 (6.44)                         | 1,809 (6.73)                 | 1,809 (6.73)                      |
| 2             | 2,564 (10.58)                        | 2,791 (10.38)                | 3,848 (7.68)                      |
| 3             | 1,390 (5.73)                         | 1,365 (5.08)                 | 2,690 (5.37)                      |
| 4             | 1,205 (5.10)                         | 1,165 (4.33)                 | 2,370 (4.73)                      |
| 5             | 1,296 (5.35)                         | 1,355 (5.04)                 | 2,732 (5.45)                      |
| 6             | 1,990 (8.21)                         | 2,235 (8.31)                 | 3,810 (7.60)                      |
| 7             | 1,558 (6.43)                         | 1,643 (6.11)                 | 3,163 (6.31)                      |
| 8             | 1,080 (4.46)                         | 1,179 (4.38)                 | 2,389 (4.77)                      |
| 9             | 0 (0.00)                             | 581 (2.16)                   | 1,046 (2.09)                      |
| 10            | 1,259 (5.20)                         | 1,438 (5.34)                 | 2,650 (5.29)                      |
| 11            | 1,424 (5.88)                         | 1,538 (5.72)                 | 3,167 (6.32)                      |
| 12            | 1,228 (5.11)                         | 1,423 (5.29)                 | 2,491 (4.97)                      |
| 13            | 0 (0.00)                             | 690 (2.57)                   | 1,245 (2.48)                      |
| 14            | 754 (3.11)                           | 781 (2.90)                   | 1,580 (3.15)                      |
| 15            | 819 (3.40)                           | 888 (3.30)                   | 1,667 (3.33)                      |
| 16            | 1,025 (4.23)                         | 1,273 (4.73)                 | 2,142 (4.27)                      |
| 17            | 1,261 (5.20)                         | 1,672 (6.22)                 | 2,695 (5.38)                      |
| 18            | 329 (1.41)                           | 299 (1.11)                   | 0 (0.00)                          |
| 19            | 1,211 (5.10)                         | 1,432 (5.33)                 | 2,291 (4.57)                      |
| 20            | 515 (2.13)                           | 611 (2.27)                   | 1,070 (2.14)                      |
| 21            | 234 (0.01)                           | 223 (0.01)                   | 440 (0.88)                        |
| 22            | 370 (0.02)                           | 501 (1.86)                   | 798 (1.59)                        |
**Leukocytopenia vs. normal control.** Signaling KEGG pathways with the most significant P-values were pathways in cancer, Axon guidance, focal adhesion and PI3K-Akt signaling pathway (Fig. 5). P-values with the greatest significance included cancer, Down syndrome, diabetes mellitus, cholelithiasis, urinary depression, brain disease and drug abuse (Fig. S2).

**Leukocytosis vs. normal control.** Signaling KEGG pathways with the most significant P-values were associated with Epstein-Barr virus infection, viral myocarditis, focal adhesion, and small cell lung cancer (Fig. 6). Diseases with P-values with the greatest significance included cancer, adenovirus infection, brain tumor, leukoencephalopathy, neurodegenerative disorder and deafness (Fig. S3).

**Key site screening.** The analysis of the leukocytopenia group vs. the leukocytosis group and the leukocytopenia group vs. the normal control group showed a total of 66 differential sites, which corresponded to 37 genes; both the leukocytopenia group vs. the leukocytosis group and the leukocytopenia group vs. the normal control group had 28 sites with hypermethylation and 38 sites with hypomethylation (Table S1). The analysis for the leukocytopenia group vs. the leukocytosis group, and the leukocytosis group vs. normal control group...
showed 27 significantly different sites, which corresponded to 18 genes. A total of 13 hypermethylated fragments were observed in the leukocytopenia group compared with the leukocytosis group, and 14 were observed in the leukocytosis group compared with the normal control group.
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leukocytopenia group compared with the leukocytosis group, and 13 were observed in the leukocytosis group compared with the normal control group (Table SII).

Based on P-values and the functions of related genes according to GenBank (https://www.ncbi.nlm.nih.gov/; accessed on February 18, 2018), four genes were ultimately selected for verification (Table VI). WWP2 is an E3 ubiquitin protein ligase containing a WW domain that participates in the ubiquitination process and viral entry into host cells (28). AKT2 has shown nucleotide binding, protein serine, threonine kinase, protein binding, a TP-binding and transferase activities, and participates in protein amino acid phosphorylation and diabetes development (29). NOX5 has calcium ion binding, hydrogen ionic channel, oxidation reduction enzyme, hemocrystallin-binding and NADP-binding activities, and participates in cytokinesis, electron transfer, apoptosis, hypoxode release and cytokine release. CBARA1 has been demonstrated to have calcium ion-binding activity and participate in defense responses (30).

**Key gene verification**

**WWP2 gene.** A total of five CpG sites in WWP2 fragments (CpG_1, CpG_2/3, CpG_5/6 and CpG_7) were investigated, and four (CpG_1, CpG_2/3, CpG_5/6 and CpG_7) were detected (Fig. 7). Although no significant differences in methylation rates at CpG_2/3 were observed among the three groups (P=0.350), significant differences were observed at CpG_1, CpG_5/6 and CpG_7 (P=0.010, P=0.002 and P<0.001, respectively). The methylation rate at CpG_1 in the leukocytopenia group was significantly higher than that in the leukocytosis and normal control group (P=0.020 and P=0.035, respectively), although no significant difference was observed between the leukocytosis and normal control groups. In the leukocytopenia group, the methylation rate at CpG_7 was significantly higher than that in the leukocytosis group and the normal control group (P<0.001 and P=0.017, respectively), and the rate in the leukocytosis group was significantly lower than that in the normal control group (P=0.013).

**AKT2 gene.** A total of five CpG sites among AKT2 fragments (CpG_1, CpG_2, CpG_3, CpG_4 and CpG_5) were investigated, and three (CpG_1, CpG_3 and CpG_5) were detected.
No significant differences in methylation levels at the CpG_3 and CpG_5 sites were observed among the three groups (P=0.581 and P=0.215, respectively); however, the levels of CpG_1 methylation in the leukocytopenia group were significantly higher than those in the leukocytosis and normal control groups (P<0.001 and P=0.045, respectively; Fig. 8), and the leukocytosis group had a significantly lower methylation level at this site than the normal control group (P=0.001).

CBARA1 gene. A total of four CpG sites among CBARA1 fragments (CpG_1, CpG_2, CpG_3 and CpG_4) were investigated and detected. No significant differences in methylation at the CpG_1, CpG_3 and CpG_4 sites were found among the three groups (all P>0.05), whereas a significant difference was observed for CpG_2 (P=0.011; Fig. 9). The CpG_2 methylation level in the leukocytosis group was significantly lower than that in the normal control group (P=0.001).

NOX5 gene. For NOX5 fragments (CpG_1, CpG_2, CpG_3 and CpG_4), four CpG sites were investigated and three (CpG_2, CpG_3 and CpG_4) were detected. No significant differences in methylation levels at the CpG_2 and CpG_4 sites were observed among the three groups (both P>0.05), whereas a significant difference was observed for CpG_3 (P=0.005; Fig. 10). The methylation level at the CpG_3 in the leukocytosis group was significantly lower than that in the normal control group (P=0.004).

Discussion

Death due to infection is currently the cause of ~25% of all deaths caused by diseases worldwide (1). In recent years, the incidence rate of infection has remained high for various reasons, including disease complications, the use of glucocorticoids, immunocompromise, bacterial resistance and invasive operations (1). Regardless of leukocytopenia or leukocytosis, infections are primarily caused by pathogens, commonly viruses and bacteria, which initiate infection by activating innate immunity. Non-specific innate immunity (30), also known as natural immunity, plays an important role in acute infection, and neutrophils, lymphocytes, eosinophils and basophils are important immunocytes involved in this process (31). These cells reach the site of infection via adhering, binding, cellular escape and cellular migration (32), and release reactive oxygen species and antimicrobial lytic proteins, such as peroxidase, lysozyme, alkaline phosphatase and acid hydrolase (33). These cells engulf, kill and digest microorganisms, thereby playing a direct role in the immune system (34). In clinical settings, post-infection leukocytopenia occurs frequently; the symptoms of patients with
post-infection leukocytopenia are generally severe and difficult to treat, and require an extensive anti-infection treatment course (35). Furthermore, abnormalities such as cell migration disorders, myelosis, reduced chemotactic factor secretion and immunocyte surface receptor abnormalities may exist in these patients (36). In the present study, the general clinical data of patients with leukocytopenia, including age, sex, infection site, routine blood tests, and levels of IL-6 and serum complement, were analyzed and whole-genome methylation analysis was performed. Key sites of methylation were then screened with the aim of exploring post-infection leukocytopenia-associated molecular mechanisms in order to provide a novel avenue for gene diagnosis and treatment of such diseases.

The general data showed higher levels of PCT and IL-6 in the leukocytosis group compared with the leukocytopenia group. Furthermore, patients in the leukocytopenia group had a higher APACHE II score than those in the leukocytosis group, indicating a worse prognosis. Analysis of serum complement C3 and C5 levels revealed significant decreases in the leukocytopenia group compared with the leukocytosis and normal control groups. This finding suggested that post-infection leukocytopenia may be complicated by abnormally low expression of chemotactic factors, and that abnormal expression of chemotactic factors may partly contribute to the pathogenesis of infection-associated leukocytopenia. Genome methylation detection was then performed to further determine the mechanisms via which chemotactic factors are abnormally expressed, as well as to find other mechanisms underlying post-infection leukocytopenia.

DNA methylation is classically studied in epigenetic research and is the most representative feature of epigenetics. Moreover, changes in methylation may underlie the pathogenesis of a variety of diseases (37). In the present study, whole-genome methylation detection was performed, revealing significant enrichment in various pathways and disease-causing functions. The significantly enriched pathways and functions closely associated with the development of post-infection leukocytopenia included receptor activity, transmembrane signal transduction and potassium ion channels. These results indicated that multiple pathways are responsible for the development of this disease via network interactions. Diseases with significant enrichment included cancer, neurodegenerative disorders, intestinal diseases, cardiovascular diseases, endocrine diseases and urinary diseases, although most of these were not associated with the disease investigated in the present study. This may be partially due to the fact that genome methylation detection is primarily applied in research into tumors, as well as cardiovascular and nervous system diseases (38,39), whereas its application in investigating infectious disease has not been reported to date. In addition, this finding may also be due to limitations such as small sample size and experimental error. Therefore, experiments with a larger sample size should be performed in the future to obtain more accurate functional enrichment analysis data.

Intersection analysis of differentially methylated gene sites between the leukocytopenia and leukocytosis groups, and the leukocytopenia and normal control groups may suggest the cause of leukocytopenia, and that between the leukocytopenia and leukocytosis groups, and the leukocytosis and normal control groups may indicate the cause of leukocytosis. Based on reports from the literature and P-values obtained, NOX5, WWP2, AKT2 and CBARA1 were selected for further verification. NOX5 is expressed in the spleen and lymph node cells (40), and its functions primarily include calcium ion binding, hydrogen ion channels, superoxide production, NADPH oxidase, oxidoreductase, flavin adenine dinucleotide binding and NADP binding (41). NOX5 protein primarily participates in cytokinesis, electron transfer, apoptosis, cellular proliferation, proton transportation regulation, superoxide release, cytokine secretion and cell fusion regulation (42). According to a previous study (43), monocytes can differentiate into various subsets of dendritic cells, and NOX5-p22phox complexes may promote the differentiation of monocytes into dendritic cells via Janus kinase/signal transducers, and activators of transcription/MAPK and NF-κB pathways. NOX5 may be able to reduce oxidative stress and may be associated with calcium ion channel-related oxidative stress in lymph nodes (44). WWP2 also exhibits protein binding and ligase activities (28); it is involved in the ubiquitination process, circulation and viral entry into host cells (28). WWP2 has a negative regulatory effect on Toll-like receptor 3-mediated innate immunity (45). CBARA1 has calcium ion-binding activity and participates in the immune response (26). AKT2 shows nucleotide-binding, protein serine, threonine kinase, protein-binding, ATP-binding and transferase activities, and participates in protein amino acid phosphorylation as well as the development of diabetes (46,47). Moreover, it has been shown that AKT2 is associated with dendritic cell migration and the functions of neutrophils, as well as vascular permeability, edema and leukocytic emigration in acute inflammation (48). AKT2 knockout can ameliorate pulmonary injury by reducing alveolar cell depolarization (49), and AKT2 may also participate in the development of pulmonary fibrosis via transforming growth factor-β1 and IL-13 (50).

The present study has a number of limitations. First, the sample used for genome methylation detection was small, which may have resulted in false positive results. Therefore, these findings need to be further verified by experiments with larger sample sizes. Second, as most studies into methylation and genomes have focused on tumor research, signaling pathway analysis and disease enrichment analysis in this study, based on the current datasets, did not identify the direct association of the methylated genes with leukocytopenia. Therefore, studies involving animal experiments and gene knockout need to be performed in the future.

In conclusion, four genes may participate in the occurrence of leukocytopenia after infection: NOX5, WWP2, AKT2 and CBARA1. Based on the results of this study, multiple experiments with larger sample sizes should be carried out and more genes should be selected for verification. These experiments may be useful for clarifying the molecular mechanisms underlying leukocytopenia following infection, and identifying novel strategies for diagnosis and treatment.

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Availability of data and materials

The datasets used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors' contributions

All authors have made contributions to the current work. CW and XY designed the study; CW led the writing of the article. XM, WW, MD, and RJ conducted the experiments and collected the data. ZL, DL and YL conducted data analysis. XY supervised the process and offered constructive advice. All authors read and approved the final manuscript.

Ethics approval and consent to participate

This study was conducted in accordance with the Declaration of Helsinki. Approval was obtained from the Ethics Committee of People's Hospital of Xinjiang Uygur Autonomous Region (policy no. 2016056). Written informed consent was obtained from each participant.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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