



Comprehensively analyze the expression and prognostic role for ten-eleven translocations (TETs) in acute myeloid leukemia

Yan Huang^{1#}, Jie Wei^{1#}, Xunjun Huang¹, Weijie Zhou¹, Yuling Xu², Dong-Hong Deng², Peng Cheng²

¹Department of Hematology and Rheumatology, People's Hospital of Baise, Baise, China; ²Department of Hematology, the First Affiliated Hospital of Guangxi Medical University, Nanning, China

Contributions: (I) Conception and design: J Wei, Y Huang; (II) Administrative support: X Huang, Y Xu; (III) Provision of study materials or patients: W Zhou; (IV) Collection and assembly of data: J Wei, Y Huang; (V) Data analysis and interpretation: DH Deng, P Cheng; (VI) Manuscript writing: All authors; (VII) Final approval of manuscript: All authors.

[#]These authors contributed equally to this work.

Correspondence to: Yuling Xu. Department of Hematology, The First Affiliated Hospital of Guangxi Medical University, Nanning, China. Email: xuyulinggx@163.com; Dong-Hong Deng. Department of Hematology, The First Affiliated Hospital of Guangxi Medical University, Nanning, China. Email: ddh_gx@163.com; Peng Cheng. Department of Hematology, The First Affiliated Hospital of Guangxi Medical University, Nanning 530000, China. Email: gxchengpeng@163.com.

Background: The ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs) and promote the locus-specific reversal of DNA. The role of TETs in acute myeloid leukemia (AML) is mostly unknown.

Methods: TETs mRNA expression levels were analyzed via Gene Expression Profiling Interactive Analysis (GEPIA). The association TETs expression levels and methylation with prognosis by UALCAN GenomicScape, and METHsurv. We analyzed TETs' aberration types, located mutations, and structures via cBioPortal. GeneMANIA performed the functional network. Gene ontology (GO) enrichment was analyzed via LinkedOmics. MiWalk identified miRNAs, miTarbase, and TargetScan. Transcription factor (TF) targets were analyzed via ChEA3. GSCALite analyzed the role of these defined genes in cancer pathways and potential drug targets. Finally, we selected AML patients in our department to investigate the mutated types of TETs.

Results: TETs expression level results showed TET1 (P=0.003) and TET2 (P=0.004) overexpressed in Haferlach leukemia samples, TET3 (P=4.04e-8) downregulation in Andersson leukemia samples. TET2 and TET3 overexpression but TET1 downregulation in the GEPIA database. Overexpression of TET2 leads to positive outcomes (P=0.0091). The upregulation of TET2 led to poor survival for CN-AML patients, but downregulation of TET3 indicated a satisfactory prognosis. The hypermethylation of TETs like cg24705708 (P=0.036), cg05976228 (P=0.022), cg19127638 (P=0.022), cg15254238 (P=0.025), cg07669489 (P=0.037) indicate poor outcomes. Overexpression of GALNS (P=0.024) as an adverse biomarker, downregulation of E2F5 (P=0.037), MAP7 (P=0.019), and NRIP1 (P=0.0013) indicated good prognosis. Regulatory network analysis indicated TETs' functions, including covalent chromatin modification, histone modification, DNA methylation, or demethylation. Enrichment functions involving TETs participate in several cancer pathways, including DNA repair response and receptor tyrosine kinase (RTK) signaling pathway. TETs are sensitive to belinostat, ceranib-2, docetaxel, tivantinib, and vincristine.

Conclusion: Present study showed that TETs have different expressions in AML, and the expression levels of TETs lead to different outcomes of AML. The TETs cancer pathway analysis will also provide potential therapy methods for AML patients with TETs aberrations.

Keywords: Acute myeloid leukemia (AML); bioinformatics analysis; ten-eleven translocation (TETs); prognosis values

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Introduction

Acute myeloid leukemia (AML) is a heterogeneous disease of the blood system characterized by the clonal expansion and differentiation arrest of myeloid progenitor cells. The incidence of AML increases by age, and the older patients, the more mortality (1). An estimated 19,940 new diagnostic AML patients, and 11,180 people die of AML in the United States (2). However, intensive chemotherapy is not a suitable option for many older patients with significant comorbidities, baseline organ dysfunction, or poor performance status, in whom the risk of complications and treatment-related mortality is unacceptably high. The worse thing is that relapsed AML patients only have less than 25% complete remission rate (3). Since the more efficient salvage regimens have been into practice, the complete response rate has increased, but these patients cannot obtain substantial CR duration (4.9 to 9.8 months) or survival (6.2 to 8.7 months) (4), this result did not be satisfied. Furthermore, the numbers of AML patients gradually increase in times. Thus, we urgently need more sensitive and specific markers for early diagnosis, more efficient but less side effect therapy for AML patients.

Although the advance of cell biology and comprehensive genomic analyses has shown the possible leukemogenesis mechanisms, it is still incompletely understood. The genes aberrations have been confirmed, one of the essential leukemogenesis drivers, even can as biomarkers indicated the favorable or adverse outcome of AML patients. Many studies showed that some improved cytogenetic changes play crucial roles in tumorigenesis and prognosis of AML, including aberration of TP53 (5), WT1 (6), double CEBPA mutation (7), RUNX1 (8), DNMT3A (9), FLT3 (10), and so on. DNA's methylation has appeared relatively stable epigenetic process. However, with recent studies improve the function of ten-eleven translocation (TET) family proteins as 5-methylcytosine oxidases, the view has been reversed. The TET family, including TET1, TET2, and TET3. Over the last decade, TET protein or family functions and differential expression in various types of cancers have further insight. Many studies explore the TETs role in different cancers. For solid tumors, TET1 accelerates the triple-negative breast cancer exacerbation (11), but TET2 as inhibitors tumorigenesis of breast cancer cells in another study (12). In ovarian cancer, high expression of TET3 as an adverse biomarker (13). About hematological malignancies, TET1 mutations have been observed in chronic lymphocytic leukemia (CLL) (14) and T-cell acute lymphoblastic leukemia

(T-ALL) (15), and AML (16). TET2 mutations also were found in MDS (17) and AML (18). However, TET3 changes less be found in hematological malignancies. In different cancer types, the TETs functions may be different. For hematological cancers, promoting DNA demethylation and regulate immunity is the most important biological function of TETs (19). Significant gene methylation, immunity tolerance, and immunity evasion significantly influence tumorigenesis, therapy resistance, and tumor progress. From these, we use public databases to analyse the prognostic value of TETs in AML and observing whether there are significant expression levels between AML samples and normal samples. We present the following article in accordance with the MDAR reporting checklist (available at <http://dx.doi.org/10.21037/tcr-20-3149>).

Methods

Expression analysis

The OncoPrint database (20) and GEPIA (Gene Expression Profiling Interactive Analysis) (21) were preliminarily used to analyze the TETs expression levels between tumor samples and normal samples. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

Analyzing the aberrations types, location of mutations, and structure of TETs

We used cBioPortal to learn the types of aberrations of TETs, perform the aberration locations, and construct the three-dimension structure of TETs (22).

Survival analysis

To investigate the relation TETs expression levels with the prognosis of AML patients, we use the UALCAN to analyze the TCGA data of AML patients (23). Further, we use GenomicScape (<http://www.genomicscape.com>) that data (GSE12417) derived from the GEO database to validate the survival results of AML (24). Also, we analyze the significantly correlated genes prognostic value of AML. Finally, we explore the relation TETs methylation with the prognosis of AML patients via METHSurv (25).

GeneMANIA analysis

GeneMANIA is a commonly used website for performing

protein-protein interaction (PPI) network analysis and predicting the function of preferred genes (26). This user-friendly online tool can display gene or gene lists using bioinformatics methods, including gene co-expression, physical interaction, gene co-location, gene enrichment analysis, and website prediction. We predicted the TETs family's function and significantly correlated genes and visualizing the gene networks via GeneMANIA.

Constructing the miRNA networks

We search for the miRNAs that target TETs via miTarbase (27), miRTarBase (27), and miWalk (28) in various databases to find out. Further, we performed the Venn diagram to find the TETs' co-targets miRNA.

LinkOmics and ChEA3 analysis

The LinkedOmics database (<http://www.linkedomics.org/login.php>) is a Web-based platform for analyzing 32 TCGA cancer-associated multi-dimensional datasets (29). Data from the LinkFinder results were signed and ranked, and GSEA was used to analyze GO (CC, BP, and MF), KEGG pathways, and kinase-target enrichment. We used LinkOmics to analyze the CC, BP, and MF, kinase-target enrichment, and the significantly correlated genes of TETs. Further, we used the ChEA3 to find the TETs' significant transcription factors (TFs) annotating the potential biological functions, and constructing the network of top 10 TFs (29).

Analyzing the cancer pathway activity and drugs sensitivity of defined genes

Cancer Analysis (GSCALite) analyzes Gene expression associated cancer pathway activity and Drug sensitivity for genes (30). We used the GSCALite dataset to explore the role of defined genes in the cancer pathway, further investigate the drug sensitivity of defined genes.

Validation of mutated types of TETs in AML patients

We selected the AML patients with TETs mutations in The Department of Hematology, the First Affiliated Hospital of Guangxi Medical University, to find the mutated types of TETs in our department.

Statistical analysis

The gene expression, survival analysis, and the protein-protein network relations P value <0.05 is considered a significant difference in our study.

Results

Transcriptional expression levels of TETs in AML patients

To identify the differential expression levels of TETs in AML and normal samples, the TETs mRNA levels in leukemia samples and control samples were analyzed by the Oncomine (*Figure 1A,B,C*) and GEPIA (*Figure 1D,E,F*) databases. The results of Oncomine analysis showed that the expression of TET1 (P=0.003) and TET2 (P=0.004) was significant overexpression in AML patients in Haferlach leukemia, but the TET3 (P=4.04e-8) was downregulation in Andersson leukemia samples. To further validate to TETs expression level, we used the GEPIA to perform analysis. The results indicated that TET2 and TET3 were significantly upregulated expression, but TET1 was significantly downregulated expression.

Aberrations types and the structure of TETs

To investigate the aberrations types, aberrations locations, 3D structure of TETs, we used the C-BioPortal database to analyze. As the results have shown the top 10 co-expression of TETs including FLT3, NPM1, DNMT3A, CEBPA, NRAS, IDH2, SRSF2, RUNX1, STAG2, and ASXL1 (*Figure 2A*). The most standard mutated types of TET2 and TET1 were deep deletion and missense mutations (*Figure 2B*). The locations of TETs were shown in *Figure 2C*.

The relation between TETs expression levels and the prognosis of AML patients

To understand whether the expression levels of TETs would have a significant influence on AML patients' survival, we used the UACLAN database to clarify the relation between TETs expression and prognostic role of AML. As the results have shown, the higher expression TET2 may lead to positive outcomes of AML patients (P=0.0091). However, TET1 and TET3 expression levels did not impact the prognosis of AML patients (*Figure 3A,B,C*). To further

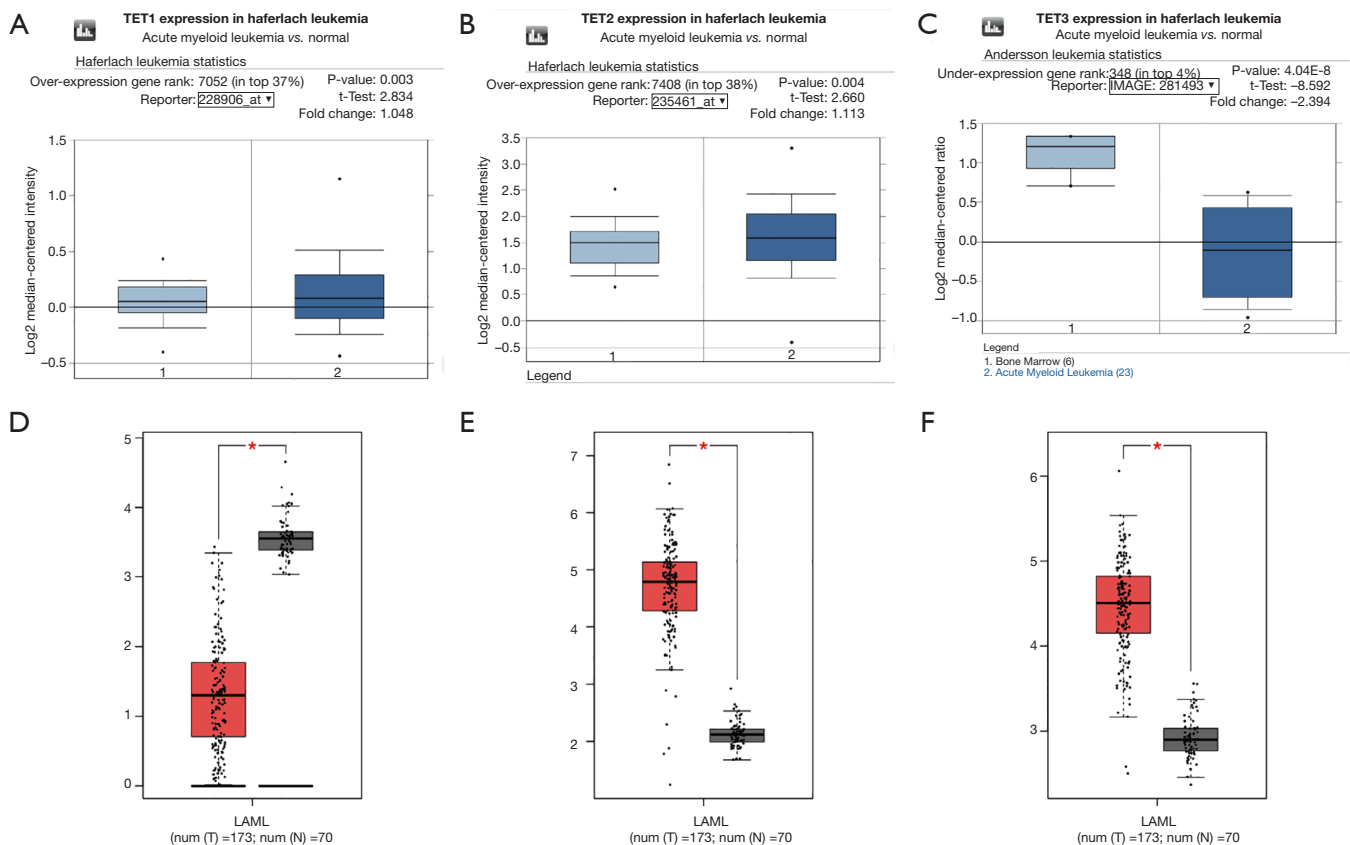


Figure 1 The expression levels of TETs between AML and normal samples. (A) The AML samples with the higher expression of TET1 in Haferlach leukemia (Oncomine). (B) TET2 overexpresses in AML samples in Haferlach leukemia (Oncomine). (C) TET3 downregulated in tumor samples in Andersson leukemia samples (Oncomine). (D) TET1 was downregulated in AML samples (GEPiA). (E) TET2 was an upregulation in AML samples (GEPiA). (F) Overexpression of TET3 was detected in AML patients (GEPiA). TETs, a ten-eleven translocation (TET) family, oxidize 5-methylcytosines (5mCs); AML, acute myeloid leukemia; GEPiA, Gene Expression Profiling Interactive Analysis.

validate the impact of TETs expressions for AML patients, we used the GSE12417 data to analyze. The results showed that high expression TET2 leads to poor survival for CN-AML patients, but high expression TET3 indicated a protective factor (Figure 3D,E,F).

Prognostics role of significantly correlated genes in AML

To further investigate the potential role of differentially expressed TETs in AML, the LinkFinder module of LinkedOmics was used to analyze mRNA sequencing data from 173 patients in TCGA. In the volcano plot, the dark red dots show a significantly positive correlation with TET1, TET2, and TET3, whereas dark green dots show a significantly negative correlation (false discovery rate, FDR <0.01). The top 50 significant gene sets positively

and negatively associated with TET1, TET2, TET3, as shown in the heat maps (Figure 4A,B,C,D,E,F,G,H,I). Meanwhile, the statistical scatter plots show the association TETs expression with significantly correlated genes (Pearson's correlation ≥ 0.6). From the results of association genes, we can observe APT11C, CD63, OS9, GALNS, NRIP1, MAP7, and E2F5 were strong associations with TET1; ZSWIM6, QKI, SNX13, and SSFA2 were strong associations with TET2; PDPK1 and NCOA6 were strong associations with TET3 (Figure 5A,B,C,D,E,F,G,H,I,J,K,L,M). To explore whether the strong association with TETs genes that have a significant impact on AML patients, we used UALCAN to analyze the TET1, TET2, TET3 strong related genes in OS of AML patients. There were four correlated significant genes have a significant impact on the OS of AML patients,

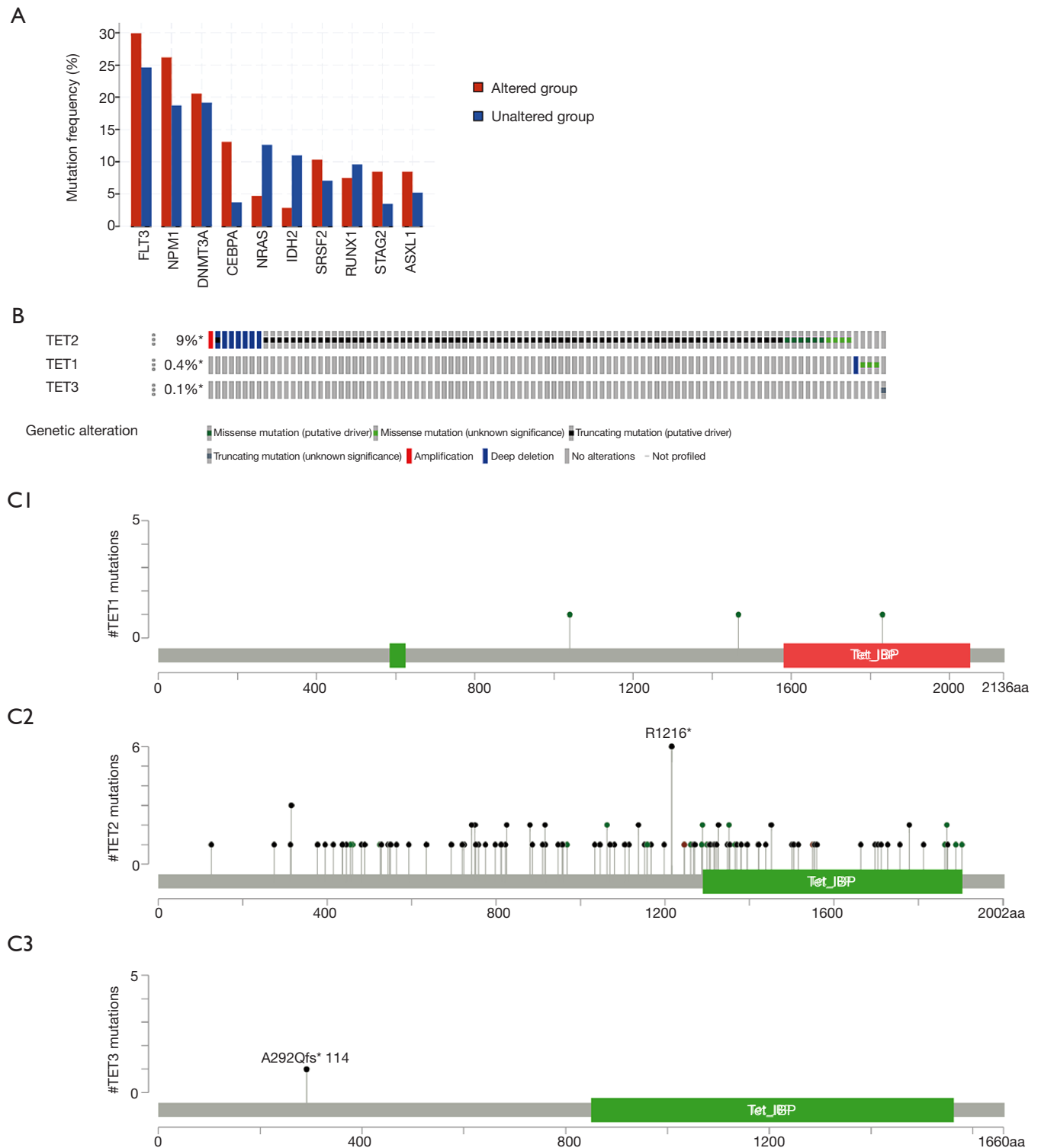


Figure 2 Aberration types, located mutation, the co-mutation status of TETs in AML. (A) The co-mutation frequency in TETs aberration groups and unmutated group. (B) Aberration types and frequency of TETs in AML; truncating mutations are the most common, 9%, 0.4%, and 0.1% mutations of TETs in AML, respectively. (C) The mutated locations of TETs. TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs); AML, acute myeloid leukemia.

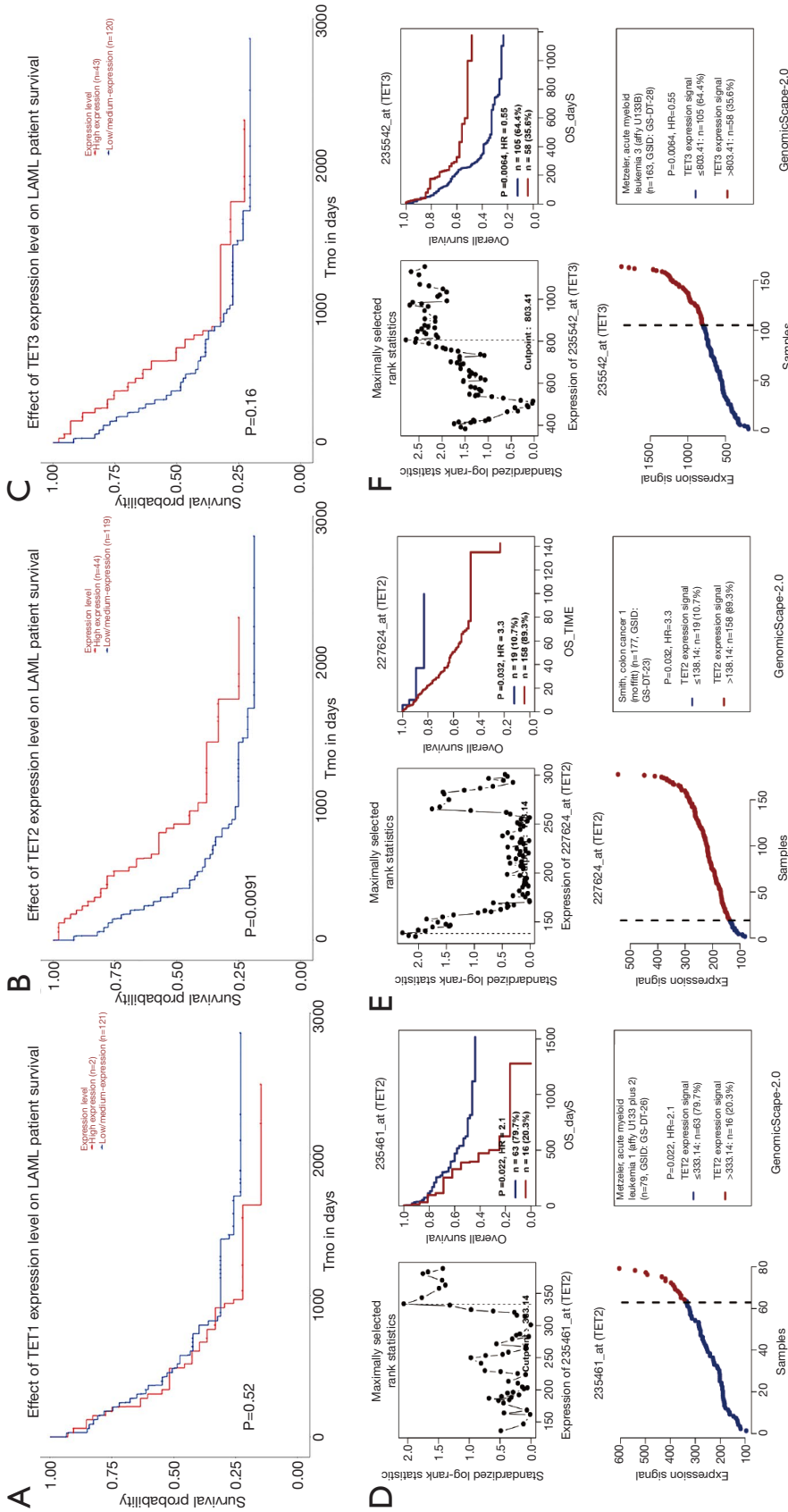


Figure 3 The prognostic role of the TETs gene (UALCAN and GenomicScape). (A,B,C) High TET2 expression was correlated with better prognosis of OS in AML, but the expression levels of TET1 and TET3 do not significantly influence AML. (D,E,F) Overexpression TET2 indicates the poor outcome of CN-AML, but overexpression of TET3 shows the better prognosis of CN-AML. TET1, ten-eleven translocation 1; TET2, ten-eleven translocation 2; TET3, ten-eleven translocation 3; AML, acute myeloid leukemia; CN-AML, cytogenetically normal acute myeloid leukemia.

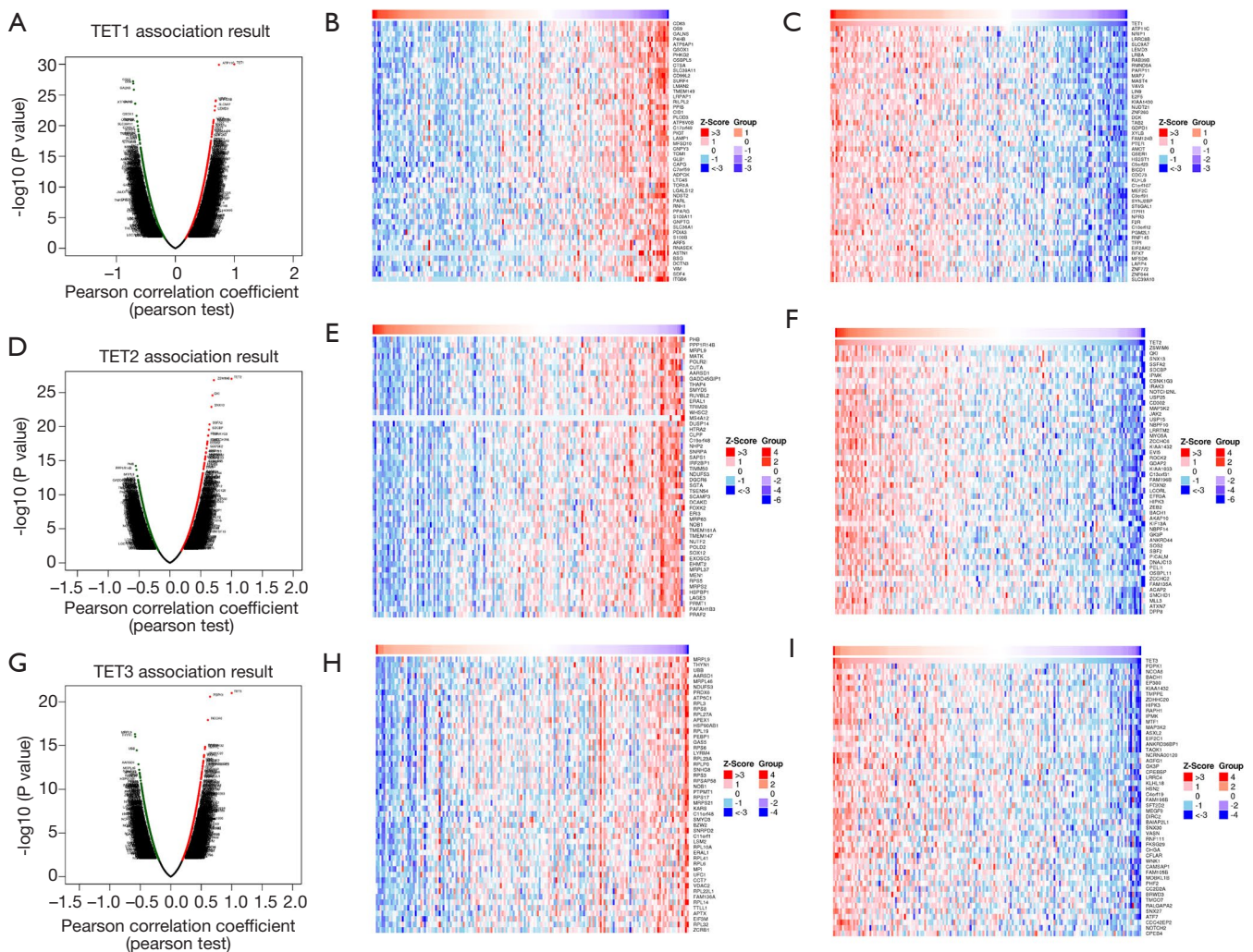


Figure 4 Correlated significant genes of TETs (LinkedOmics). (A,B,C,D,E,F,G,H,I) Volcano plots and heat maps showed genes positively and negatively correlated with TETs in AML, respectively (top 50). (D,F) Red suggests positively correlated genes, and green shows negatively correlated genes. TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs); AML, acute myeloid leukemia.

the high expression GALNS (P=0.024) leads to an adverse outcome, the downregulation of E2F5 (P=0.037), MAP7 (P=0.019), and NRIP1(P=0.0013) indicated the excellent prognosis (Figure 6A,B,C,D).

The relationship TETs methylation with the prognosis of AML

From the important role of methylation in tumorigenesis and prognosis of AML patients, we used METHSurv to analyze the TETs methylation status and the association with prognosis for AML patients, respectively. The locations of TETs methylation were shown in the heatmap

(Figure 7A,B,C) and the prognosis of TETs methylation for AML patients was shown in Figure 8A,B,C,D,E,F,G,H,I. The hypermethylation of TET1 including cg19127638, cg12548760, cg18515801, cg01093854, cg13810683, and cg27426824, and the hypermethylation of TET2 including cg20586654, cg08530497, cg22794775, cg09666717, and cg17862558. TET3 has the most frequency of hypermethylation including cg02956499, cg13808088, cg00755592, cg15827185, cg21855109, cg05976228, cg25299214, cg17213010, cg24705708, cg02237855, cg01244346, and cg01355757 (Figure 7A,B,C). Further, we analyzed the prognostics value of TETs methylation in AML patients. The results indicated that

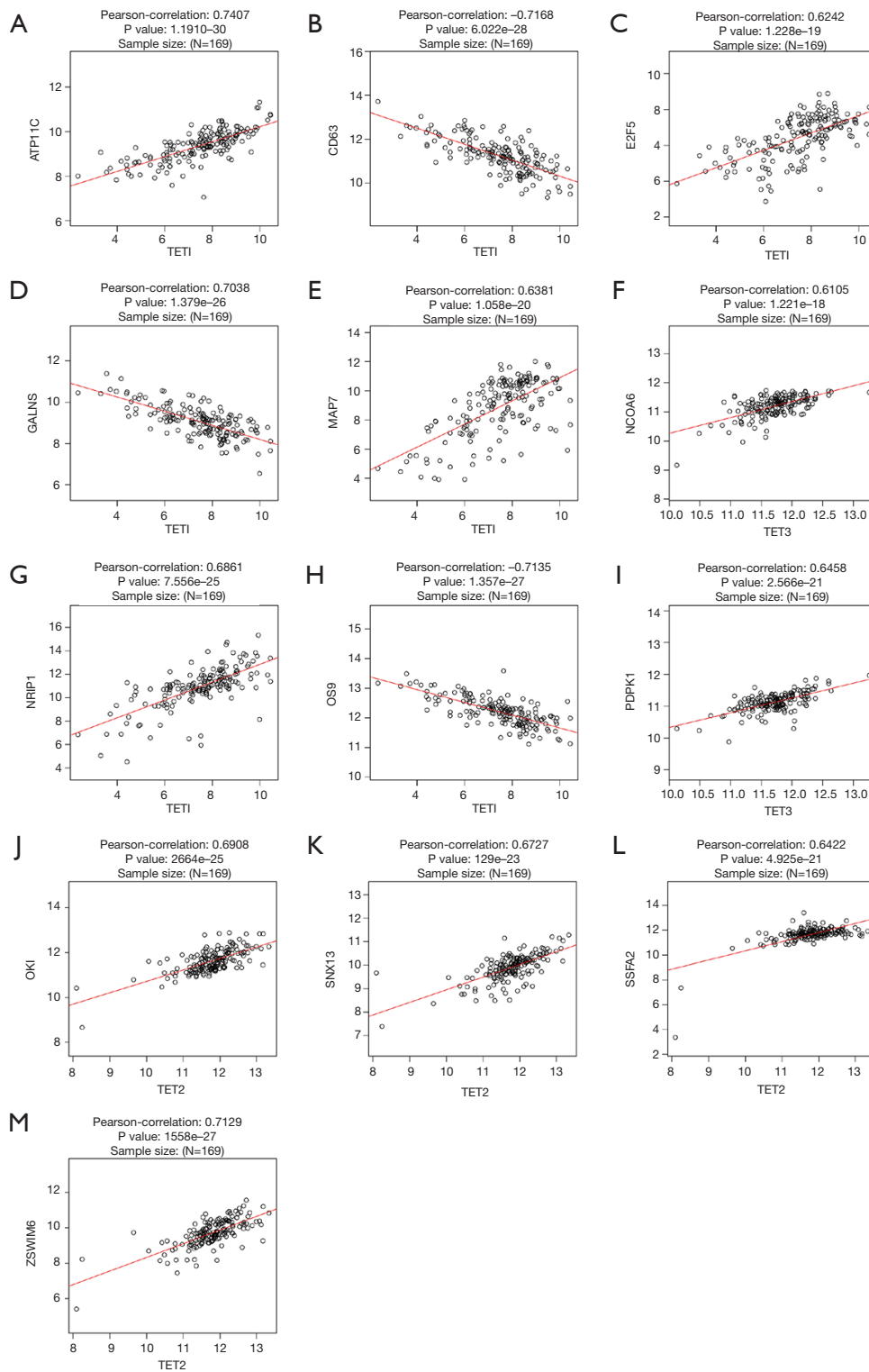


Figure 5 Gene correlation expression analysis for TETs (LinkedOmics). (A,B,C,D,E,F,G) The scatter plots show the spearman-correlation of TET1 expression with expression of in AML (H,I,J,K). The scatter plots show the spearman-correlation of TET2 expression with expression of in AML (L,M). The scatter plots show spearman-correlation of TET3 expression with expression of in AML. TET1, ten-eleven translocation 1; TET2, ten-eleven translocation 2; TET3, ten-eleven translocation 3; AML, acute myeloid leukemia.

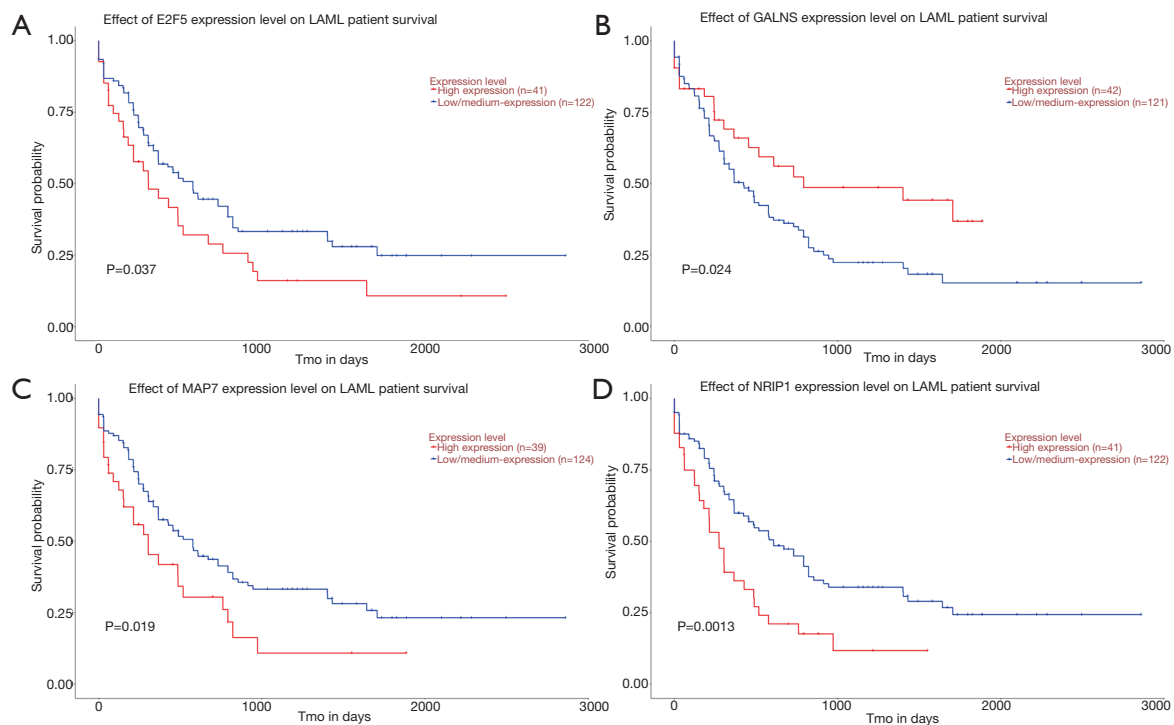


Figure 6 Prognostic analysis of genes correlated with TETs in AML (UALCAN). (A) The overall survival curves of E2F5. (B) The overall survival curves of GALNS. (C) The overall survival curves of MAP7. (D) The overall survival curves of NRIP1. TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs); E2F5, E2F transcription factor 5 protein; GALNS, N-acetylgalactosamine-6-sulfatase; MAP7, ensconsin, E-MAP-115; NRIP1, nuclear receptor-interacting protein 1; AML, acute myeloid leukemia.

hypermethylation of cg24705708 ($P=0.036$), cg05976228 ($P=0.022$), cg21855109 ($P=0.022$), and cg25299214 ($P=0.0028$), cg17862558 ($P=0.0073$), cg13810683 ($P=0.013$), cg19127638 ($P=0.022$), cg15254238 ($P=0.025$), and cg07669489 ($P=0.037$) were the adverse biomarkers for AML patients (Figure 8A,B,C,D,E,F,G,H,I).

Functional enrichment analysis of TETs

To investigate the potential functional enrichment among TETs, we used the LinkedOmics database to analyze the significant enriched GO biological process (BP), cellular component (CC), molecular functions (MF), and target kinase. The main enrichments result in DNA transcription, RNA production, energy metabolism pathway, proteolysis, angiogenesis, and so on (Figures 9-11). The significant kinase targets of TET1 were CDK2 and TGFBR2. For TET2, there were CSNK1D, IRAK4, SYK, EGFR, MAP3K4, and NEK6 kinase targets. TET3 has the most numbers of kinase targets that include MYLK3, MYLK4,

SGK1, CSNK1D, PLK2, CDK2, ILK, PRK CZ, PRKCE, MAPK10, MYLK2, and MYLK. Among the kinase targets, CDK2 was the TET1 and TET3 kinase co-targets, and CSNK1D was the kinase co-targets of TET2 and TET3 (Figures 9D,10D,11D).

Further, the protein-protein network of TETs and the significantly correlated genes was constructed via GeneMANIA. Both of TETs interacted with CXXC4, but only TET3 and TET1 interacted with DNMT1, there still were many other significant interactions. The PPI network showed that the functions of TETs were covalent chromatin modification, demethylation, histone modification, histone H3-K4 methylation, histone lysine methylation, and DNA methylation or demethylation (Figure 12).

The regulated miRNA network of TETs

We used miTarbase, TargetScan, and miWalk databases to find the regulated miRNA of TETs, there were 690, 298, and 68 regulated miRNAs for TET2, TET1, and

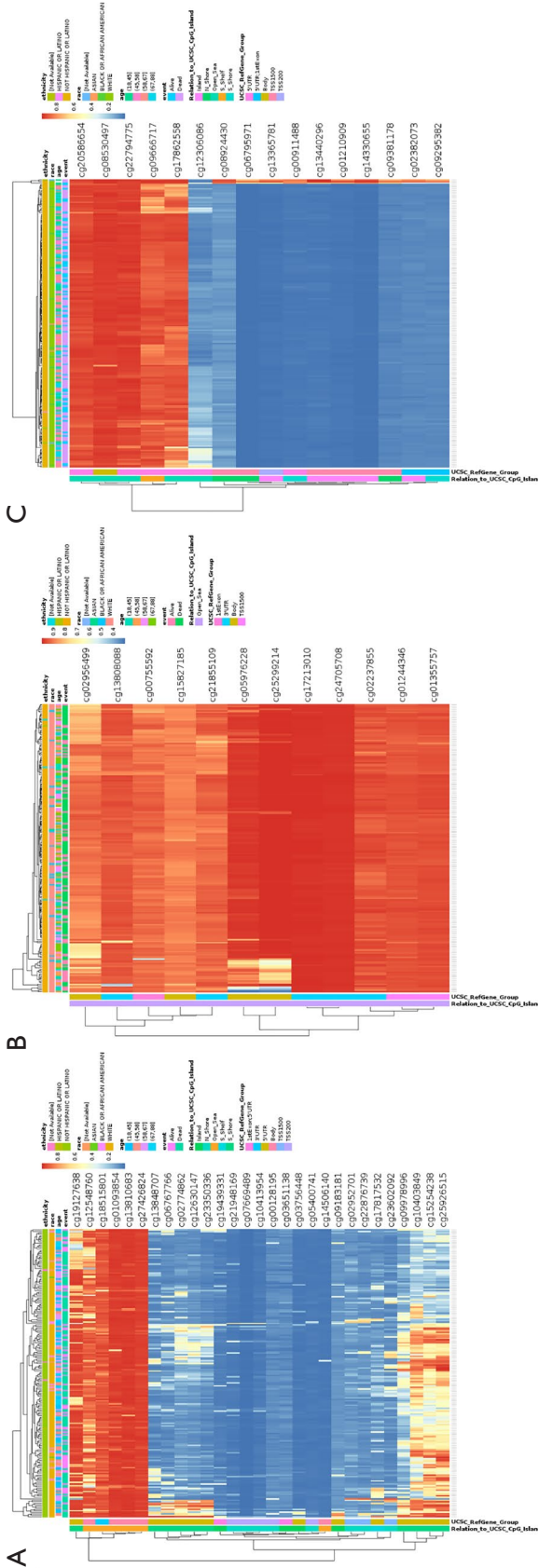


Figure 7 The heatmap methylation of TEIs for AML. (A,B,C) Heat maps showing methylation of TEIs for AML. Red suggests hypomethylation; blue indicates hypomethylation. TEIs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs); AML, acute myeloid leukemia.

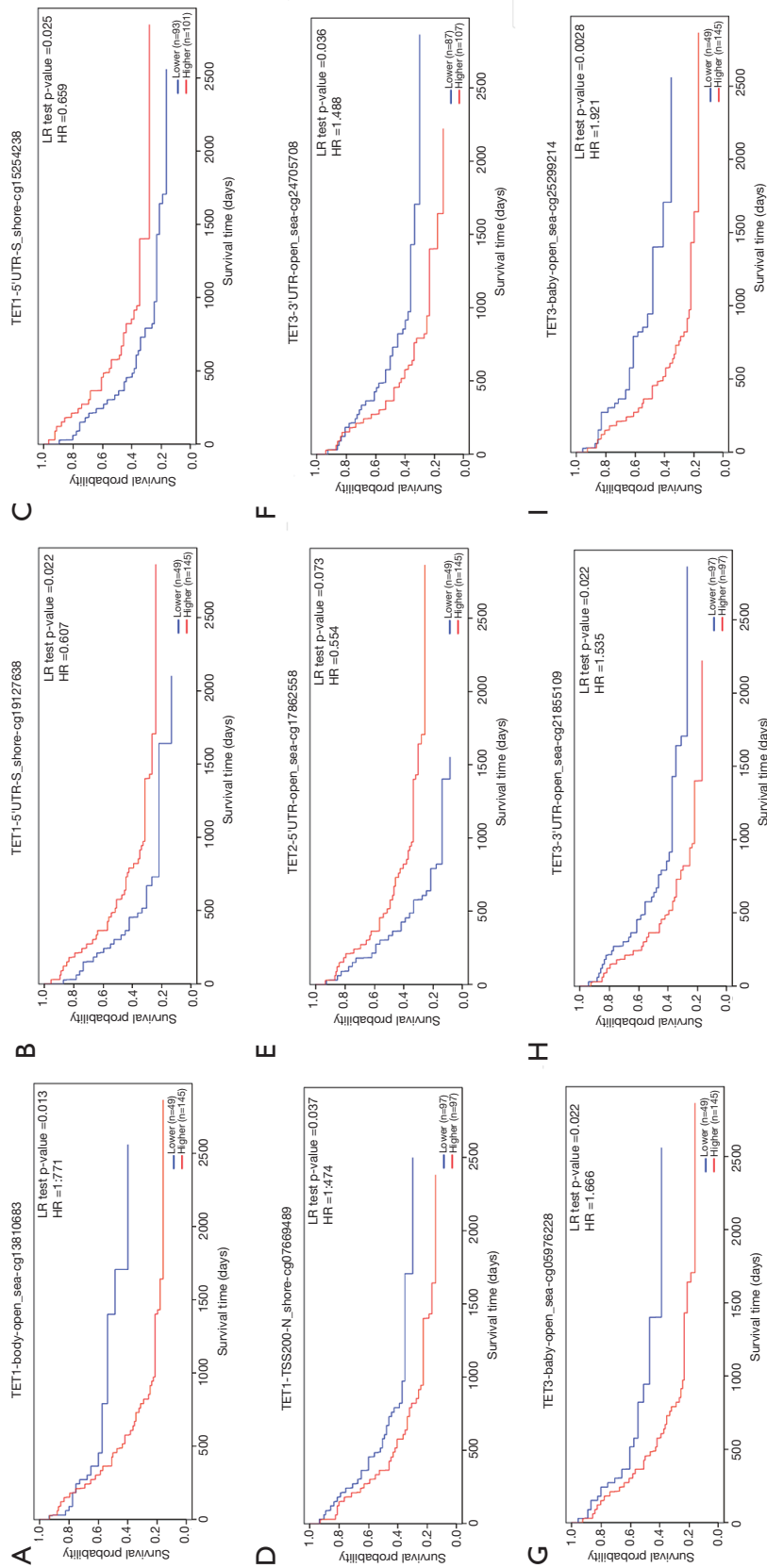


Figure 8 The prognostics role of TETs methylation in AML. (A,B,C,D) The prognostics role of TET1 methylation for AML. (E) The prognostics role of TET2 methylation for AML (F,G,H,I) The prognostics role of TET3 methylation for AML. TET1, ten-eleven translocation 1; TET2, ten-eleven translocation 2; TET3, ten-eleven translocation 3; AML, acute myeloid leukemia.

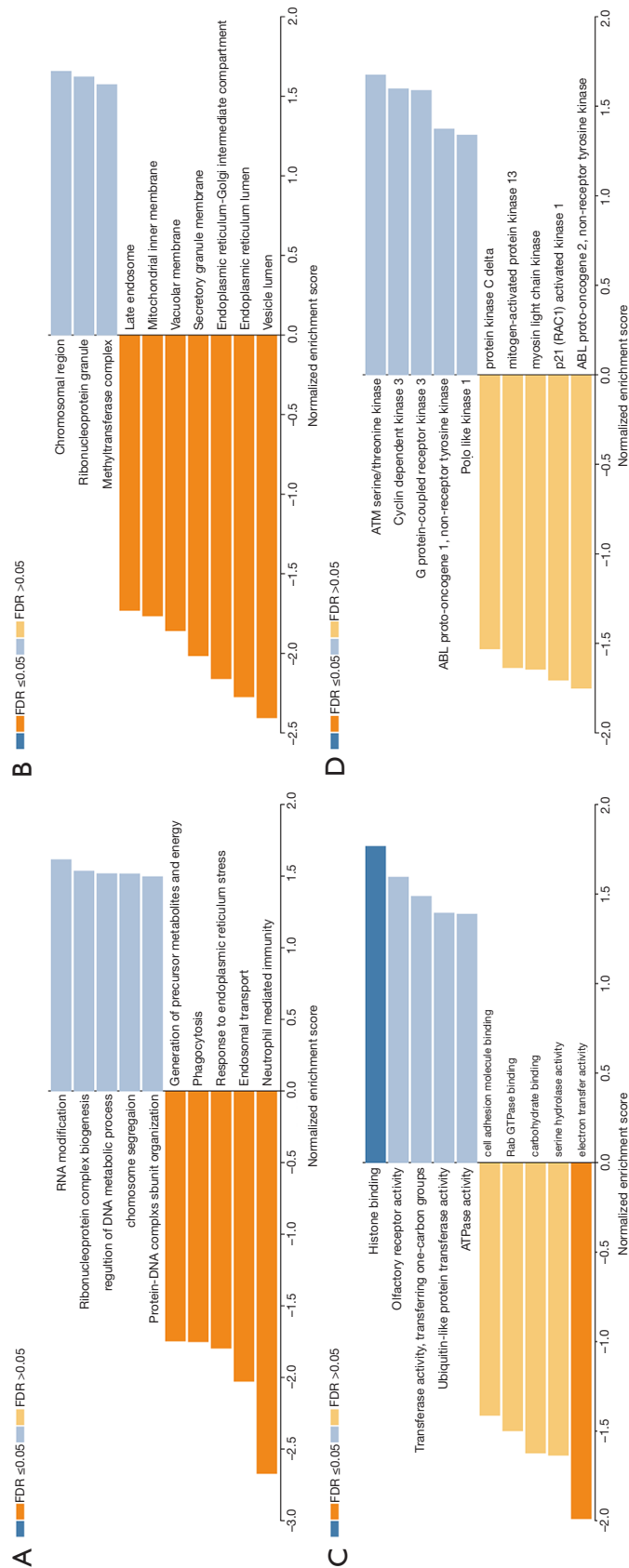


Figure 9 Functions enrichment of TET1 in AML (LinkedOmics). (A) BP of TET1 in AML. (B) CC of TET1 in AML. (C) MF of TET1 in AML. (D) Targets kinases of TET1 in AML. TET1, ten-eleven translocation 1; AML, acute myeloid leukemia; BP, biological process; CC, cellular component; MF, molecular functions.

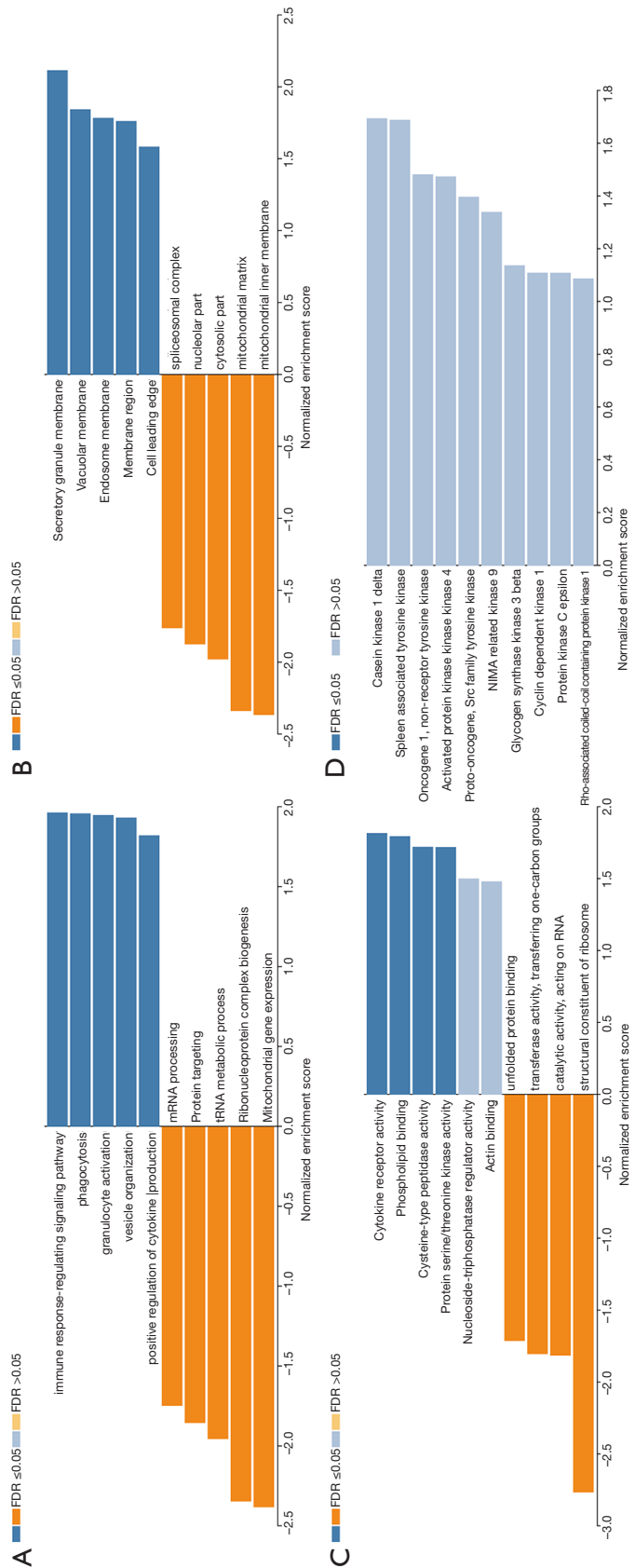


Figure 10 Functions enrichment of TET2 in AML (LinkedOmics) (A) BP of TET2 in AML. (B) CC of TET2 in AML. (C) MF of TET2 in AML. (D) Targets kinases of TET2 in AML. TET2, ten-eleven translocation 2; AML, acute myeloid leukemia; BP, biological process; CC, cellular component; MF, molecular functions.

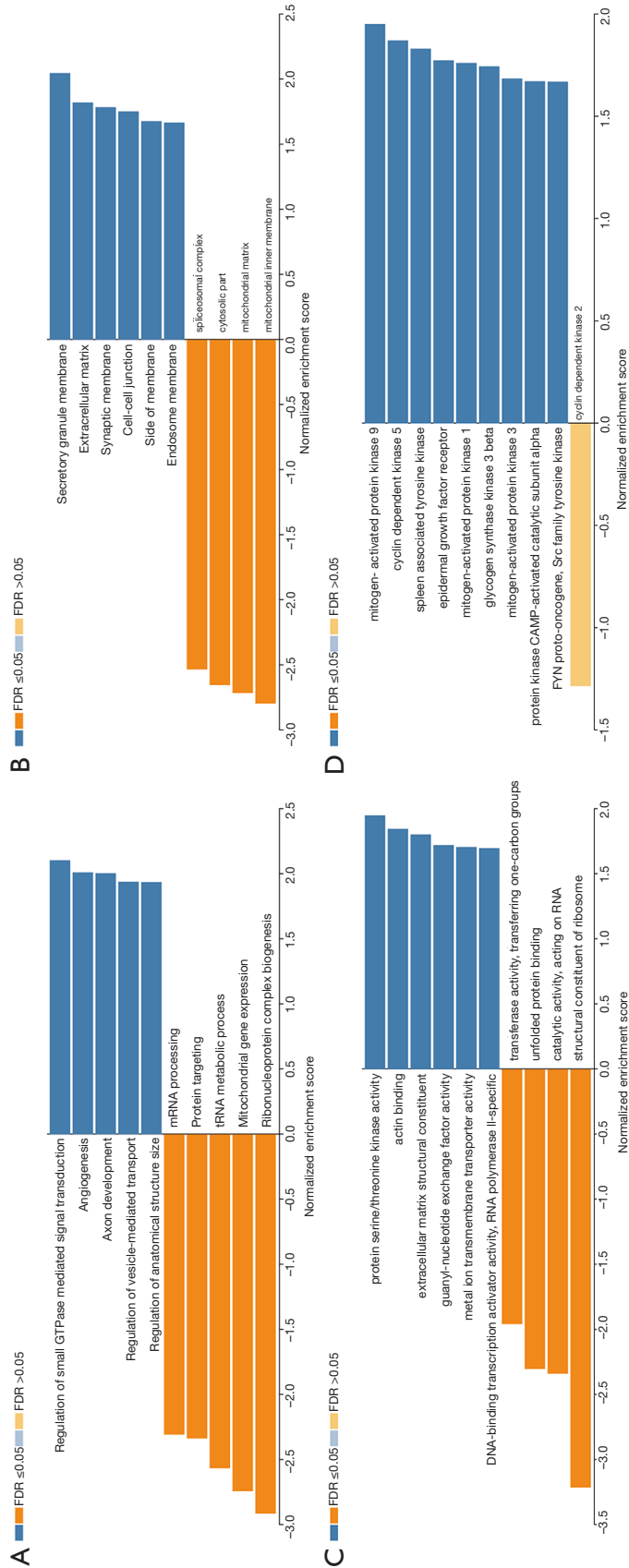


Figure 11 Functions enrichment of TET3 in AML (LinkedOmics) (A) BP of TET3 in AML (B) CC of TET3 in AML (C) MF of TET3 in AML (D) Targets kinases of TET3 in AML. AML, acute myeloid leukemia. TET3, ten-eleven translocation 3; AML, acute myeloid leukemia; BP, biological process; CC, cellular component; MF, molecular functions.

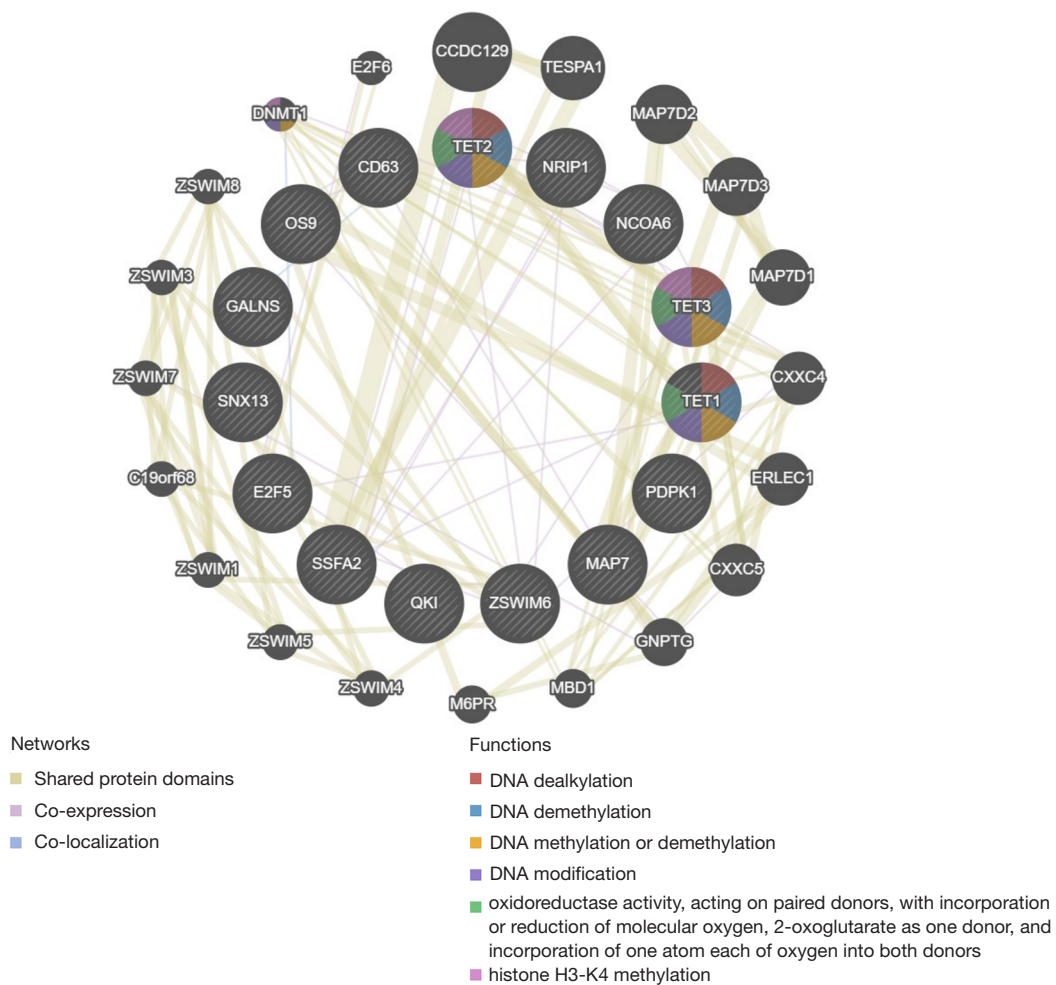


Figure 12 Protein-protein interaction network of TETs (GeneMANIA). PPI network and functional analysis showing the gene set enriched in the target network of TETs. The network edge’s distinct colors indicate the bioinformatics methods applied: physical interactions, co-expression, predicted, co-localization, pathway, genetic interactions, and shared protein domains. The distinct colors for the network nodes show the biological functions of the sets of enrichment genes. TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs); PPI, protein-protein interaction.

TET3, respectively. The network of TETs and miRNAs was constructed by miWalk (Figure 13A). There were 9 miRNAs including hsa-miR-2278, hsa-miR-105-3p, hsa-miR-6882-5p, hsa-miR-4732-3p, hsa-miR-4524a-5p, hsa-miR-6866-5p, hsa-miR-5580-3p, hsa-miR-4692, and hsa-miR-649 that their co-target genes were TETs (Figure 13B).

TF targets of TETs

Further, we used the ChEA3 to analyze the TF targets of TET1, TET2, and TET3. Also, the analysis results of

TFs indicated TFs of TETs were scattered in the nervous system, digestive system, and peripheral blood, further enrichment TFs functions they are the regulators of the immune response, cell differentiation, organ development, DNA transcription, and so on (Figure 14A,B). Further, we construct the encode TFs network and show the top 10 rank TFs, and the top 10 TFs of TETs are JUN, FOSL1, ZNF263, CTCF, MYOD1, TCF12, SP1, JUND, ETS1, and NFIC (Figure 14C,D). From these biological functions of kinase targets and TFs, the TETs may play essential roles in tumorigenesis and prognosis.

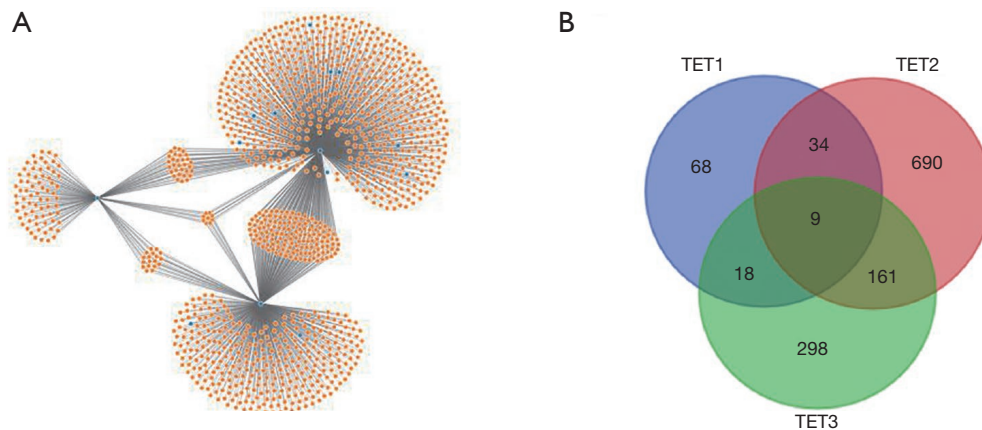


Figure 13 The co-targeted miRNAs of TETs. TET family oxidize 5mCs. TET, ten-eleven translocation; 5mCs, 5-methylcytosines.

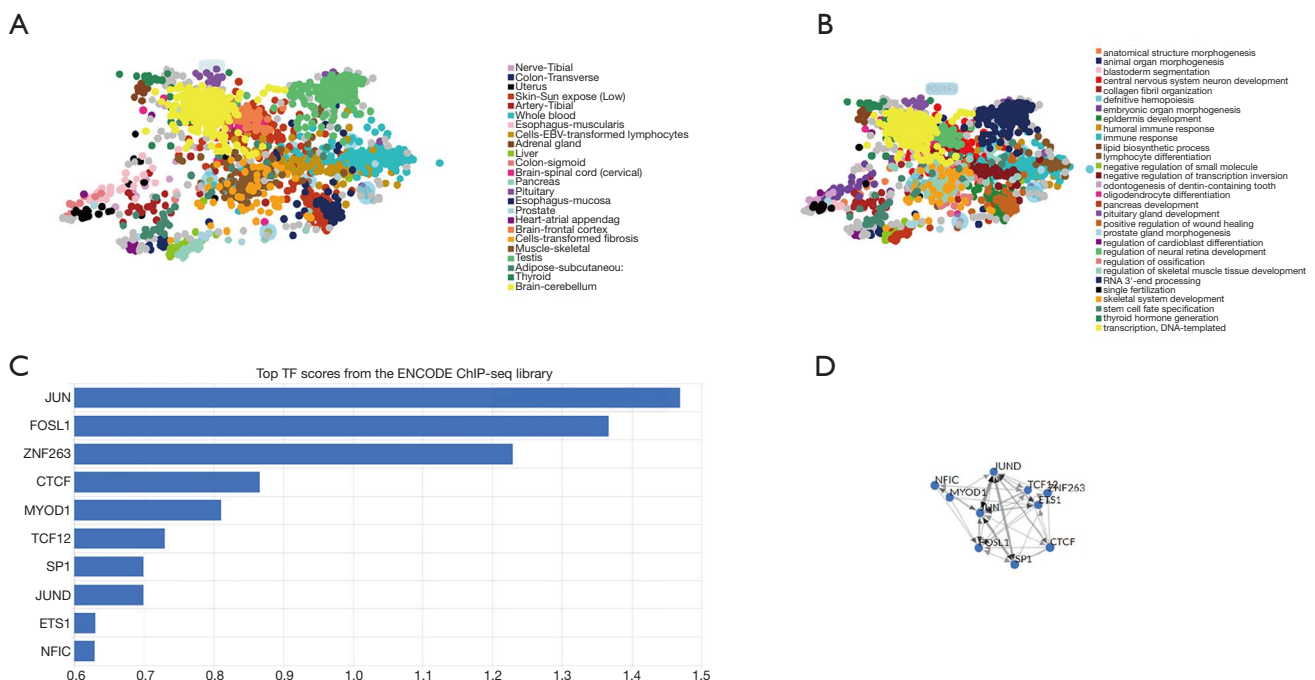


Figure 14 Transcription factors targets of TETs (CHEA3). (A) Tissue distribution of the transcription factors targets for TETs. (B) The biological function of the transcription factors targets for TETs. (C) The top 10 transcription factors targets for TETs. (D) The network of top 10 transcription factors targets for TETs. TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs).

Cancer pathway activity and drugs sensitivity of TETs and correlated genes

To incite the role of defined genes in the cancer pathway, the GSCALite was used to perform the cancer pathway activity, and further investigate whether have drugs target the genes. The cancer pathway activity indicated TETs activated the PI3K/AKT signal pathway, but have

a bidirectional function (activation and inhibition) in the DNA repair response, receptor tyrosine kinase (RTK), and cell cycle. The significantly correlated genes have an essential role in these cancer pathways as well, including GALNS, MAP7, and ZSWIM6 as the activators in TSC/mTOR; NCOA6, NRIP1, SNX13 are the inhibitors of apoptosis, and other genes also as the crucial parts in these

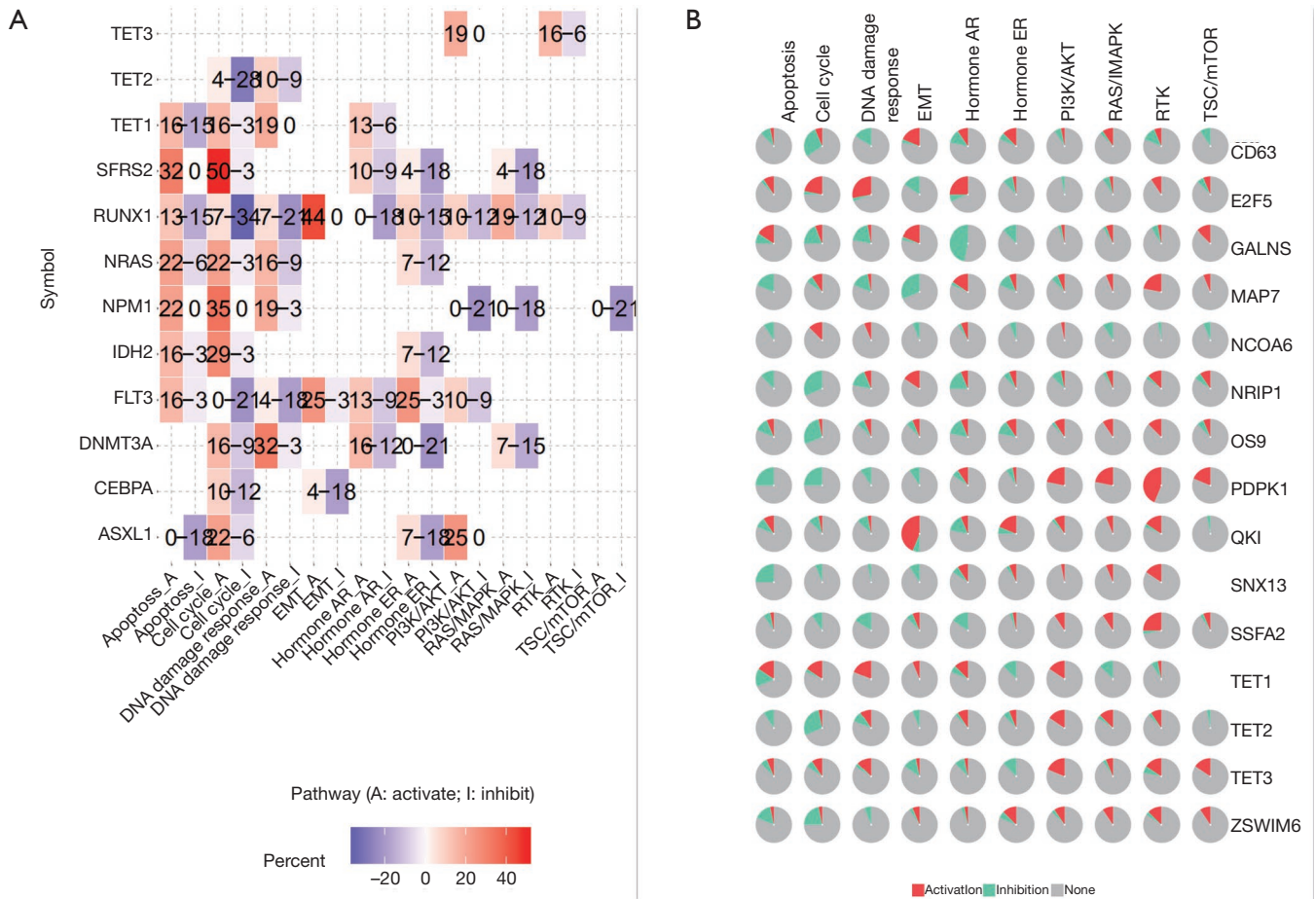


Figure 15 The role of TETs and significantly correlated genes in several cancer pathways (GSCAlite) (A,B) These defined genes serve as the inhibition or activation in several cancer pathways. TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs).

cancer pathways (Figure 15A,B). The final goals of much cancer research are that it can provide useful information for therapeutically cancer patients, so we further analyze the potential drug targets of these defined genes. We used GSCAlite to construct the sensitivity of the drug of these genes that drugs sensitivity data from Cancer Therapeutics Response Portal (CTSP) and Genomics of Drug Sensitivity in Cancer (GDSC). The Spearman correlation represents the gene expression correlates with the drug. The positive correlation means the gene's high expression is resistant to the drug. The results from GDSC shown trametinib was the most resistant for TET1 and TET2, but vorinostat and VNLG/124 were the most sensitive of TET1 and TET2, respectively. CD63 has the largest numbers of resistant drugs, including navitoclax, GSK1070916, belinostat, tubastatin A, vorinostat, phenformin, but

QKI has the most numbers of sensitive drugs including TW37, piperlongumine, bortezomib, and bleomycin (Figure 16). Besides, the results of drug sensitivity from CTSP indicated they have many sensitive drugs for TETs, including alisertib, alvocidib, AT13387, belinostat, BI-2536, ceranib-2, CR-131-B, docetaxel, tivantinib, vincristine, but CD63 has the most significant number of resistant drugs (Figure 17).

The mutated types of TETs in our department

We selected the AML patients in the Department of Hematology, the First Affiliated Hospital of Guangxi Medical University from January 2019 to January 2020. It only had three patients with TET2 mutations—also, the mutated type, including missense and truncating mutation

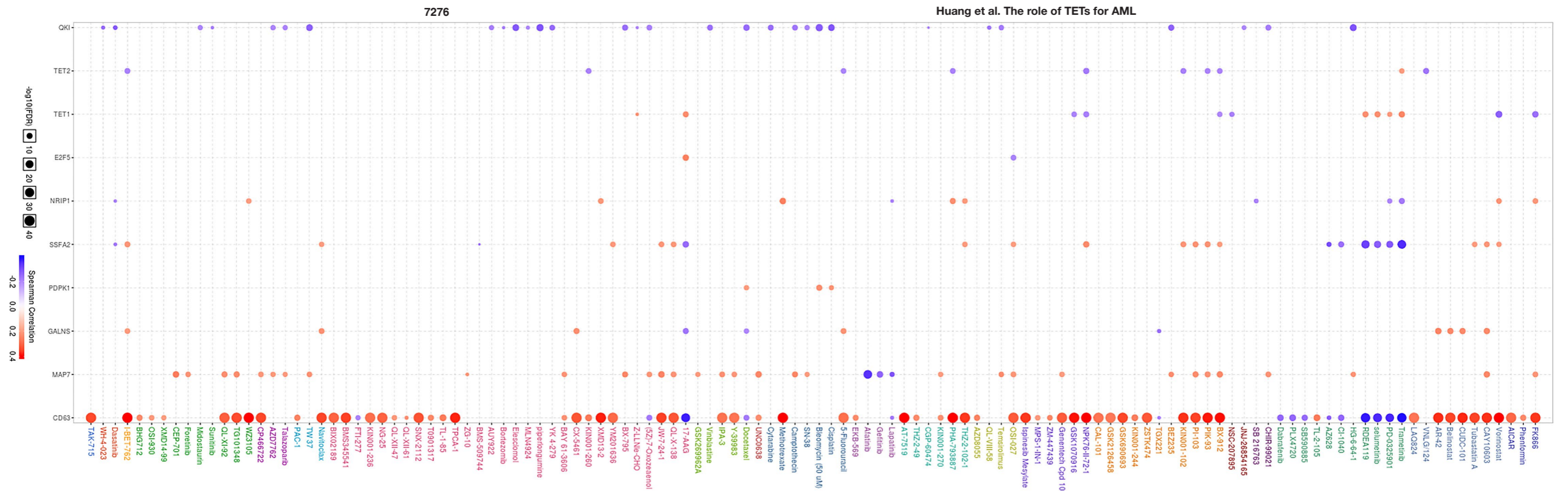


Figure 16 The drug sensitivity of selected genes (GSCAlite) based on GDSC. GDSC, Genomics of Drug Sensitivity in Cancer.

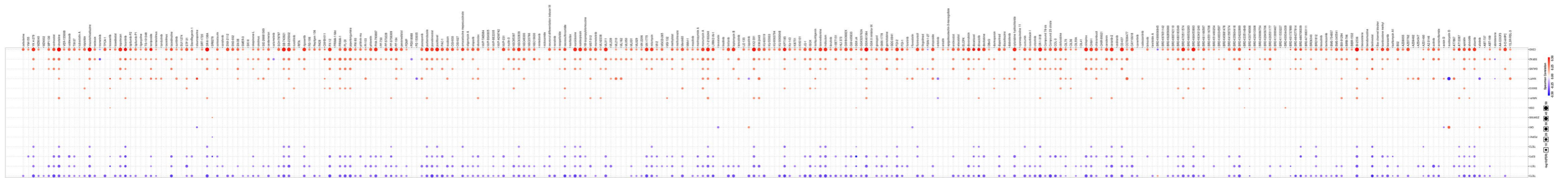


Figure 17 The drug sensitivity of selected genes (GSCAlite) based on CTRP. CTRP, The Cancer Therapeutics Response Portal.

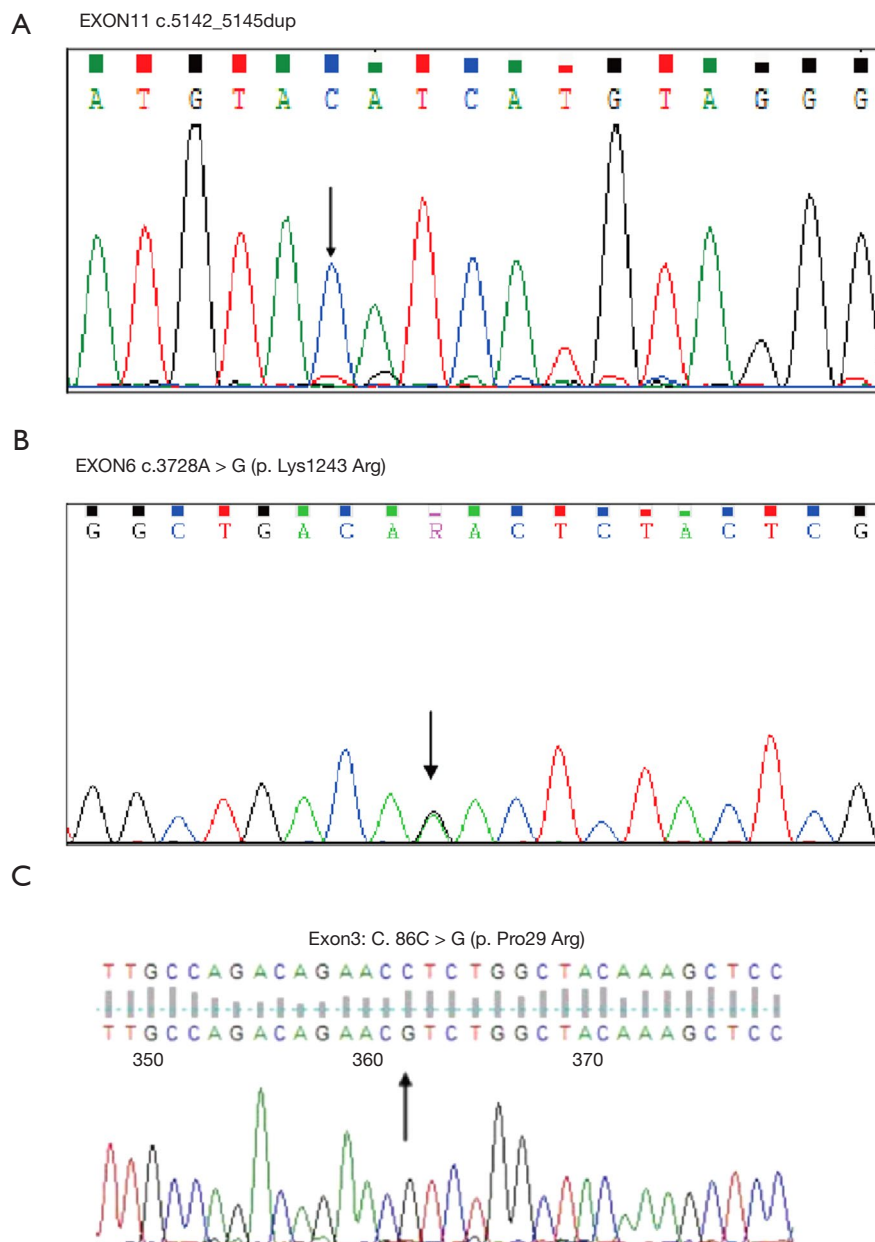


Figure 18 The AML patients with TETs mutations in our department. AML, acute myeloid leukemia; TETs, ten-eleven translocation (TET) family oxidize 5-methylcytosines (5mCs).

(Figure 18A,B,C).

Discussion

Many cancers have aberrant promoter methylation, and it is considered to have a significant relation with tumorigenesis. For recent studies, TETs have seemed crucial regulators of the DNA demethylation pathway in various cancers. The functions of TET2 aberrant mutations in AML have been

well elaborated; the *TET2* gene has a relationship with leukemogenesis, therapy response, and prognosis. However, the other member of TETs' (*TET1* and *TET3*) role in AML remains unclear. Thus, we used bioinformatics to investigate the TETs expression levels, prognostics values, and the potential biological functions in AML.

The aberrant mutations of TETs not only have been studied in hematological cancer but also solid tumors, including breast cancers (11), prostate cancer (31), colon

cancer (32), and melanoma (33). TET1 was an activator in TNBC by upregulating the oncogenic signaling, and the high TET1 expression level may lead the drug resistance by targeting the PI3K-mTOR pathway (11), so the high expression TET1 may indicate the adverse prognosis of TNBC. The research observed that TET1 has a relationship with cancer patients' immunity regulating (34). To date, TET2 has seemed like the one that closest interaction with hematological malignancies. Cimmino *et al.* revealed TET2 as a tumor suppressor in leukemia, the restoration of the TET2 function could inhibit the cancer cell self-renewal and leukemia progression (35).

Further, Xu *et al.* study showed that activated the TET2 may improve the efficacy of immunotherapy efficacy and enhance the immunity of solid cancer patients (36). According to TET3, Cao *et al.* research showed that TET3 high expression indicated the poor prognosis in ovarian cancer patients (13). However, Ye *et al.* revealed TET3 is a tumor suppressor of ovarian cancer through inhibiting epithelial-mesenchymal transition. These studies revealed that the expression of TETs could be detectable in multiple cancer types, and the functions of TETs may be different in different cancer types or even subtype of the same cancer.

In this study, TETs expression levels showed TET1 significantly decreased expression, but the TET2 and TET3 were significant upregulation in AML samples via GEPIA analysis. The Oncomine analysis showed that TET1 and TET2 were significant overexpression in Haferlach Leukemia, but TET3 was a significant low expression in Andersson Leukemia. The TETs subgroups analyses indicated that TETs have significant differential expression levels between different subtypes, age, and other clinical features, these results may be one reason to explain the TETs different expression in various leukemia samples. Some studies approved the expression of TET1 and TET2 can impact the prognosis of AML patients. In our study, the OS results showed that only the TET2 expression level as a significant factor for AML prognosis via UALCAN, and the TET2 high expression leads to positive outcomes of AML patients.

Further, the validation OS results indicated TET2 and TET3 expression levels have a significant relation with AML prognosis, the high expression of TET2 indicating the poor prognosis, but the high expression levels of TET3 may be the protective factor for AML patients. One of the reasons for explaining this is that the validation of AML patients is the cytogenetically normal. These results indicated that the TET2 and TET3 expression levels might

have different functions in different AML subtypes. Few studies investigated the relation between TET3 and AML. So, the underlying mechanisms of TETs in leukemogenesis need to be further explored.

Many studies approve DNA methylation as a critical part of tumorigenesis, progress, and metastasis in solid tumors. Emerging evidence reveals that the downregulation of DNA methylation may be a critical event for leukemogenesis and progression (37). TET one of the main roles is demethylation, but few studies explored the association TET methylation with the prognosis of AML. Herein, this study showed that some located TETs methylations significantly impact AML patients' outcomes.

The regulation of TETs in AML further interested us. We constructed the gene interaction network of TETs and explored their GO enrichment and KEGG pathway analysis. As expected, these genes' functions were primarily related to covalent chromatin modification, demethylation, histone modification, histone H3-K4 methylation, histone lysine methylation, and DNA methylation or demethylation.

These results indicated that the TETs as the vital regulator of DNA transcription and methylation. Both of these biological functions that take part in the tumorigenesis, progress, and metastasis. These results provide more information to explore further the TET's biological role in leukemogenesis and progression.

Dysregulation of miRNAs has been found in multiple cancers; miRNAs can act as promoters or inhibitors of tumors. Hematologic malignancies are no exception; the miRNA's aberrant expression may lead to hematologic cancer, different hematopoietic lineages. Distinctive miRNA profiles were observed in multiple cytogenetic subtypes of AML, and the miRNA expression levels were associated with the prognosis of AML (38). In our study, we finally acquired nine miRNAs that targeted the TETs. Kaymaz *et al.* revealed the has-miR-2278 act as the tumor suppressor; high expression miR-2278 can inhibit leukemic cell proliferation and induce apoptosis (39). To date, there are no studies that explore the other eight miRNAs' role in any disease.

Tumorigenesis involves the changes of multi-omics that include gene aberration, changes of TF and the kinases, and so on. Single gene aberration may not be attributed to cancer. So, we find the significantly correlated genes of TETs, exploring their association and investigating the prognostic role in AML patients, finding the significant TFs and kinases, and incite their functions. For TETs,

there are four significant correlated genes (GALNS, E2F5, MAP7, and NRIP1) that significantly impact the prognosis of AML patients. High expression GALNS indicated the poor outcomes of AML patients, downregulation of E2F5, MAP7, and NRIP1 as the protective factors of AML patients. Ho *et al.* study approved GALNS can as the diagnostics biomarker of multiple cancers (40), but no study of GALNS in AML. Aberrational E2F5 expression has been observed in various cancers. Kothandaraman *et al.* study revealed the E2F5 could act as a biomarker for diagnosing epithelial ovarian cancer (41). A study showed that E2F5 accelerated prostate cancer cell migration and invasion by regulating TFPI2, MMP-2, and MMP-9 (42). Many studies result revealed that targeted E2F5 can inhibit cancer development by various miRNAs (43-45). Research indicated that the E2F5 upregulation in AML, but the E2F5 function in AML, did not further be explored. The BP of E2F5 is related to cell cycle progression and transformation, and acting downstream of the TGF pathway (46). Aberrant, the changes of MAP7 have been observed in multiple cancers as well. Zhang *et al.* revealed MAP7 corporates with RC3H1 as the regulator of cell-cycle progression in cervical cancer cells by activating NF- κ B signaling (47).

Further, Zhang *et al.* study showed MAP7 was a promotor of migration and invasion in cervical cancer by regulating autophagy (48), this result indicated the high expression MAP7 might lead to a poor prognosis for the patients with cervical cancer. According to AML, a study observed that high expression MAP7 indicated the short OS of young patients with CN-AML (49). There was a study that found that suppressed NRIP1 can inhibit breast cancer growth and cell-induced apoptosis (50). Both of the selected correlated significant genes indicate that high expression as an adverse biomarker for AML. However, the role of the selected significant genes in AML and the relation with TETs still unclear. It is needed to be further investigated.

The kinases play critical roles in gene expression and signaling transduction. For TETs, there are so many significant kinases, including CDK2, TGFBR2, CSNK1D, IRAK4, SYK, EGFR, MAP3K4, MYLK3, MYLK4, SGK1, PLK2, ILK, PRKCZ, PRKCE, MAPK10, MYLK2, and MYLK. Their main functions are the regulators of the cell cycle, cell growth, energy metabolism. Among the kinases, only CDK2 has a significant relation with AML. Shao *et al.* research showed that inhibiting the CDK2 can enhance all-trans-retinoic acid efficacy in AML cells (51). Further, Ran *et al.* also revealed CDK2 interacted with

CR6-interacting factor 1 (CRIF1) to inhibit the leukemia cell cycle (52). The function of CDK2 is a regulator of the cell cycle. Dysregulation of the cell cycle is an essential part of tumorigenesis and drug resistance for some cancer types. Some research has already investigated the anticancer agents, targeting therapy the cell cycle (53,54). Regrettably, no study investigates the other kinases in AML. However, most of these kinases work as the promotor in various solid cancers, such as MYLK, promoting the migration and invasion of bladder cancer (55) and gastric cancer (56). PRKC as the accelerator of colon cancer (57). Inhibiting CSNK1D can weaken the migration and invasion ability of TNBC (58), and so on. These studies indicated that most of these kinases might act as an essential role in tumorigenesis and prognosis.

Many factors regulate gene expression; the TFs may be one of the most critical regulatory factors because most regulators impact gene expression by directly or indirectly affecting TFs. Our study used ChEA3 to find the TETs' TFs and annotated them. The GO enrichment of ENCODE TFs revealed they are the regulators of DNA and RNA production. Among the TFs, Jun is the one most significantly related to TETs. Most of these TFs play a key role in different cancer types. Aberrant jun (including c-jun, jun-B, and jun-D) expression has been observed in multiple cancers. Activation of C-Jun can promote liver cancer development and sorafenib resistance (59). The expression of jun has been observed in AML, as well as. Inhibiting the jun can induce the death of leukemia cells by regulating unfolded protein response (60). The ETS1 proto-oncoprotein is a member of the Ets family of transcription factors that share a unique DNA binding domain, the ETS domain. ETS1 regulates the leukemic progenitor cell by effecting the autocrine GM-CSF transcription (61). Sp1 has a significant relation with drug resistance in AML by regulating the ERK-MSK MAPK signaling pathway to impact the survivin expression (62). For this, high expression Sp1 may lead to the poor outcome of AML. The other top TFs do not have any study related to AML, but most of them serve as the promotor of tumorigenesis and progression in multiple solid cancers. These clues indicate that the related TFs of TETs also play a key role in AML as well.

All cancer research aims to provide the underlying mechanisms of tumorigenesis, a biomarker of diagnosis and prognosis, and even find the new therapy target for cancer patients. Thus, patients with cancer can lengthen

their lifespan, even cure the disease. Most cancer-related gene aberrant mutations act as the critical parts of multiple cancers' tumorigenesis and tumor progression via the cancer pathway. In the present study, we used the GSCALite to explore the role of TETs and the significantly correlated genes in cancer pathways, further investigating the potential drug targets of these genes. Among the cancer pathway, the PI3K/AKT signaling pathway has a significant relation with drug resistance in AML (63).

Furthermore, the PI3K/AKT signaling pathway is the regulator of leukemic cell fate that inhibits it can promote leukemic cell apoptosis (64). From these, a study explored the efficacy of targeting the PI3K/Akt signaling pathway in AML (65). Since the TETs are the positive regulator of the PI3K/Akt signaling pathway, we may conclude TETs are the promotor for AML. In AML, the oncofusion proteins, self-renewal activity, and leukemogenic program in hematopoietic cells have a significant association with a dysregulation in the DNA repair pathway, the accumulation of DNA damage that seems like the oncogenic driver of AML. Through these mechanisms, the DNA repair pathway may be the novel potential therapeutic target of AML. Yang *et al.* revealed that combinatorial therapy with AT-101 and IDA could eliminate leukemia stem-like cells via blockage of DNA damage repair (66). In our study, TET1 as activator but TET2 and TET3 have a bidirectional role in the DNA repair pathway.

Further research should focus on the underlying relation TET2 and TET3 with DNA repair response. About 40–60% AML patients with receptor tyrosine kinases (RTKs) or their downstream effector's mutations (67,68). Among the RTKs, FLT3 and C-KIT have been well explored in AML. Most RTKs indicate the poor prognosis of AML. The inhibitors of FLT3 (69) and c-KIT (70) have been introduced to the practice. Both of these TETs acts as the activator or inhibitor in the RTK pathway. So, the TETs may have a different role in different AML types. The TSC/mTOR signaling pathway's role is not unclear in cancer, the primary function of the TSC/mTOR signaling pathway is that it regulates the inflammatory responses (71), but there is not cancer research. Apoptosis is one of the essential types of cancer cell death. Many chemotherapy drugs achieve efficacy by inducing cancer cell apoptosis. The significantly correlated genes that NCOA6, NRIP1, and SNX13 inhibit the apoptosis. These genes may be as possibly a factor for the drug resistance of cancer patients. However, no study investigated the functions of NCOA6 and SNX13 in cancer. For the vital role of TETs and the

significantly correlated genes in the cancer pathway, we use GSCALite to explore them' potential target drugs. Finally, we found the alisertib, alvocidib, AT13387, belinostat, BI-2536, ceranib-2, CR-131-B, docetaxel, tivantinib, and vincristine are the drugs that TETs sensitive. These results can provide useful information for the clinician to choose the drugs for therapy, TETs mutation, or correlate significant AML patients' genes.

Conclusions

In the present study, we comprehensively analyze the TET's expression levels, prognostic role, methylation, biological functions with AML patients. Also, further explored the significant TFs, kinases, cancer pathways, and drug targets of TETs. Our results suggest that TET1, TET2, and TET3 were differentially expressed in AML, CN-AML patients were benefited from overexpression of TET3, but overexpression of TET2 leads to adverse prognosis for CN-AML and other AML patients. The hypermethylation cg24705708, cg05976228, cg21855109, cg25299214, cg17862558, cg13810683, cg1912763, cg15254238, and cg07669489 of TETs were the adverse biomarkers for AML patients. We found that significantly correlated genes, including GALNS, E2F5, MAP7, and NRIP1, can be the novel prognostic biomarker for AML. Regulatory network analysis suggested TETs regulate covalent chromatin modification, demethylation, histone modification, histone H3-K4 methylation, cell cycle, and immune response through pathways involving the significant kinases, TFs, and miRNA. Finally, cancer pathway analysis of TETs activates the PI3K/AKT signal pathway, but have a bidirectional function (activation and inhibition) in the DNA repair response, receptor tyrosine kinase (RTK), and cell cycle. Drug sensitivity analysis results reveal that trametinib was the most resistant for TET1 and TET2, vorinostat, and VNLG/124 were the most sensitive of TET1 and TET2 by GDSC, respectively. Both TETs are sensitive for alisertib, alvocidib, AT13387, belinostat, BI-2536, ceranib-2, CR-131-B, docetaxel, tivantinib, vincristine by CTSP.

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013).

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