Proteomic analysis of cerebrospinal fluid in pediatric acute lymphoblastic leukemia patients: a pilot study

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Purpose: Involvement of central nervous system in acute lymphoblastic leukemia (CNSL) remains one of the major causes of pediatric acute lymphoblastic leukemia (ALL) treatment failure. However, the current understanding of the pathological process of CNSL is still limited. This study aimed to better understand the protein expression in cerebrospinal fluid (CSF) of ALL and discover valuable prognostic biomarkers.

Materials and methods: CSF samples were obtained from ALL patients and healthy controls. Comparative proteomic profiling using label-free liquid chromatography-tandem mass spectrometry was performed to detect differentially expressed proteins.

Results: In the present study, 51 differentially expressed proteins were found. Among them, two core clusters including ten proteins (TIMP1, LGALS3BP, A2M, FN1, AHSG, HRG, ITIH4, CF I, C2, and C4a) might be crucial for tumorigenesis and progression of ALL and can be potentially valuable indicators of CNSL.

Conclusion: These differentially expressed proteins of ALL children with central nervous system involvement and normal children may work as diagnostic and prognostic factors of ALL patients.

Keywords: ALL, central nervous system leukemia, CSF, mass spectrometry, proteomics

Introduction

Acute lymphoblastic leukemia (ALL) is a malignancy derived from the medullary or lymphocyte precursor cells.1 ALL is the most common cancer in children, accounting for 25%–35% of pediatric cancers.2,3 With the understanding of the pathogenesis of ALL and individualized stratified therapy, the clinical efficacy of children with ALL has been significantly improved.4–8 However, central nervous system (CNS) involvement remains one of the major causes of ALL treatment failure.9–11 About 5% of newly diagnosed ALL children are found to be with CNS infiltration.9,12 Without specific treatment for these children, the prevalence rate of central nervous system leukemia (CNSL) will increase to 50%–70% during the whole disease process.13–15 Currently, the mechanism of CNSL is still not clear, and therefore, more molecular level-based research studies are needed to clarify the development and treatment of CNSL.

Proteomic technology enables people to observe the development and progression of diseases dynamically and to carry out quantitative analysis.16–18 Proteomic analysis of biological fluids has been widely used to screen and identify disease markers.19 Recently, various groups have attempted to discover potential protein biomarkers of pediatric leukemia.18–21 One study applied surface-enhanced laser desorption/
ionization time-of-flight mass spectrometry (SELDI-TOF MS) to analyze the proteomes of cell lysates from childhood leukemia cell lines and childhood leukemia bone marrow samples of different subtypes.22 To discover potential protein biomarkers for pediatric ALL, research groups measured serum proteomic profiles between pediatric ALL patients and healthy controls by using SELDI-TOF-MS. In addition, they identified candidate protein biomarkers by using liquid chromatography-tandem mass spectrometry (LC-MS/MS). Notably, the obtained data revealed platelet factor, connective tissue activating peptide III, and two fragments of C3a as potential biomarkers to distinguish pediatric ALL patients from healthy controls and pediatric AML patients.23

Cerebrospinal fluid (CSF) analysis provides important and valuable information about process occurring in the CNS. CSF proteomics has been applied to identify biomarkers of various diseases. In normal physiological conditions, 80% of the protein in the CSF is blood-borne.24,25 The change of protein content in CSF provides a new approach to study malignant hematological disease. Researchers conducted quantitative CSF proteomic analysis in newly diagnosed pediatric ALL and lymphoblastic lymphoma (LL) patients. Notably, in this pilot study, the expression of antithrombin III and plasminogen decreased over time in one child who developed CNS thrombosis, compared to other subjects. The results of this study successfully demonstrated the feasibility of CSF quantitative proteomics in ALL and LL.26

However, to the best of our knowledge, the CSF proteomic profiling of CNSL has not been reported. In this prospective pilot study, we performed quantitative proteomics by using LC-MS/MS to discover differential expression of CSF protein in newly diagnosed pediatric CNSL patients and healthy controls to discover possible prognostic biomarkers.

Materials and methods

Sample collection

The CSF of six pediatric ALL with CNS involvement was obtained from inpatients of Cancer Center, and CSF samples of six controls were obtained from inpatients of pediatrics department, West China Hospital, Sichuan University (2015). All enrolled patients were at the age of 1–11 years. ALL patients with CNS involvement were confirmed by CSF examination. The control group consisted of children with suspected encephalitis, but CSF examination turned out normal. After standard laboratory checks, the CSF samples were centrifugated at 3,000×g at 4°C for 5 minutes to remove the cells, and then stored at −80°C immediately. Our study was approved by the Medical Ethics Committee of West China Hospital, Sichuan University. All patients’ legal guardians signed informed consent, and the study was conducted following the Declaration of Helsinki.

Protein digestion

After thawed at room temperature, a fixed volume of 30 μL CSF was slightly diluted in RIPA buffer (150 mm NaCl, 50 mm Tris–HCl pH 7.61, NP-40, 1% deoxycholic acid) with protease and phosphatase inhibitor on ice for 10 min. Then the CSF samples were centrifugated at 2,000×g at 4°C for 5 minutes. After that, the supernatant was determined by Bradford protein assay. Buffer containing 100 mM NH₄HCO₃ was added to the centrifuged CSF to create an alkaline environment for trypsin digestion. Then 5 mM D,L-dithiothreitol was added to reduce disulfide bonds for 1 hour at 37°C, and 55 mM iodoacetamide (IAA) was used to alkylate the cysteines and reacted in the dark for 45 minutes. Finally, 30 mM L-cysteine was needed for the blockade of redundant IAA, and trypsin (Sequencing Grade Modified Trypsin; Promega Corporation, Fitchburg, WI, USA) was added for the digestion of protein samples overnight at a 1:50 (trypsin to protein) ratio and pH 8.0 at 37°C. The samples were heated to 95°C for inactivating the enzyme, which stopped the digestion reaction. Before the LC-MS/MS analysis, the peptides were desalted by C18 ZipTip (ZTC18S096; EMD Millipore, Billerica, MA, USA).

LC-MS/MS analysis

The procedure for LC-MS/MS analysis in the experiment is similar to that in the previous report, with only minor modifications.27 All peptide samples were lyophilized and suspended in buffer A (2% acetonitrile, 0.1% formic acid) before MS/MS. An EASY-nLC 1000 nanoflow LC instrument coupled to a Q Exactive quadrupole-orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) was adopted for LC-MS/MS analysis. Magic C18 AQ resin (200A, 5 μm; Michrom Bioresources, San Diego, CA, USA) was filled with a 100 μm × 2 cm trap column and a 75 μm × 12 cm analytical column. The mobile phases were composed of buffer A and buffer B (95% ACN, 0.1% formic acid). LC gradient elution conditions initially lasted for 3 minutes for 4% B, from 3 to 43 minutes for 22% B, followed by 8 minutes for 22%–30%, B to 90% for 52–60 minutes and lasted for the last 5 minutes at a flow rate of 300 nL/min. Data-dependent acquisition was conducted in positive ion mode. MS spectra were obtained from 350 to 1,800 m/z with a resolution of 70,000 at m/z =200. The automatic gain control (AGC) value was set to 3e6 with a maximum fill time of 20 ms. The top 20 most intense parent ions were selected.
Hierarchical clustering in Mev software was utilized for presenting global protein expression. Protein–protein interaction (PPI) network analysis was performed using the online Search Tool for the Retrieval of Interacting Genes database (STRING) (http://string-db.org). Interactions with a combined score >0.4 were considered significant. Further, MCODE was used to discover the core module in the PPI network. Finally, GO and REACTOME pathway analyses of proteins in the core module were done online.

Results
CSF protein profiles and differential protein expression
All CSF protein samples were collected from six selected pediatric ALL patients and six healthy pediatric controls in 2015 from pediatrics department of West China Hospital. Age and gender distributions were similar in the two groups. After LC-MS/MS experiments, 455 non-redundant proteins were identified. All the proteins were statistically tested for differential expression. Fifty-one differentially expressed proteins with statistical significance (P<0.05) were found. Among them, 32 proteins were upregulated, and 19 proteins were downregulated. Furthermore, hierarchical clustering was utilized to present protein expression profiles (Figure 1).

![Hierarchical clustering of global proteins.](https://www.dovepress.com/10.2147-ott(s215099.2019.129639)0.jpg)
Patients’ detailed information were shown in Table S1 and all identified proteins by LC-MS/MS analysis were shown in Table S2.

**GO and REACTOME pathway analyses**

To further explore the function of the obtained proteins, GO and REACTOME pathway analyses were performed. A total of 49 gene IDs were found in DAVID. The results of the GO analysis showed that 47 differentially expressed genes were related to 57 terms on the category of “Biological Process.” Among biological processes, the overrepresented terms include cell adhesion (13 genes, \( P = 2.53 \times 10^{-9} \)), negative regulation of endopeptidase activity (10 genes, \( P = 3.41 \times 10^{-11} \)), platelet degranulation (8 genes, \( P = 1.17 \times 10^{-8} \)), signal transduction (8 genes, \( P = 0.0373 \)), receptor-mediated endocytosis (7 genes, \( P = 1.10 \times 05 \)), regulation of complement activation (6 genes, \( P = 1.67 \times 08 \)), CNS development (5 genes, \( P = 3.21 \times 04 \)), regulation of cell growth (4 genes, \( P = 0.0014 \)), complement activation, classical pathway (4 genes, \( P = 0.0025 \)), and axon guidance (4 genes, \( P = 0.0094 \)). The results indicated that platelet degranulation, regulation of immunity, cell growth, and CNS might play an important role in the development of pediatric ALL. Top ten biological processes obtained from GO analysis of all differentially expressed proteins were shown on Figure 2A.

As REACTOME pathway analysis revealed, the differentially expressed genes were associated with a total of 189 pathways. The most representative pathways were immune system (17 genes, \( P = 0.0419 \)), post-translational protein modification (12 genes, \( P = 0.0241 \)), regulation of insulin-like growth factor (IGF) transport and uptake by insulin-like growth factor binding proteins (IGFBPs) (10 genes, \( P = 1.55 \times 10^{-10} \)), hemostasis (10 genes, \( P = 0.0017 \)), innate immune system (10 genes, \( P = 0.042 \)), post-translational protein phosphorylation (9 genes, \( P = 9.17 \times 10^{-10} \)), platelet degranulation (8 genes, \( P = 1.19 \times 7 \)), response to elevated platelet cytosolic Ca\(^{2+} \) (8 genes, \( P = 1.73 \times 7 \)), platelet activation, signaling, and aggregation (8 genes, \( P = 4.15 \times 5 \)). The results indicated that the immune system, cell growth, and platelet function might be essential for the development of pediatric ALL. Top ten pathways obtained from REACTOME pathway analysis of all differentially expressed proteins are shown on Figure 2B.

**PPI network analysis**

To examine the interaction between these differentially expressed genes, we used the STRING database to perform PPI network analysis. The results revealed that an interaction network comprising 31 genes and 64 gene interaction modifications (12 genes, \( P = 0.0094 \)) and axon guidance (4 genes, \( P = 0.0094 \)). The results indicated that platelet degranulation, regulation of immunity, cell growth, and CNS might play an important role in the development of pediatric ALL. Top ten biological processes obtained from GO analysis of all differentially expressed proteins were shown on Figure 2A.

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**PPI network analysis**

To examine the interaction between these differentially expressed genes, we used the STRING database to perform PPI network analysis. The results revealed that an interaction network comprising 31 genes and 64 gene interaction relations was found (Figure 3C).
Further, we used MCODE to discover core module in the PPI network. The results showed that two core clusters including 10 genes were found. Cluster 1 contained seven genes with 21 gene interaction relations (MCODE score = 5.25, node = 7, edges = 21). Proteins involved in cluster 1 were LGALS3BP, TIMP1, A2M, AHSG, FN1, HRG, and ITIH4 (Figure 3A). Cluster 2 included 3 genes with 3 interaction relations (MCODE score = 1.5, node = 3, edges = 3). Proteins involved in cluster 2 were CFI, C2, and C4A (Figure 3B).

**GO function and REACTOME pathway analysis of cluster proteins**

For cluster 1 proteins, GO analysis revealed that seven terms were on the category of “Biological Process” (Figure 2C). Platelet degranulation (7 genes, $P=4.60\times 10^{-14}$), negative regulation of endopeptidase activity (5 genes, $P=3.81\times 10^{-8}$), acute-phase response (3 genes, $P=7.84\times 10^{-5}$), and extracellular matrix disassembly (3 genes, $3.00\times 10^{-4}$) were the most representative terms. REACTOME pathway analysis showed that...

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*Figure 3* Protein–protein interaction (PPI) and MCODE analysis of differentially expressed proteins. (A) Proteins and interactions of cluster 1 identified by MCODE analysis. (B) Proteins and interactions of cluster 2 identified by MCODE analysis. (C) Proteins and interactions of PPI network.
a total of 57 pathways were involved, with platelet degranulation (7 genes, \( P=2.82 \times 10^{-14} \)), response to elevated platelet cytosolic Ca\(^{2+} \) (7 genes, \( P=3.70 \times 10^{-14} \)), platelet activation, signaling and aggregation (7 genes, \( P=6.60 \times 10^{-12} \)), hemostasis (7 genes, \( P=6.27 \times 10^{-9} \)) as the most predominant. Notably, results of cluster 1 proteins were highly consistent with the results of all the differentially expressed proteins. Top 10 pathways obtained from REACTOME pathway analysis of cluster 1 proteins are shown in Figure 2D.

Regarding cluster 2, CF I, C2, and C4A were all related to immune system especially the activity of complement according to GO and REACTOME pathway analysis.

**Discussion**

ALL is the main type of leukemia that causes CNSL.\(^{28} \) The incidence of CNSL in pediatric ALL patients is much higher than that in adults.\(^{28,29} \) CNSL is a special extramedullary infiltrating form of leukemia cells that invade the CNS.\(^{30} \) Leukemia cells may enter the CNS (including the meninges) through direct dissemination or blood circulation.\(^{29,31} \) Few studies have investigated the molecular mechanisms and pathways for CNSL. Currently, a mechanism of ALL cell entry into the CNS by targeting VEGF signaling has been identified by animal model.\(^{32} \) However, the results still need more experiments for validation. In addition, irreversible damages to the CNS remain one of the major causes of ALL treatment failure.\(^{29} \) Once CNS infiltration occurs, it is extremely difficult to control with poor clinical outcomes. Therefore, there is an urgent demand to better understand the pathogenesis of CNSL and identify potential protein markers in the diagnosis and treatment of ALL with CNS infiltration.

CSF is an excellent indicator of brain activity and processes.\(^{33} \) There is also a consensus that CSF is one of the most suitable proximal fluids for analysis of biomarkers in neurological disorders.\(^{34,35} \) It contains normal and disease-related protein components which provide insight into the function of the brain. By analyzing the proteomics of the collected CSF, we were able to identify proteins that were significantly changed in ALL patients with CNS infiltration. In our current study, CSF protein samples were collected from six selected pediatric ALL patients and six control groups. Using LC-MS/MS, 51 proteins were identified to be significantly different between the two groups (\( P<0.05 \)) including 32 proteins upregulated and 19 proteins downregulated. Out of these 51 genes, 49 were found in DAVID. Using GO analysis, 47 differentially expressed genes were related to 57 terms in the biological processes category. The results showed that the differentially expressed proteins were mainly grouped in platelet degranulation, regulation of immunity, cell growth, and CNS.

To better understand the interaction between these differentially expressed proteins, we performed PPI analysis. Two different clusters were identified as shown in Figure 3. These results indicate that the proteins we have identified were not random; rather they have intricate interaction networks. More specifically, cluster 1 proteins including TIMP1, LGALS3BP, A2M, AHSG, FN1, HRG, and ITIH4 were all related to cancer.\(^{36-61} \) Cluster 2 proteins including CF I, C2, and C4A were all related to immune system especially the activity of complement.\(^{62-70} \)

TIMP1 is a glycoprotein that inhibits the proteolytic activity of matrix metalloproteinases in the extracellular space.\(^{36} \) Previous studies have reported that TIMP1 had a positive effect on tumor metastases and progression inhibition.\(^{37,38} \) However, new pieces of evidence have indicated that the overexpression of TIMP1 was associated with poor prognosis in several tumors such as non-Hodgkin lymphomas,\(^{39} \) breast cancer,\(^{40,41} \) colorectal cancer, and papillary thyroid cancer.\(^{41} \) Recently, increased TIMP1 expression has been associated with unfavorable clinical outcomes in children with ALL. The proposed mechanism is the anti-apoptotic function in different cancer cell lines of TIMP1 by activating the FAK/Akt axis and the Bcl-2 survival signaling pathway.\(^{42} \) In the present study, higher levels of TIMP1 expression were also detected. Therefore, we assumed TIMP1 might be a key factor involved in the pathogenesis of CNSL.

LGALS3BP is also known as galectin-3-binding protein, which modulates cell–cell and cell–matrix interactions.\(^{43} \) LGALS3BP has been found to have a complex function in the development of human cancers. High LGALS3BP expression has been implicated with the presence of metastasis in some solid tumors such as non-small-cell lung cancer\(^{43,44} \) and breast cancer.\(^{45} \) However, one study reported that the overexpression of the protein in a series of tumor-derived cell lines reduced tumorigenic potential in athymic mice.\(^{45} \) To the best of our knowledge, literature evidence associated with ALL is limited. Further studies are needed to investigate the role of LGALS3BP in tumors.

As a protease inhibitor and cytokine transporter, alpha-2-macroglobulin (A2M) was associated with inflammation and cancer.\(^{46,47} \) Previous study has reported that A2M was identified as a candidate biomarker for the diagnosis of B-ALL by serum proteomic analysis.\(^{48} \) In addition, altered expression of A2M and FN1 (fibronectin 1) showed
discriminating signatures in breast cancer subtypes.\textsuperscript{49} FN1 is a glycoprotein present in the extracellular matrix and at the cell surface, which plays an important role in cancer progression by promoting cell adhesion and migration.\textsuperscript{50,51} In the present study, A2M and FN1 were also found to be differentially expressed between the two groups which might be of potential diagnostic and prognostic significance for CNSL.

AHSG is a glycoprotein produced by hepatocytes, which has been reported to be downregulated in serum in certain solid tumors, as well as hematological malignancies including acute lymphocytic, acute nonlymphocytic, chronic granulocytic, and chronic myelomonocytic leukemias.\textsuperscript{52,53} The reduction of AHSG is mainly considered to be caused by the infiltration of liver and increased consumption.\textsuperscript{52} Our study found that the concentration of AHSG in CSF decreased in ALL patients as compared to that in healthy controls. In addition, AHSG is found to be involved in brain development, and it may function as a transforming growth factor-\(\beta\) (TGF-\(\beta\)) receptor analog.\textsuperscript{54} It is known that TGF-\(\beta\) regulates a wide range of physiological and pathological processes in the brain including brain inflammatory response and extracellular matrix regulation.\textsuperscript{55} Therefore, the change of AHSG concentration might be a useful laboratory parameter for the assessment of ALL clinical status, especially for CNSL.

Histidine-rich glycoprotein (HRG) is an abundant glycoprotein with multidomain structure.\textsuperscript{56} It modulates various biological processes including cell adhesion, immunity, angiogenesis, and coagulation, most of which involved in tumor metastasis and progression. Studies have suggested that HRG suppressed tumor metastasis and growth by promoting antitumor immune response and tumor vasculature dysfunction.\textsuperscript{57} In vivo, HRG was demonstrated to control tumor metastasis in mouse models of ductal adenocarcinoma, insulinoma, and fibrosarcoma.\textsuperscript{56-58} In the present study, the level of HRG decreased, which is consistent with the lowered expression of HRG in ovarian cancer and liver cancer. Based on this, we wondered HRG might be crucial for CNSL development.

ITIH4 is a glycoprotein secreted by liver in blood with the main function of regulating extracellular matrix.\textsuperscript{59} It has been found to be involved in genesis, metastasis, and development of several solid tumors.\textsuperscript{60} Research groups have observed the dynamic change of serum ITIH4 concentration in the developmental progress of HCC.\textsuperscript{61} It is confirmed that the decreased ITIH4 concentration was related to the genesis of HCC and high HCC risk.\textsuperscript{62} Declined ITIH4 expression was also detected in the present study. Hence, ITIH4 might serve as a warning marker of CNSL genesis.

The complement system consists of more than 30 soluble proteins and membrane binding proteins, which are widely involved in the body’s anti-microbial defense reaction and immune regulation, and can also mediate the damage response of immunopathology.\textsuperscript{62} The complement system has three activation pathways and eventually forms membrane attack complex (MAC). MAC can dissolve tumor cells and therefore inhibit tumor progress.\textsuperscript{63} Complement component 2 (C2) is involved in the classical and lectin pathway of complement activation, and it is required for the formation of MAC.\textsuperscript{64} Complement activation product C4a can stimulate inflammatory reaction to eliminate pathogenic microorganisms.\textsuperscript{65} In the present study, the concentrations of C2 and C4a in CSF were decreased. The underlying mechanism might be the activation of the immune system by tumor cells leading to an increase in C2 and decrease in C4a.

Complement factor I (CFI) is of vital importance in the regulation of complement activation by cleaving C3b and C4b with cofactors complement receptor I (CD35), membrane cofactor protein (CD46), and complement factor H (CFH).\textsuperscript{66} Expression of CFI has been observed in breast cancer cells,\textsuperscript{67} NSCSC cells,\textsuperscript{68} cutaneous squamous cell carcinoma (cSCC) cells,\textsuperscript{69} and glioma cells.\textsuperscript{70} Current evidence suggested that CFI had an important role in promoting cSCC growth and invasion, possibly by using complement inhibitors for immune escape.\textsuperscript{69} In addition, high CFI expression has been reported to be positively correlated with breast cancer tumor size and associated with poor clinical outcomes.\textsuperscript{67} In the present study, we observed an elevated concentration of CFI. Therefore, CFI may participate in the development of CNSL and be a valuable indicator of CNSL diagnosis.

We have a few limitations in our current study. First, we have a relatively small sample size, so results showed in this study may or may not be a good representative of the whole. However, the 12 samples (2 groups of 6) that we obtained were relatively consistent in terms of analysis which gave us some confidence in our results. In addition, our results are consistent with published results in the literature.\textsuperscript{52,55,61,69} Next, the goal of the current study is a pilot study, so the depth of is not great. But publishing our pilot results will allow other groups to analyze some of these specific targets in depth. We are also analyzing some of these proteins more specifically and in detail. Moreover, due to ethical concerns and difficulties in collecting pediatric samples, our study lacked the distinction between ALL with and without CNS involvement. But our study design was in accordance with
previous high-quality studies on CSF research, which was relatively rigorous. Finally, all our samples came from the Chinese Han population, which may or may not be a good representative for other ethnic groups. Due to the limitation to obtaining samples, Chinese Han population were the only samples that were available in our regions.

**Conclusion**

We have used CSF samples from six ALL patients and six normal individuals and discovered significant protein changes which may aid as potential biomarkers in the diagnosis and prognosis of childhood ALL with CNS infiltration. We hope our pilot study can provide an excellent foundation for the future investigators to further analyze these changes in proteins and identify their potential role in the etiology of ALL with CNS infiltration.

**Author contributions**

All authors contributed to data analysis, drafting and revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.

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