Prognostic significance of CNSL at diagnosis of childhood B-cell acute lymphoblastic leukemia: A report from the South China Children’s Leukemia Group

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Objectives: The prognostic significance of acute lymphoblastic leukemia (ALL) patients with central nervous system leukemia (CNSL) at diagnosis is controversial. We aimed to determine the impact of CNSL at diagnosis on the clinical outcomes of childhood B-cell ALL in the South China Children’s Leukemia Group (SCCLG).

Methods: A total of 1,872 childhood patients were recruited for the study between October 2016 and July 2021. The diagnosis of CNSL depends on primary cytological examination of cerebrospinal fluid, clinical manifestations, and imaging manifestations. Patients with CNSL at diagnosis received two additional courses of intrathecal triple injections during induction.
Results: The frequency of CNLS at the diagnosis of B-cell ALL was 3.6%. Patients with CNSL at diagnosis had a significantly higher mean presenting leukocyte count \((P = 0.002)\) and poorer treatment response \((P < 0.05)\) compared with non-CNSL patients. Moreover, CNSL status was associated with worse 3-year event-free survival \((P = 0.030)\) and a higher risk of 3-year cumulative incidence of relapse \((P = 0.008)\), while no impact was observed on 3-year overall survival \((P = 0.837)\). Multivariate analysis revealed that CNSL status at diagnosis was an independent predictor with a higher cumulative incidence of relapse \((\text{hazard ratio} = 2.809, P = 0.016)\).

Conclusion: CNSL status remains an adverse prognostic factor in childhood B-cell ALL, indicating that additional augmentation of CNS-directed therapy is warranted for patients with CNSL at diagnosis.

KEYWORDS
acute lymphoblastic leukemia, central nervous system leukemia, childhood, prognosis, relapse

Introduction

Acute lymphoblastic leukemia (ALL) is a heterogeneous disease that accounts for approximately 80% of all leukemias in children. Based on the immunophenotype, ALL cases can be classified as B-cell or T-cell ALL, with B-cell ALL comprising 85%–90% of newly diagnosed cases (1, 2). With recent advances in risk-directed treatment, 5-year event-free survival (EFS) has exceeded 85% and 5-year overall survival (OS) has surpassed 90% for pediatric patients with B-cell ALL in many clinical trials (3, 4). However, central nervous system involvement remains one of the most important challenges in the treatment of childhood B-cell ALL (5).

The prognostic significance of patients with central nervous system leukemia (CNSL) at diagnosis is controversial. A report from the European Organization for Research and Treatment of Cancer (EORTC) Children’s Leukemia Group study 58881 showed that the presence of initial CNS involvement had no prognostic significance for survival (6). In contrast, another report from the EORTC Children’s Leukemia Group study 58951 showed that CNSL at diagnosis remained an independent adverse prognostic factor in children with ALL (7). Additionally, investigators from the Children’s Oncology Group (COG) reported that CNSL at diagnosis predicted inferior outcome and higher rates of CNS relapse in patients with B-cell ALL (8). The resulting differences may be due to different therapeutic protocols.

The South China Children’s Leukemia Group (SCCLG)-ALL-2016 is a prospective, multi-institutional clinical trial involving 18 major hospitals/medical centers (9, 10). We designed the SCCLG-ALL-2016 collaborative protocol based on the backbones of Berlin-Frankfurt-Münster (BFM) ALL-IC-2009 and Guangdong (GD)-2008-ALL (11, 12). In this study, we aimed to determine the impact of CNSL at diagnosis on the clinical outcomes of childhood B-cell ALL in the SCCLG-ALL-2016 trial.

Materials and methods

Patients

A total of 1,872 pediatric patients who were diagnosed with B-cell ALL between October 2016 and July 2021 were recruited for the study. The inclusion criteria were: age ≤ 18 years, treated according to the SCCLG-ALL-2016 protocol. The exclusion criteria were: mature B-ALL, mixed phenotype leukemia, condition secondary to immunodeficiency disease, presence of Down’s syndrome, non-primary, untreated according to the protocol, and incomplete data for the information of CNS status. The study was conducted in accordance with the principles set down in the Declaration of Helsinki and was approved by the Ethics Committee of Sun Yat-sen Memorial Hospital, Sun Yat-sen University. All patients, or the parents/guardians of patients, provided written informed consent. The trial is registered with the Chinese Clinical Trial Registry (https://www.chictr.org.cn/; ChiCTR2000030357).

CNS status

CNS status was classified into two groups (CNSL and non-CNSL) based on clinical manifestations, imaging (CT/MRI)
findings, cerebrospinal fluid (CSF) white blood cells (WBCs), red blood cells (RBCs), and leukemia blasts. Clinical manifestations of CNSL include general neurologic symptoms, signs of cranial nerve palsy, and spinal involvement. CNSL: CSF WBC >5/μl with blasts in non-traumatic lumbar puncture (TLP, >10 erythrocytes/μl), or TLP with a positive Steinherz/Bleyer result (CSF WBC/RBC ≥2×peripheral blood WBC/RBC), or any clinical or imaging evidence of CNSL. (13, 14). Otherwise, the other status excluding CNSL was defined as non-CNSL. According to the SCCLG-ALL-2016 protocol, CNSL at diagnosis stratified the patients into the intermediate-risk or high-risk groups. Patients with CNSL at diagnosis received two additional courses of intrathecal triple treatment (methotrexate, dexamethasone, and cytarabine) during induction. During continuation therapy, intrathecal therapy was administered according to risk features. Cranial irradiation was omitted for all patients.

**Treatment response**

Early response to treatment was measured as the absolute number of peripheral lymphoblasts at induction therapy. Peripheral blood blasts <1.0 × 10⁹/L on day 8 was considered prednisone good response (PGR), otherwise it was considered prednisone poor response (PPR). Minimal residual disease (MRD) was analyzed at the central protocol laboratory by flow cytometry. MRD positive was defined as ≥0.1% and ≥0.01% on day 15 and day 33, respectively. Complete remission (CR) was defined as less than 5% lymphoblasts in active hematopoietic bone marrow in the absence of clinical evidence of disease at the end of induction. Relapse was defined as the presence of lymphoblasts (≥25%) in the BM or on histological documentation of blasts in extramedullary sites after the achievement of CR.

**Statistical analysis**

In cases with small numbers, chi-squared analysis and Fisher’s exact test were used to compare categorical variables. The non-parametric Mann–Whitney U-test was applied to continuous variables. EFS was calculated from the time between diagnosis and the first event, including induction failure, relapse, death of any cause, or to the point of last follow-up. OS was measured from the start of initial therapy to death from any cause or to the point of last follow-up. The cumulative incidence of relapse (CIR) was defined as the time from the achievement of CR until the occurrence of the first relapse. The Kaplan–Meier model was applied to calculate 3-year EFS and OS, which were compared using the log-rank test. The three-year CIR was estimated by the Fine–Gray subdistribution hazard model, and differences were analyzed by Gray’s test. The competing event for relapse was death during remission. Prognostic factors were examined by multivariate Cox regression analysis. P-values of <0.05 were considered statistically significant (two-tailed testing). The data were analyzed with the Statistical Package for the Social Sciences (SPSS®) version 24.0 (IBM Corporation, Armonk, NY, USA).

**Results**

**Patients’ characteristics**

Overall, among 1,872 patients with childhood B-cell ALL, 68 (3.6%) demonstrated CNSL status at diagnosis. Neurological symptoms such as headaches, vomiting, and convulsions were observed in eight patients among the patients with CNSL. However, the clinical features of CNSL were not associated with imaging or cytology positivity in our study. As shown in Table 1, there were no differences in gender and age distribution between CNSL and non-CNSL patients. Nevertheless, compared with non-CNSL, childhood B-cell ALL with CNSL at diagnosis had a significantly higher median leukocyte count (22.8 × 10⁹/L vs. 9.0 × 10⁹/L, P = 0.002). Among all the patients with CNSL at diagnosis, 24 cases (35.3%) and 44 cases (64.7%) were stratified into the intermediate-risk and high-risk groups, respectively. Additionally, no significant differences were found in the frequencies of fusion genes including BCR/ABL, MLL-r, ETV6/RUNX1, and E2A/PBX1 between patients with CNSL and non-CNSL. In total, 27 (1.4%) patients received hematopoietic stem cell transplantation in our study. However, the percentage of hematopoietic stem cell transplantation was equally distributed between the CNSL group and the non-CNSL group (P = 0.257).

**Clinical outcomes**

Treatment response was measured by the prednisone response on day 8, as well as MRD on both days 15 and 33. The rate of PGR for patients with CNSL and non-CNSL was 83.6% and 92.8%, respectively, and the difference was statistically significant (P = 0.005). Moreover, CNSL at diagnosis was associated with a higher rate of MRD positive on day 15 (85.3% vs. 73.8%, P = 0.033) and day 33 (84.8% vs. 73.8%, P = 0.044), respectively.

We then evaluated the survival data for all childhood patients. As shown in Figures 1A, B, patients with CNSL at diagnosis had a significantly worse 3-year EFS (83.2 ± 5.2%) than non-CNSL patients (91.3 ± 0.8%, P = 0.030). However, we found that CNSL at diagnosis yielded no significant
differences in 3-year OS (94.3 ± 3.4% vs. 95.1 ± 0.6%, \(P = 0.836\)) between these two groups. We further evaluated the prognostic impact of CNSL and the risk group in patients by stratification analysis. In terms of 3-year EFS (Figure 1C), CNSL had an adverse impact trend on the survival of patients restricted to the intermediate-risk group (88.2 ± 5.6% vs. 93.5 ± 0.9%, \(P = 0.256\)) or the high-risk group (73.4 ± 10.8% vs. 80.6 ± 2.5%, \(P = 0.345\)), but the difference was not statistically significant. In terms of 3-year OS (Figure 1C), we found that CNSL had no significant impact in the survival of patients restricted to the intermediate-risk group (\(P = 0.751\)) or the high-risk group (\(P = 0.905\)).

**TABLE 1** Characteristics of childhood B-ALL with or without CNSL at diagnosis.

| Characteristics                        | Total 1,872 | Non-CNSL (n = 1,804) | CNSL (n = 68) | P-value |
|----------------------------------------|-------------|----------------------|--------------|---------|
| Gender, n (%)                          |             |                      |              |         |
| Male                                   | 1,073 (57.3%) | 1,033 (57.3%) | 40 (58.8%) | 0.798   |
| Female                                 | 799 (42.7%)  | 771 (42.7%) | 28 (41.2%) |         |
| Age (y), median (range)                |             |                      |              |         |
| Age (y), n (%)                         | 4.4 (0.5–17.4) | 4.4 (0.5–17.4) | 4.9 (0.6–14.1) | 0.317   |
| <10                                    | 1,618 (86.4%) | 1,562 (86.6%) | 56 (82.4%) |         |
| ≥10                                    | 254 (13.6%)  | 242 (13.4%) | 12 (17.6%) |         |
| Risk group, n (%)                      |             |                      |              | <0.001  |
| Low risk                               | 454 (24.3%)  | 454 (25.2%) | 0 (0%)     |         |
| Intermediate risk                      | 994 (53.1%)  | 950 (52.7%) | 44 (64.7%) |         |
| High risk                              | 424 (22.6%)  | 400 (22.2%) | 24 (35.3%) |         |
| WBC (>10^9/L), median (range)          |             |                      |              | 0.002   |
| WBC (>10^9/L), n (%)                   | 9.3 (0.16–1095) | 9.0 (0.16–1095) | 22.8 (0.93–618.7) | 0.001   |
| <50                                    | 1,540 (82.3%) | 1,494 (82.8%) | 46 (67.6%) |         |
| ≥50                                    | 332 (17.7%)  | 310 (17.2%) | 22 (32.4%) |         |
| ALL, acute lymphoblastic leukemia; CNS, central nervous system; CNSL, central nervous system leukemia; EFS, event-free survival; HSCT, hematopoietic stem cell transplantation; MRD, minimal residual disease; WBC, white blood cell.
CIR and distribution of relapses

The 3-year CIR for patients with CNSL and non-CNSL was 12.8 ± 4.9% and 4.7 ± 0.6%, respectively, and the difference was statistically significant (P = 0.008) (Figure 2A). By stratification analysis (Figure 2B), we found that patients with CNSL at diagnosis had a high-risk trend of 3-year CIR in the intermediate-risk group (9.6 ± 5.3% vs. 3.2 ± 0.7%, P = 0.060). However, when restricted to the high-risk group, no significant difference was found between CNSL and non-CNSL patients in terms of 3-year CIR (19.9 ± 10.6% vs. 11.4 ± 2.2%, P = 0.259).

We further investigated the distribution of relapses according to CNS status at diagnosis. As shown in Table 2, patients with CNSL at diagnosis had a higher risk of relapse (8.8% vs. 3.5%, P = 0.036). Notably, CNSL at diagnosis was associated with a higher risk of relapse with CNS involvement (4.4% vs. 0.6%, P = 0.010) and isolated CNS relapse (2.9% vs. 0.4%, P = 0.040), respectively.

Multivariate analyses

Table 3 summarizes the results of multivariate analyses for the complete cohort of patients with childhood ALL. In the multivariable survival analysis for EFS, OS, and CIR, we included CNS status with other risk factors in the Cox model, including sex, age, WBC, prednisone response on day 8, and MRD on both days 15 and 33 as covariates. Our results showed that...
CNSL at diagnosis did not reach significance either for EFS ($P = 0.217$) or OS ($P = 0.323$). However, we identified that CNSL status was an independent predictor with a higher CIR (hazard ratio = 2.809, $P = 0.016$). In this model, age ($\geq 10$-year-old) and WBC ($\geq 50 \times 10^9/L$) were associated with worse EFS, while age ($\geq 10$-year-old) and sex (male) were associated with higher CIR.

### Discussion

CNSL is caused by the infiltration of leukemia cells into the CNS. The diagnosis of CNSL was made by primary cytological examination of the CSF, clinical manifestations, and imaging manifestations. In our SCCIG cohort, the frequency of CNLS at diagnosis among 1,872 patients with childhood B-cell ALL was 3.6%. This result was higher than that of other childhood B-cell ALL reports (range 1.3%–1.7%) (6, 7). This difference may be explained in part by the different classification of CNS status and various study populations. Moreover, we found that childhood B-cell ALL with CNSL at diagnosis had a significantly higher median leukocyte count. However, in our study, no association was found between the CNS status and the frequency of fusion genes, including $BCR/ABL$, $MLL-r$, $ETV6/RUNX1$, and $E2A/PBX1$. Currently, hyperleukocytosis upon diagnosis is considered a risk factor for CNS involvement in B-cell ALL (15). Nevertheless, some reports have shown that $BCR/ABL$, $MLL-r$, and E2A/PBX1 are associated with a higher incidence of CNSL in B-cell ALL (16–18).

There has been controversy about the significance of CNS involvement in childhood ALL. A report from the Nordic Society of Paediatric Haematology and Oncology (NOPHO) showed that the post induction bone marrow MRD did not differ between patients with CNS involvement and those without CNS involvement. Moreover, the 12-year EFS for patients with CNSL
at diagnosis did not differ from that in patients with non-CNSL (19). Another report from the Dutch Childhood Oncology Group also showed no significant difference in 10-year EFS between patients with CNSL and non-CNSL at diagnosis (20). These findings were in accordance with the results of the EORTC Children’s Leukemia Group study 58881 (6). However, a study from the ALL-BFM-95 trial showed that patients with CNSL at diagnosis had the worst prognosis, with a 5-year EFS estimate of 50%, and these patients had a higher risk of CNS relapses (21). Another trial from the EORTC Children’s Leukemia Group study 58881 showed that the prognosis was worse for patients with CNSL in terms of 3-year survival, while it was not at long-term follow-up (6). Thus, a long-term follow-up study is needed to further address the prognostic significance of CNSL at diagnosis. Second, we did not classify CNSL at diagnosis into different groups, such as abnormal cytological examination of CSF or imaging manifestations. However, Levinsen et al. demonstrated that the long-term EFS for patients with leukemia mass on neuroimaging did not differ from CNSL patients with negative scans (19). Third, our study was a lack of data regarding the immunophenotype of CNS status. Recently, it has been demonstrated that flowcytometric evaluation of CSF in childhood ALL better identifies CNS involvement than conventional cytomorphology. Further studies with multiparametric flow cytometry are warranted to obtain higher quality evidence.

In summary, we analyzed the prognostic impact of CNSL for the diagnosis of childhood B-cell ALL in the SCCLG trial. Our findings showed that patients with CNSL at diagnosis had an
independent adverse prognostic factor for CNS relapse. Additional augmentation of CNS-directed therapy is warranted for managing CNS involvement.

Data availability statement

The original contributions presented in the study are included in the article/supplementary material. Further inquiries can be directed to the corresponding author.

Ethics statement

The studies involving human participants were reviewed and approved by the Ethics Committee of Sun Yat-sen Memorial Hospital, Sun Yat-sen University. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

Author contributions

L-HX and YL participated in project design, analysis, interpretation, and manuscript drafting. XG, NL, L-HY, H-RM, W-QW, L-BH, and M-CZ obtained and assembled data. CT, H-QC, Q-WC, X-JL, Z-JZ, R-YL, Q-RL, B-YW, L-NW, X-LK, G-HC, and J-PF analyzed and interpreted the data. All authors contributed to the article and approved the submitted version.

References

1. Teachey DT, Pui CH. Comparative features and outcomes between pediatric T-cell and B-cell acute lymphoblastic leukaemia. Lancet Oncol (2019) 20(5):e142–54. doi: 10.1016/S1470-2045(19)30031-2
2. Hunger SP, Mullighan CG. Acute lymphoblastic leukaemia in children. N Engl J Med (2015) 373(16):1541–52. doi: 10.1056/NEJMra1406972
3. Pui CH. Precision medicine in acute lymphoblastic leukemia. Front Med (2020) 14(6):689–700. doi: 10.1007/s11684-020-0759-8
4. Inaba H, Pui CH. Advances in the diagnosis and treatment of pediatric acute lymphoblastic leukaemia. J Clin Med (2021) 10(9):1926. doi: 10.3390/jcm10091926
5. Inaba H, Mullighan CG. Pediatric acute lymphoblastic leukaemia. Haematologica (2020) 105(11):2524–39. doi: 10.3324/haematol.2020.247031
6. Sirvent N, Suciu S, Riafland X, Millot F, Benoit Y, Plantaz D, et al. Prognostic significance of the initial cerebro-spinal fluid (CSF) involvement of children with acute lymphoblastic leukaemia (ALL) treated without cranial irradiation: results of European organization for research and treatment of cancer (EORTC) children leukaemia group study 58881. Eur J Cancer (2011) 47(2):239–47. doi: 10.1016/j.ejca.2010.01.019
7. Sirvent N, Suciu S, De Moerloose B, Ferster A, Mazingue F, Plat G, et al. CNS-3 status remains an independent adverse prognosis factor in children with acute lymphoblastic leukaemia (ALL) treated without cranial irradiation: Results of EORTC children leukaemia group study 58951. Arch Pediatr (2021) 28(5):411–6. doi: 10.1684/arcped.2021.004.009
8. Winick N, Devidas M, Chen S, Maloney K, Larson E, Mattano L, et al. Impact of initial CSF findings on outcome among patients with national cancer institute standard- and high-risk b-cell acute lymphoblastic leukemia. A report from the children’s oncology group. J Clin Oncol (2017) 35(22):2527–34. doi: 10.1200/JCO.2016.71.4774
9. Qiu KY, Xu HG, Luo XQ, Mai HR, Liao N, Yang LH, et al. Prognostic value and outcome for ETV6/RUNX1-positive pediatric acute lymphoblastic leukemia: A report from the south China children’s leukaemia group. Front Oncol (2021) 11:797194. doi: 10.3389/fonc.2021.797194
10. He YY, Wu XL, Zhou DH, Yang LH, Mai HR, Wan WQ, et al. A nomogram for predicting event-free survival in childhood acute lymphoblastic leukaemia: A multicenter retrospective study. Front Oncol (2022) 12:854798. doi: 10.3389/fonc.2022.854798
11. Radu LE, Colita A, Pasca S, Tomulesca C, Popa C, Serban C, et al. Day 15 and day 33 minimal residual disease assessment for acute lymphoblastic leukemia patients treated according to the BFM ALL IC 2009 protocol: Single-center experience of 133 cases. Front Oncol (2020) 10:923. doi: 10.3389/fonc.2020.00923
12. Li XY, Li QJ, Luo XQ, Wu XD, Sun X, Xu HG, et al. Reduced intensity of early intensification does not increase the risk of relapse in children with standard risk acute lymphoblastic leukemia - a multi-centric clinical study of GD-2008-ALL protocol. BMC Cancer (2021) 21(1):59. doi: 10.1186/s12885-020-07752-x
13. Larson RA. Managing CNS disease in adults with acute lymphoblastic leukemia. Leuk Lymphoma (2018) 59(1):3–13. doi: 10.1080/10428194.2017.1326597
14. Jastaniah W, Eliman N, Abdalla K, Khattab TM, Felimban S, Abrar MB. Does the early intensification of intrathecal therapy improve outcomes in pediatric acute lymphoblastic leukemia patients with CNS2/TLP+ status at diagnosis? Hematology (2015) 20(10):561–6. doi: 10.1179/1607845415Y.0000000020

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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15. Frishman-Levy I, Izraeli S. Advances in understanding the pathogenesis of CNS acute lymphoblastic leukaemia and potential for therapy. Br J Haematol (2017) 176(2):157–67. doi: 10.1111/bjh.14411

16. Sanchez R, Ayala R, Alonso RA, Martinez MP, Ribera J, Garcia O, et al. Clinical characteristics of patients with central nervous system relapse in BCR-ABL1-positive acute lymphoblastic leukaemia: the importance of characterizing ABL1 mutations in cerebrospinal fluid. Ann Hematol (2017) 96(7):1069–75. doi: 10.1007/s00277-017-3002-1

17. Shen S, Chen X, Cai J, Yu J, Gao J, Hu S, et al. Effect of dasatinib vs imatinib in the treatment of pediatric Philadelphia chromosome-positive acute lymphoblastic leukaemia: A randomized clinical trial. JAMA Oncol (2020) 6(3):358–66. doi: 10.1001/jamaoncol.2019.5868

18. Winters AC, Bernt KM. MLL-rearranged leukemias—an update on science and clinical approaches. Front Pediatr (2017) 5:4. doi:10.3389/fped.2017.00004

19. Levinsen M, Taskinen M, Abrahamsson J, Forestier E, Frandsen TL, Harila-Friis F, et al. Diagnostic cerebrospinal fluid examination in children with acute lymphoblastic leukaemia without pleiocytosis and normal cerebrospinal fluid. J Clin Oncol (2003) 21(2):184–8. doi:10.1200/JCO.2003.04.096

20. Dutch Childhood Oncology G, te Loo DM, Kamps WA, van der Does-van den Berg A, van Wering ER, de Graaf SS. Prognostic significance of blasts in the cerebrospinal fluid without pleiocytosis or a traumatic lumbar puncture in children with acute lymphoblastic leukaemia: experience of the Dutch childhood oncology group. J Clin Oncol (2006) 24(15):2332–6. doi: 10.1200/JCO.2005.03.9727

21. Burger B, Zimmermann M, Mann G, Kuhl J, Loning L, Rehm H, et al. Diagnostic cerebrospinal fluid examination in children with acute lymphoblastic leukaemia: significance of low leukocyte counts with blasts or traumatic lumbar puncture. J Clin Oncol (2003) 21(2):184–8. doi:10.1200/JCO.2003.04.096

22. Pui CH, Howard SC. Current management and challenges of malignant disease in the CNS in paediatric leukaemia. Lancet Oncol (2008) 9(3):257–68. doi:10.1016/S1470-2247(08)70070-6

23. Vora A, Andreano A, Pui CH, Hunger SP, Schrappe M, Moeriicke A, et al. Influence of cranial radiotherapy on outcome in children with acute lymphoblastic leukaemia treated with contemporary therapy. J Clin Oncol (2016) 34(9):919–26. doi: 10.1200/JCO.2015.64.2850

24. Zajac-Sypucha O, Pawlak M, Karmelita-Katuska K, Pilarsczyk J, Jonczyk-Potoczna K, Przezioara A, et al. Anti-leukemic treatment-induced neurotoxicity in long-term survivors of childhood acute lymphoblastic leukaemia: impact of reduced central nervous system radiotherapy and intermediate- to high-dose methotrexate. Leuk Lymphoma (2018) 59(10):2342–51. doi: 10.1080/10428194.2018.1434879

25. Lenk L, Alsdadj A, Schewe DM. Involvement of the central nervous system in acute lymphoblastic leukaemia: opinions on molecular mechanisms and clinical implications based on recent data. Cancer Metastasis Rev (2020) 39(1):173–87. doi:10.1007/s10555-020-09848-z

26. Hiniker SM, Agarwal R, Modlin LA, Gray CC, Harris JP, Million L, et al. Survival and neurocognitive outcomes after cranial or craniospinal irradiation plus total-body irradiation before stem cell transplantation in pediatric leukemia patients with central nervous system involvement. Int J Radiat Oncol Biol Phys (2014) 89(1):67–74. doi:10.1016/j.ijrobp.2014.01.056

27. Jha S, Pei D, Choi J, Cheng C, Sandlund JT, Coustan-Smith E, et al. Improved CNS control of childhood acute lymphoblastic leukemia without cranial irradiation: St Jude total therapy study 16. J Clin Oncol (2019) 37(35):3377–91. doi:10.1200/JCO.19.01692

28. Bhoywani D, Sahin ND, Pei D, Yang JJ, Khan RB, Panetta JC, et al. Methyltherate-induced neurotoxicity and leukoencephalopathy in childhood acute lymphoblastic leukaemia. J Clin Oncol (2014) 32(9):949–59. doi:10.1200/JCO.2013.53.0808

29. Jabbour E, O’Brien S, Kantarjian H, Garcia-Manero G, Ferrajoli A, Ravandi F, et al. Neurologic complications associated with intrathecal liposomal cytarabine given prophylactically in combination with high-dose methotrexate and cytarabine to patients with acute lymphocytic leukemia. Blood (2007) 109(8):3214–8. doi:10.1182/blood-2006-08-043646

30. Shaper A, Stark R, Kornreich L, Cohen JJ, Arivner S, Steinmetz A, et al. Methyltherate treatment protocols and the central nervous system: significant care with significant neurotoxicity. J Child Neurol (2000) 15(9):573–80. doi:10.1177/088307380015009002

31. Qi Y, Zhao M, Hu Y, Wang Y, Li P, Gao J, et al. Efficacy and safety of CD19-specific CAR T-cell-based therapy in b-cell acute lymphoblastic leukemia patients with CNSL. Blood (2022) 139(23):3376–86. doi:10.1182/blood.2021013733

32. Tang J, Yu J, Cai J, Zhang L, Hu S, Gao J, et al. Prognostic factors for CNS control in children with acute lymphoblastic leukemia treated without cranial irradiation. Blood (2021) 137(4):311–43. doi:10.1182/blood.2020014338

33. Zhou F, Wen Y, Jin R, Chen H. New attempts for central nervous infiltration of pediatric acute lymphoblastic leukemia. Cancer Metastasis Rev (2019) 38(4):657–71. doi:10.1007/s10555-019-09827-z

34. Paul S, Short NJ. Central nervous system involvement in adults with acute leukaemia: Diagnosis, prevention, and management. Curr Oncol Rep (2022) 24(4):427–36. doi:10.1007/s11912-022-01220-4

35. McNeer JL, Schmiegelow K. Management of CNS disease in pediatric acute lymphoblastic leukaemia. Curr Hematol Malig Rep (2022) 17(1):1–14. doi:10.1007/s11899-021-00640-6