

CASE REPORT

Refractory Chronic Lymphocytic Leukemia with Central Nervous System Involvement: A Case Report with Literature Review

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Abstract: There have been few reports on central nervous system (CNS) involvement in chronic lymphocytic leukemia (CLL). This is an extremely rare disease with poor prognosis, owing to resistance to various treatments. We describe a 33-year-old man with intractable CLL with CNS involvement. He was diagnosed with CLL, with diplopia as the first manifestation. Magnetic resonance imaging revealed a contrast-enhancing tumor in the right temporal lobe, which was diagnosed as CNS involvement in CLL on brain biopsy. High-dose methotrexate therapy was ineffective for this lesion, which was also resistant to subsequent whole-brain irradiation, treatment with fludarabine-cyclophosphamide-rituximab chemoimmunotherapy, and ibrutinib administration. Because no standard protocol exists for CLL with CNS involvement, it is important to accumulate case data to verify the choice of new drugs for administration at an early stage. Therefore, we also conducted a literature review of 50 case reports of CNS lesions in the last 10 years to consider the pathophysiology, diagnosis, and treatment of CNS involvement in CLL. The possibility of new therapeutic agents, eg, ibrutinib and venetoclax, or a combination of these agents and methotrexate, can be envisioned as a treatment strategy for CLL with CNS involvement.

Keywords: chronic lymphocytic leukemia, central nervous system involvement, literature review

Introduction

Chronic lymphocytic leukemia (CLL) is the most frequent adult leukemia in the US and Europe, but is a rare disease in Japan, with a frequency 10% that in the US. The disease typically occurs in older patients, and the median age at diagnosis is 72 years.² Generally, CLL progresses slowly, but some cases progress rapidly and aggressively.3 Furthermore, CLL has a highly variable clinical course, and neurological complications arising from direct leukemic involvement in the central nervous system (CNS) are reported in only 1% of patients with CLL. 4,5 Here, we present a rare case of a young CLL patient with CNS involvement that was resistant to various therapies. CLL treatment has improved considerably in the last decade; however, it remains unclear which the best treatment for CNS involvement in CLL is. Therefore, in this case report, we also conducted a comprehensive literature review of 50 case reports with CNS involvement in the last 10 years in which the clinical course was described.

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Case Presentation

A 33-year-old man with diplopia was referred to our hospital. He had a 9 month history of asymptomatic revised Rai low- and Binet A-stage CLL that had been diagnosed owing to an increase in lymphocyte count at a medical checkup, but he had not come to the hospital at his own discretion. Thereafter, he developed diplopia and was referred to neurosurgery by an ophthalmologist. Except for double vision and intracranial hypertension-related headaches, the neurological examination was unremarkable, and he had no other symptoms or lymph-node swelling. Magnetic resonance imaging (MRI) revealed a 5×3.5 cm nonuniformly contrasted mass in the right temporal lobe that appeared hypointense on T_1 -weighted and hyperintense on T_2 -weighted images (Figure 1A). In this case, because there was a risk of cerebral hernia owing to a bulky CNS lesion, lumbar puncture could not be performed.

A diagnostic cranioscopic biopsy was performed, which revealed infiltration of small monoclonal lymphocytes with expression of CD5, CD20 (Figure 2), and CD79A, but without CD10, CD23, cyclin D1, or evidence of transformation. Similarly, his blood showed CLL-cell clonality, with

expression of CD5, CD19, CD20 (dim), CD22, and cellsurface Ig, but no expression of CD10, CD23, or IgH-BCL1 on fluorescence in situ hybridization. Bone marrow (BM) specimens revealed 96.6% of lymphocytes had the same flow-cytometry appearance as peripheral blood (PB). BM lymphocytes had a normal karyotype without poor prognostic factors, deletion 17p, deletion 11q, or transformation (Figure 3), which was compatible with a diagnosis of CLL. These findings were indicative of leukemic involvement in the CNS, and the patient was eventually transferred to hematology. In this case, Richter's syndrome was initially suspected from the symptoms and course, but CNS-infiltrating cells were small lymphoid cells similar to those of PB and BM, and transformation to a diffuse large-cell type was ruled out by brain biopsy. Therefore, we diagnosed CNS involvement in CLL.

Laboratory data (Table 1) were significant for a white blood–cell count of 464,200/μL (98.5% lymphocytes and 1.5% neutrophils). Hemoglobin level and platelet count were 11.7 g/dL and 305,000/μL, respectively. Lactate dehydrogenase was 262 IU/L (normal range 112–230 IU/L) and soluble IL2R 11,000 IU/L (normal range 124–466 IU/dL).

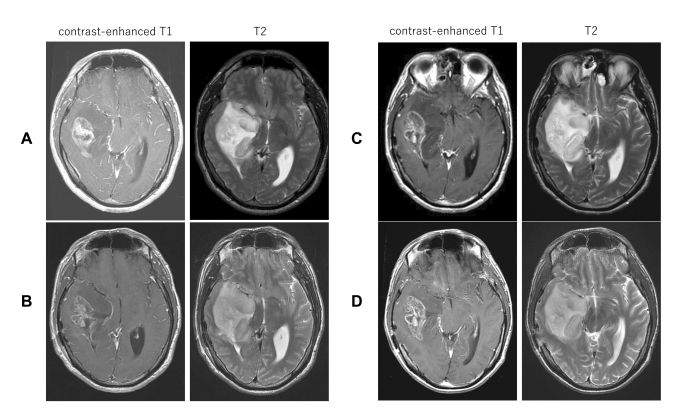


Figure 1 Magnetic resonance imaging (MRI) showing 5×3.5 cm abnormal nonuniformly contrasted mass with hypointensity on T_1 -weighted image (left) and with hyperintensity on T_2 -weighted image (right) in the right temporal lobe. (**A**) MRI at first consultation; (**B**) MRI after MPV administration (at day 17 after admission); (**C**) MRI after FCR administration (at day 34 after admission); (**D**) MRI after Ibr administration (at day 54 after admission).

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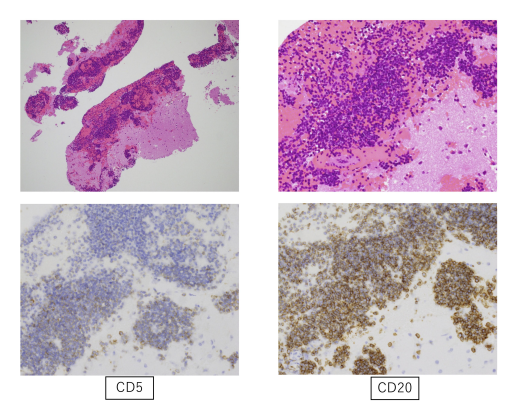


Figure 2 Brain specimens (cranioscopic biopsy) showing infiltration of small monoclonal lymphocytes with expression of CD5 and CD20 (upper left, H&E ×40; upper right, H&E ×100; lowerleft CD5 ×100; lower right, CD20 ×100).

 β_2 -microglobulin was 2.1 mg/L. Evaluation with thoracoabdominal computed tomography (CT) revealed splenomegaly and mild systemic lymphadenopathy.

Treatment with 2 mg betamethasone for 7 days transiently improved the diplopia and headaches, but tumor size evaluated by CT/MRI remained unchanged. No standard protocol exists for CLL with CNS involvement, because it is an extremely rare disease condition. Therefore, according to the treatment strategy of primary CNS lymphoma, MPV chemotherapy (methotrexate 3.5 mg/m² on day 1, vincristine 1.4 mg/m² [max 2.8 mg on day 1], and procarbazine 100 mg/m² per day on days 1-7) was started. Ten days after treatment, intracranial hypertension-related symptoms, such as diplopia and headaches, recurred and performance status was decreased. MRI showed that the tumor size remained unchanged (Figure 1B) and PB-lymphocyte reduction was poor (Figure 4), indicating resistance to the MPV treatment. Therefore, rituximab (Rtx) 375 mg/m² and subsequent whole-brain radiotherapy (30 Gy/15 fr) plus simultaneous in-field boost (10 Gy/5 fr) were administered.

After Rtx administration, the diplopia and headaches improved and lymphocyte reduction was observed.

Therefore, treatment with one cycle of FCR chemotherapy (fludarabine 25 mg/m² per day and cyclophosphamide 250 mg/m² per day for the first 3 days, with addition of Rtx 375 mg/m²) was started. Although the PB lymphocytes decreased steadily (Figure 4) without recurrence of intracranial hypertension-related symptoms, no reductive effect on the intracranial tumor was observed on contrast-enhanced MRI (Figure 1C). Because the effects of ibrutinib (Ibr) on the CNS have been reported in CLL and mantle-cell lymphoma,6 we next selected Ibr 420 mg/day for treatment. However, 2 weeks later, contrast-enhanced MRI revealed no reductive effect, and diplopia and headaches had recurred (Figure 1D). Finally, the patient refused subsequent treatment and was self-discharged from the hospital. He died at home 9 weeks after the onset of initial symptoms (48 weeks after the diagnosis of CLL).

Discussion

Diagnostic cranioscopic biopsy was performed in our case, but many cases were diagnosed by cerebrospinal fluid (CSF) analysis in a retrospective cohort of 30 CLL patients with CNS involvement.⁷ In that cohort, biopsies

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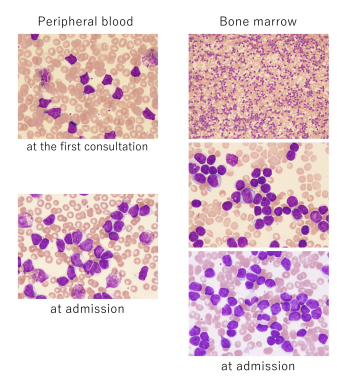


Figure 3 CLL cells from peripheral blood at first consultation and admission and BM at admission showed mature small monoclonal lymphocytes with narrow cytoplasm, concentrated nuclei, and partially aggregated chromatin without transformation to a large cell type (H&E ×40 and ×200).

were performed in only five cases, of which only one was diagnosed by brain biopsy. Our review of the literature revealed diagnostic biopsies had been performed in 12 of 50 cases (not including surgical resection). Ten of the 50 cases were diagnosed as Richter's syndrome, and 11.3% of Richter's transformation with intracranial involvement was found in an old literature review of CLL (before 2011).8 By contrast, there were no cases of Richter's syndrome in the 30 cases of the retrospective cohort. It has been reported that Richter's transformation occurs in approximately 5%–10% of the CLL population; therefore, it is still difficult to conclude whether there is an intimate correlation between CNS involvement and Richter's transformation.

The 50 reported cases of CNS involvement in CLL had diverse and uncharacteristic symptoms, such as headaches, convulsions, diplopia, ataxia, facial paralysis, and cognitive dysfunction (Table 2). It is difficult to identify the risk factors for CNS involvement in CLL. 10,11 Our literature review confirmed this, because we could not find a common feature in cases of CNS involvement. There are cases in which CNS involvement develops when the stage is not necessarily progressive (on Rai or

Table I Hematologic Assessment of Patient

White blood Cells/μL	464,200/μL	Na	I4I mEq/L
Neutrophils	1.5%	K	4.2 mEq/L
Basophils	0	Cl	103 mEq/L
Eosinophils	0	BUN	12 mg/dL
Lymphocytes	98.5%	Cr	0.83 mg/dL
Monocytes	0	TP	6.5 g/dL
Others	0	Alb	4.1 g/dL
Plt	30.5×10 ⁴ /μL	AST	26 U/L
RBC	449×10 ⁴ /μL	ALT	37 U/L
Hb	II.7 g/dL	T-Bil	0.4 mg/dL
Ht	42.1%	D-Bil	0 mg/dL
MCV	93.8 fL	ALP	544 U/L
MCH	26.1 pg	γGTP	87 U/L
MCHC	27.8 g/dL	LDH	262 U/L
APTT	25.7 seconds	CRP	0.072 mg/dL
PT	107.4%		
FBG	241 mg/dL	IgG	955 mg/dL
AT-III	109%	lgA	83 mg/dL
HBs-Ag	_	IgM	29 mg/dL
HCV-Ab	_	sIL2R	11,000 U/mL
HTLV-I	_	BMG	2.1 mg/L
HIV	_	ANA	_
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Binet staging) or without high-risk chromosomal abnormality, such as del17p or del11q. This suggests clinical and pathophysiological heterogeneity of CNS involvement in CLL.⁷

A report summarizing the literature published before 2011 of CNS involvement in CLL⁸ showed average age 63.4 years, average latency between CLL diagnosis and first signs of CNS involvement 2.6 years, average overall survival (OS) from CLL diagnosis 3.8 years, and average OS from time of CNS onset 12 months. Our review of the 50 case reports revealed average age 62.2 years (in 49 cases) and average latency 4.9 years (in 32 cases. OS data could not be extracted. Our case showed a younger and more aggressive disease course of age 33 years, latency 9 months, OS 48 weeks, and OS from time of CNS onset 9 weeks. Our case was resistant to high-dose Mtx and whole-brain radiotherapy as standard treatments for primary CNS lymphoma. As the standard treatment for non-high risk CLL, FCR was effective in reducing the number of PB lymphocytes and improved intracranial hypertension-related symptoms; however, it had less effect on tumor shrinkage, indicating it was ineffective for the CNS lesion. Although the number of reports of CNS involvement in CLL is low, there were reports of successful treatment with FCR in some cases in our

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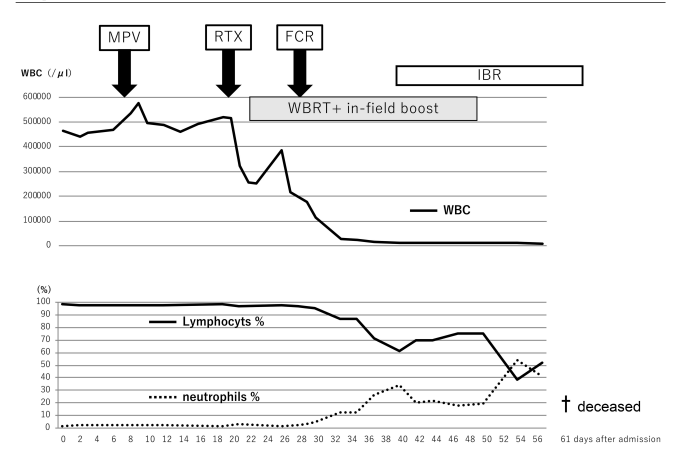


Figure 4 Clinical course of WBC and lymphocyte counts after treatment.

Abbreviations: MPV, methotrexate 3.5 mg/m² (day 1), vincristine 1.4 mg/m² (max 2.8 mg) (day 1), and procarbazine 100 mg/m² per day (days 1–7); Rtx, rituximab (375 mg/m²; FCR, fludarabine (25 mg/m² per day), cyclophosphamide (250 mg/m² per day) for the first 3 days, with addition of Rtx 375 mg/m²; WBRT, whole-brain radiotherapy (30 Gy/15 fr) plus simultaneous in-field boost (10 Gy/5 fr).

literature review (Table 2). However, in general, prognosis was poor, owing to resistance to various treatments, such as high-dose Mtx, intrathecal injection, whole-brain radiotherapy, and FCR. A similar result was obtained in our case

Ibr has been reported to be effective in CNS lesions of mantle-cell lymphoma⁶ and Waldenström macroglobulinemia. ¹² Effects of Ibr appear 1–2 weeks after administration. ¹³ Nine successful cases of Ibr treatment for CNS involvement in CLL were found in 50 cases (complete response in eight cases, partial response in one) (Table 2); therefore, Ibr may be a promising drug for CNS involvement. However, this was not found in our case. It is possible that the effective concentration of Ibr in the CNS lesion had not reached sufficient levels in our case. Concentration in CSF was reported to be 2log lower than in the plasma of 18 patients with primary CNS lymphoma treated with Ibr. ¹⁴ It has been reported that an increased dose of Ibr escalates CSF concentration

without adverse events,¹⁵ and that increasing the dose of Ibr is effective for CNS lesions in CLL.¹⁶ It will be necessary in the future to verify the optimal dose of Ibr for CNS lesions. Bulky disease in CNS lesion might also cause treatment failure. Although ofatumumab and alemtuzumab are alternative treatment options, we did not select them, because they did not show superiority to Ibr in the data or in drug penetration of the CNS. In addition, there have been two reports showing the effectiveness of venetoclax against CNS lesions (Table 2).^{17,18} One of those was a case where venetoclax was effective after Ibr resistance, and thus it may be beneficial to test venetoclax against CNS involvement in CLL.

Conclusion

Patients with CNS lesions in lymphoid tumors have a poor prognosis, but the possibility of concomitant use of Mtx and Ibr or venetoclax can be envisioned. Accumulation of

 Table 2 Literature on CNS Involvement in CLL

Reference	Age (years), sex	Symptoms	Interval from diagnosis of CLL to first CNS	Rai/ Binet	Lymphocyte	(G-band)	Method of diagnosis	Transformation to Richter's syndrome	Treatment (treatment prior to diagnosis of CNS involvement)	Response
Clin Case Rep. 2020.8.269. 17	7.1 M	Epileptic seizures	12 years	ĄV	∀ Z	l lq deletion	CSF- Clinical diagnosis		WWV/FCR/ Rtx-bendamustine— Ibr HD Mtx Rtx Venetoclax	Venetoclax PR
Mult Scler Relat Disord. 2020.37.101455	50 M	Fecal incontinence, tetraparesis	∀ Z	A N	131,000 (WBC)	∀ Z	CSF+		Rtx and cyclophosphamide IVIg	CR
Can J Neurol Sci. 2019,46.640. ²⁰	53 M	Neck pain, adenopathy, urinary retention, monocular vision loss in right eye	5 years	∀ Z	∀ Z	⋖ Z	CSF⁺		wwv Dex pulse→ Rtx and cyclophosphamide	R S
Haematologica.2019.104. e222. 18	28	٧	₹ Z	∢ Z	∢ Z	Trisomy 12	CSF⁺		Six FCR courses, six Rtx-bendamustine cycles, and four Rtx- DHAP courses lbr Venetoclax with IT chemotherapy (cytarabine plus methotrexate)	lbr PD Venetoclax CR

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ĸ.	HD Mtx with IT Mtx and AraC	Rtx, HD Mtx, Dex, and vincristine with IT Mtx	CHOP/FC HD Mtx	WW/FCR Ibr 420 mg→560 mg	Chlorambucil monoctherapy IT Mtx + liposomal AraC →lbr 420 mg →HD Mtx →WBRT	CVP, CHOP, FC, F, ofatumumab + idelalisib vs no treatment
- AZ	H e	Rtx, and v	ㅎ 보	> ₫		Of ide
						Yes
						7
Craniotomy with resection CSF	Śś t.	SS t.	, t.	+.	+.	CT Autopsy
Crani with CSF	Biopsy CSF ⁺	Biopsy CSF ⁺	Biopsy CSF ⁺	CSF ⁺	CSF ⁺	CT
(BM)	₹ Z	₹ Z	Trisomy 12 (lymph node)	∀ Z	l 3q deletion	∀ Z
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5,300 (WBC)	₹ Z	₹ Z	3300 (WBC)	¥ Z	¥Z	∢ Z
₹ Z	∢ Z	₹ Z	₹ Z	Rai I	Rai 0 Binet A	Rai II
			ars	ars	د	71 months
₹ Z	₹ Z	₹ Z	14 years	14 years	9 years	7 I m
esis,	ory of	l oast 5	oance		vical o the	of
Dysmetria, left upper— extremity paresis, apraxia, mild amnesia, and prosopagnosi	6-month history of headache	Headache and dizziness for past 5 years	Mental disturbance	iic ches	Headache complaints, photophobia, vertigo, and extensive pain (from the cervical spine down to the inferior limbs)	Disturbances of consciousness
Dysmet upper- extremi apraxia, amnesia	6-month l	Heada dizzine years	Menta	Chronic headaches	Headache complaints, photophobii vertigo, and extensive pa (from the ce spine down inferior limb	Distur
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Case Rep Hematol. 2019. 1,825,491;	BMC Neurol. 2019 0.19. 200.	BMC Neurol. 2019 0.19. 200.	Neuropathology. 2019.39.54. ²³	Cureus. 2018. 10. e2176.	Case Rep Hematol. 2018. 7,817,918. 24	Adv Clin Exp Med. 2018.27. 1683. 25
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Table 2 (Continued).

Reference	Age (years), sex	Symptoms	Interval from diagnosis of CLL to first CNS symptoms	Rai/ Binet	Lymphocyte	(G-band)	Method of diagnosis	Transformation to Richter's syndrome	Treatment (treatment prior to diagnosis of CNS involvement)	Response
Turk J Hematol 2018.35.147.	7.1 F	Expressive aphasia, memory problems, confusion, and headache	12 years	Binet A	14,652	Normal	Biopsy reject CSF		ww Rtx Ibr 420 mg	Æ
Annals of Hematology. 2018.97. 1627.		Paralysis of the left oculomotor nerve and left hemianopsia	20 months	Rai 0 Binet A	35,600	Y Z	CSF*		WWV Rex and chlorambucil with IT Mtx	æ
Annals of Hematology. 2018.97. 1627.	77 M	Apathy, urinary incontinence	9 years	Rai II Binet B	44,500	Y Z	CSF*		Fludarabine and cyclophosphamide No	Q
Medicine. 2018. 97. e12701 ²⁸	67 F	Slurred speech, headache, and left- sided hemiparesis	NA	₹ Z	14,500	p53+	Surgical resection CSF (DLBCL)	Yes	HD Mtx IT Mtxand AraC	Q
Cureus.2018.10. e3660	84 F	Mild dysmetria in the upper-left extremity	V V	Rai 0	15,311,000	∀ Z	Tumor resection	Yes	Temozolomide and WBRT	PD
Ann Indian Acad Neurol. 2018. 21. 85.	57 M	Bradypsychia, headaches, nausea, vomiting	6 months	Rai III	85,500 (WBC)	del17p	CSF*		WW HD Mtx	Q

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SD,	D	Q	წ	PR→PD	క	క	CR.
균 줬	ww DHAP	СНОР DHAP	Steroids Ibr 420 mg	FCR	lbr with IT Mtx	Eight prior lines of therapy for CLL lbr 420 mg	Four prior lines of therapy for CLL lbr 420 mg
CSF⁺	CSF-	CSF⁺	CSF ⁺	CSF⁺	Optic nerve- sheath biopsy CSF ⁺	CSF⁺	CSF⁺
Normal	13q14 nullisomy	∢ Z	₹ Z	Normal	∢ Z	del I 7 p	del I 7 p
23,000 (WBC)	103,900 (WBC)	86,500 (WBC)	94,000	13,400	2304	¥ Z	A N
Rai IV	Rai IV	Rai II	Rai III	Rai 0	₹ Z	Binet C	Binet B
62 months	9 months	63 months	₹ Z	∢ Z	2 years	∀ Z	∢ Z
Dysphasia, repeated unconsciousness, urinary incontinence	Dyslexia, lack of fine motor control, diplopia	Diplopia, bilateral eyelid swelling, and tumors	Tightness, paresthesia, and neuropathic pain in the left hand and left arm	Progressive lower- extremity weakness and urinary incontinence	Visual loss in the right eye	Dysautonomy	Headaches and cognitive disturbance
ξ Σ	72 M	4 Α	9 Σ	60 F	45 Μ	Σ8	75 M
Ann Indian Acad Neurol. 2018. 21. 85.	Ann Indian Acad Neurol. 2018. 21. 85.	Ann Indian Acad Neurol. 2018. 21. 85.	Br J Haematol. 2017.176. 829.	BMC Hematol. 2017.17.3	J Neuroophthalmol. 2016. 36.61.	Blood. 2016.127. 2356	Blood. 2016.127. 2356 34

Table 2 (Continued).

Reference	Age (years), sex	Symptoms	from diagnosis of CLL to first CNS	Rai/ Binet	Lymphocyte	(G-band)	Method of diagnosis	Transformation to Richter's syndrome	Treatment (treatment prior to diagnosis of CNS involvement)	Response
Blood. 2016.127. 2356–2358	Σ 63	Cerebellar syndrome and aphasia, confusion	₹ Z	Binet C	₹Z	∀ Z	CSF ⁺		Two prior lines of therapy for CLL lbr 420 mg	ర
Blood. 2016.127. 2356	89 F	Visual loss	₹ Z	Binet A	₹Z	dell7p	CSF ⁺		No prior lines of therapy for CLL lbr 420 mg	Z.
BMC Res Notes. 2014.7.645.	75 F	Headache, otalgia in the right ear, fever, dizziness, and dysphagia	5 years	Rai I	24,300 (WBC)	₹ Z	CSF ⁺		Chlorambucil and prednisone IT Mtx FC	ర
Leukemia Lymphoma. 2014. 55.1939 36	2 Σ	Hypoesthesia	2 months	Binet B	251,000	Normal	CSF ⁺		WW IT Mtx + AraC Rtx—bendamustine	S.
BMJ Case Rep. 2014. Bcr-2013-202,051.	45 F	Seizures, headaches, and vomiting	NA	A N	Y Z	∀ Z	Biopsy		Surgical excision RT	Relapse
Clin Lymphoma Myeloma Leuk. 2013.13. 338. ³⁸	44 F	Double vision	3 years	Rai I	98,280	Trisomy 12 and 13q-	Biopsy CSF ⁺		Rtx mPSL IT AraC FCR	೪

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Leuk Lymphoma. 2013.54. 2070. 39	Σ Σ	Gait disturbance, tremors, slurred speech, marked fatigue, intermittent confusion, and visual impairment	10 years	Rai –	2200 (WBC)	Diploid	CSF*	Ş.	Chlorambucil— fludarabine— pentostatin, cyclophosphamide, and Rtx— ofatumumab— lenalidomide IT liposomal AraC IT Rtx HD AraC WRRT	Q
J Clin Exp Hematop. 2013.53. 157	99 4	Fatigue and difficulty walking	2 years	Rai 0	27,000	₹ Z	Biopsy (CT. guided), non-GCB DLBCL	Yes	ww// cyclophosphamide + PSL IT Dex RT Rtx	Transient PR⊸relapse
J Clin Oncol. 2013.31 e280 41	Σ Σ Σ	Right-eye pain associated with blurry vision, floaters, and bright halos	l years	Rai 0	34,900	Trisomy 12	CSF ⁺		PSL Rtx—fludarabine	ర
Acta Haematol. 2012;127. 93	Σ Σ	Seizures, psychomotoric deficits, and left- sided hemiparesis	l month	Binet A	₹ Z	₹ Z	Stereotactic biopsy CSF	Yes	Systemic and intraventricular polychemotherapy regimen WBRT	Transient CR→relapse

Table 2 (Continued).

Reference	Age (years), sex	Symptoms	Interval from diagnosis of CLL to first CNS symptoms	Rai/ Binet	Lymphocyte	(G-band)	Method of diagnosis	Transformation to Richter's syndrome	Treatment (treatment prior to diagnosis of CNS involvement)	Response
J Neurooncol 2012.106. 185 8	F 33	Vision changes	4 years	Rai I	16,000,000 (WBC)	∢ Z	CSF⁺		CVP Rtx-fludarabine Alemtuzumab Rtx IT Mtx Temozolomide RT	ж
J Neurooncol 2012.106. 185	52 M	Encephalopathy, dementia, seizures	3 years	Rai IV	14,000,000 (WBC)	∢ Z	CSF⁺		Fludarabine None	РО
J Neurooncol. 2012.109. 213 43	65 F	Difficulty speaking, weakness in the right arm and leg	I month	Rai IV	600,000 (WBC)	13q14 deletion (biopsy)	Biopsy		Cyclophosphamide + steroids FCR WBRT Rtx—bendamustine	చ
Case Rep Hematol. 2012 589,718 44	99 H	Bilateral hearing loss	∀ Z	₹ Z	104,000 (WBC)	del (17p13.1) and del (13q34)	Tympanic membrane biopsy CSF*		Rtx-CVP Rtx IT liposomal cytarabine Cyclophosphamide, cladribine, and Rtx HyperCVAD	ж
Am J Hematol. 2011.86.783 45	73	Bilateral visual loss	∀ Z	Rai II Binet B	ĄV	Y N	Ethmoidectomy		Fludarabine/FCR Steroids RT	PD

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	PD 6 months			Transient CR →relapse	Transient CR → relapse		
G	PD m 9	R	r.	Tran CR	Tran CR ↓re	ర	٦.
WWV/chlorambucil/ FCR/CHOP HD Mtx	Mtx Flu RT	HD Mtx + ifosfamide HD AraC + Mit + IT Mtx Dasatinib	IT liposomal AraC RT Rtx + VCR + HD Mtx + PCBZ + HD AraC FCR	IT liposomal AraC HD Mtx + HD AraC	IT liposomal AraC Chlorambucil	IT liposomal AraC FCR	IT liposomal AraC R-CHOP
Yes				Yes			
Brain biopsy	Open-brain biopsy CSF	CSF⁺	CSF⁺	CSF⁺	CSF ⁺	CSF⁺	CSF⁺
Normal	∀ Z	Ą	Y Y	V	∢ Z	Y Z	₹ Z
A N	36,000 (WBC)	₹ Z	∀ Z	∢ Z	¥ Z	∀ Z	∀ Z
Rai III	Rai III	∀ Z	Rai Binet A	Rai IV Binet B	Rai IV Binet C	Rai 0 Binet A	Rai IV Binet C
A Z	10 years	∀ Z	61 months	34 months	25 months	25 months	13 months
Bilateral leg weakness, pain, and urinary retention	Temporary seizures, poor memory, and progressive blindness	Paraparesis of both legs, urinary and stool incontinence, and central rightsided facial nerve palsy	Headache, cognitive complaints: slow response and inattentiveness	V cranial pair palsy	Headache, chin and face dysesthesia, optic neuritis, blurred vision	Headache, nausea, weakness, somnolence, lethargy, and confusioned state	Headache and diplopia
late60sM	8ς Σ	89 Σ	Σ2 Μ	68 F	4 Σ	<u>ω</u> Σ	64 F
J Neurol Neurosurg Psychiatry. 2011.82.943	J Clin Oncol. 2010.28.e30	Blood.2010. 116. 2617 48	Br J Haematol. 2010.150.618 49	Br J Haematol. 2010.150.618 49	Br J Haematol. 2010.150.618 49	Br J Haematol. 2010.150.618 49	Br J Haematol. 2010.150.618

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Table 2 (Continued).

Reference	Age (years), sex	Symptoms	from diagnosis of CLL to first CNS	Rai/ Binet	Lymphocyte	FISH Method o (G-band) diagnosis	Method of diagnosis	Transformation to Richter's syndrome	Treatment (treatment prior to diagnosis of CNS involvement)	Response
Br J Haematol. 2010.150.618 ⁴⁹	79 M	Leg weakness, difficulty walking, upper-back pain, and VII cranial pair palsy	66 months	Rai IV Binet B	NA	NA	CSF⁺	Yes	IT Mtx and liposomal AraC HyperCVAD/MA	Transient CR →relapse
Br J Haematol. 2010.150.618 49	89 F	Headache	24 months	Rai IV Binet B	∢ Z	∢ Z	CSF⁺	Yes	IT Mtx and liposomal AraC AraC	ర

Abbreviations: NA, not available; FISH, fluorescence in situ hybridization; CSF, cerebrospinal fluid; Rxx, rituximab; FCR, fludarabine–cyclophosphamide–Rxx; MPV, methotrexate–procarbazine–vincristine; HD, high dose; Mtx, methotrexate, Mit, mitoxantrone; AraC, cytosine arabinoside (cytarabine); CVP, cyclophosphamide–vincristine–pradnisone; IT, intrathecal; RT, radiotherapy; Ibr, ibrutinib; WBRT, whole-brain radiotherapy; hyperCVAD, cyclophosphamide–hydroxydaunorubicin (doxorubicin)–oncovin (vincristine)–prednisone; DHAP, dexamethasone–HD AraC (cytarabine)–platinol (cisplatin); WW, watch and wait.

Dovepress Nakanishi et al

data from cases is important to verify the choice of new or combination drugs for administration from an early stage.

Ethics

Informed consent was provided by the patient on admission to have the case details published. The patient passed away before publication of causes not included in the case report. Institutional approval was not required for publication.

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Author Contributions

All authors contributed to data analysis, drafting or revising the article, have agreed on the journal to which the article will be submitted, gave final approval to 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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Nakanishi et al **Dove**press

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