Primary CNS Lymphoma: Analysis of Treatment by Gamma-Knife Radiosurgery and Chemotherapy in a Prospective, Observational Study

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Abstract

Background: Primary central nervous system lymphomas (PCNSL) include 4% of all primary brain-tumors. It can involve the brain parenchyma, leptomeninges, eyes and the spinal cord. Unlike systemic lymphoma, durable remissions remain uncommon. Different combinations of chemotherapy and Methotrexate are the gold standard. This is a clinical study comparing the outcomes using Methotrexate and Methotrexate with Gamma Knife Radiosurgery (GKRS).

Methods: Retrospective, comparative study. Study arms, control group performed by Methotrexate as a single agent, in dose of 8 g/m² and treatment group Methrotexate and GKRS. Strict inclusion and exclusion criteria were employed and review of histologically confirmed PCNSL. Primary objective were survival rate. Secondary outcomes were the remission or re-incurrence of a lesion.

Results: Between June 1995 and May 2015, 128 charts were evaluated, 73 subjects in Control group and 55 in GKRS group. Follow-up period 24 to 49 months (mean 36.9 months). There were no statistically significant differences in patient demographics or histology diagnosis. GKRS doses ranging from 12 Gy to 26 GY (median: 13 Gy). Median survival rate 26, 8 months control group and 47.6 in GKRS group from the 1st pharmacological treatment. All lesions had complete response after GKRS at 3-8 weeks (mean range 5.8 weeks). Control group (8-20 weeks).

Conclusion: GKRS is non-invasive, safe, and shows rapid success, improving the prognosis of the patients. This noninvasive treatment modality should be considered as an alternative or co-adjuvant option in PCNSL. Level of Evidence: Retrospective, Comparative study (Level III).

Keywords: Primary CNS lymphoma; Radiosurgery; Gamma knife; Chemotherapy; Brain tumor; Methotrexate; Lymphoma

Introduction

Primary central nervous system lymphoma (PCNSL) is a rare cancer accounting for less than 4% of brain tumors [1]. These lesions usually present as intracranial nodules with diffuse meningeval, periventricular, and perivascular spread being very common. The primary central nervous system extranodal high-grade malignant cells are usually large or immunoblastic cells of B-cell non-Hodgkin’s, tracing their origin back to the brain parenchyma, spinal cord, leptomeninges or eyes and are typically limited to the central nervous system (CNS) with infrequent extension outside the nervous system. Treatment modalities for patients with newly diagnosed PCNSL include radiation and/or chemotherapy [2]. While the role of radiation therapy for initial management of PCNSL is controversial, clinical trials will attempt to improve the therapeutic index of this modality. Routes of chemotherapy administration include intravenous, intraocular, intraventricular or intra-arterial. Multiple trials have outlined different methotrexate-based chemotherapy regimens and have used local techniques to improve drug delivery. A major challenge in the management of patients with PCNSL remains the delivery of aggressive treatment with preservation of neurocognitive function [3,4]. Routinely PCNSL present as single or multifocal, massive infiltrative lesions that may occur in the cortex, and penetrating deep into the white and/or gray matter. The lesions may show areas of necrosis especially in patients with immunodeficiencies. GKRS has not yet been used universally in the treatment of the PCNSL and is non-invasive, safe and shows prompt and successful results, improving the prognosis and quality of life of the patient. Like other brain tumors, one of the major contests in treatment with chemotherapeutic agents remains the delivery of therapeutic concentrations of drugs to the CNS. A study of blood–brain barrier (BBB) permeability in a rat brain tumor model established a large heterogeneity of microvascular leakage; the vasculature within and around the brain tumors has a wide range of permeability, from typical capillaries with no (BBB) leakage to a tumor vasculature that allows free entry of large molecules. Although MTX crosses the BBB, remote minus is quantifiable in the brain tissue than in the serum. High-dose MTX (>1 g/m²) has been shown to be an independent factor correlating with survival [5-8]. Thus, MTX is administered in high doses, up to 8g/m², in order to achieve therapeutic drug concentrations in the tumor and surrounding brain. Intravenous (IV) doses less than 1 g/m², similar to what has been used in the treatment of other malignancies outside the brain, reach CNS concentrations generally felt not to be cytotoxic. The purpose of this retrospective comparative study is to determine the effectiveness of chemotherapy using methotrexate in dose of 8g/m² + radiosurgery in the treatment of PCNSL.

Materials and Methods

This is a retrospective, comparative study evaluating the effect and

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survival rate of patient with a histological diagnosis of PCNSL treated by Methotrexate and Methotrexate + GKRS. The study received approval through the Institutional Review Board. Strict inclusion and exclusion criteria were employed (Table 1). All patient was included if they have a pathology report of histologically confirmed PCNSL. Primary inclusion criteria was subjects received Methotrexate therapy as single agent in a standard protocol dosage of 8 g/m². This chart review evaluate clinical data and MRI images from June 1995 - May 2015.

Patients with diagnosis of PCNSL by patho-histology report of the brain lesion observed on MRI were added. The presence and size of the tumor, as well as its localization were confirmed using Brain MRI Neuro- Oncologist and Radio-Oncologist Protocol. The maximal brain tumor was measured at 1 cm and 2 cm proximal to the most distal aspect of its midline. Primary outcomes was survival rate and secondary assessed by the sensibility and decrease of the brain tumor lesion as noted on MRI.

Chart review

One experienced blinded post-doctoral fellow collected the data using red cap system from eligible patients. Those patients underwent Gamma Knife Radiosurgery and/or Methotrexate dose from institutional protocol. Patients' inclusion criteria are outlined in Table 1.

Subjects was excluded on various grounds including whether or not they previously failed a surgical brain tumor removal, radiation therapy or any other chemotherapy protocol.

Statistical analysis

A power analysis by mortality rate had been evaluate by y a priori sample size calculation performed on the basis of a prior randomized study [9], which evaluated using a survival rate with a sample size of 40 patients. This study suggested a sample size of at least 31 subjects per group to detect the minimal clinically important difference in mortality rate of 1.5 standard deviations (σ = 1.5, α = 0.05, β = 0.20). Additionally, a general estimate for detecting a one-unit change on an ordinal scale of 136 (σ = 1.0, α = 0.05, β = 0.20) resulted in the same number. The Wilcoxon rank-sum test was used for comparison of continuous data between groups. Differences between means were analyzed using two-sided t-tests. Comparison of categordical data was performed using Pearson chi-square or Fisher’s exact probability test as indicated. Ordinal ranking scores were compared using the Mantel-Haenszel test. Differences less of p=0.05 or less was reported statistical significant [10].

Results

Between June 1995 and May 2015, 128 charts was evaluated. 73 subjects in control group (Methotrexate) and 55 (Methotrexate + GKRS). Follow-up period 24 to 49 months (mean 36.9 months). There were no statistically significant differences in patient demographics or histology diagnosis (Table 2). Patients was treated using GKRS doses ranging from 12 Gy to 26 Gy (median:13.6 Gy) (Figure 1). Median survival was rate 26.8 months in Control group and 47.6 months in GKRS group from the initial diagnosis. All lesions showed a complete response after Radiosurgery when evaluated using magnetic resonance imaging 3-8 weeks (mean range 5.8 weeks).

The histologic finding in all patients was the presence of large, diffuse B-cell lymphoma. The symptoms and signs of the patients remained evidently improved within 2 - 6 weeks after GKRS and 4-10 weeks after chemotherapy alone. All treated lesions by GKRS showed complete response in magnetic resonance imaging 3 weeks to 8 weeks (mean range 6.3 weeks). There was a statistical significant difference of median survival rate chemo group alone 36.3 months from the initial diagnosis and 49.8 months in GKRS group (P=0.0034) [11,12]. The doses ranged from 12 Gy to 26 Gy (median: 13.8) 50% isodose line (range: 45-85; median 50) (Table 3). The clinical data was gathered for brain tumor lesion control, disease control, toxicity, recurrence and survival. The Kanofsky performance grade was enhanced from a preoperative average of 55% to a postoperative one of 88% in GK group. Significant factors contributing to survival greater 24 months following GKRS, were increased marginal dose (Odds ratio = 6.5,

### Table 1: Inclusion and exclusion criteria.

<table>
<thead>
<tr>
<th>Patient Parameters</th>
<th>Control</th>
<th>GKRS Group</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of Patients</td>
<td>73</td>
<td>55</td>
<td>NA</td>
</tr>
<tr>
<td>Age (years)</td>
<td>58.1 ± 5.3</td>
<td>56.9 ± 3.3</td>
<td>0.685</td>
</tr>
<tr>
<td>Male</td>
<td>33 (70%)</td>
<td>26 (56%)</td>
<td>0.51</td>
</tr>
<tr>
<td>Female</td>
<td>40 (30%)</td>
<td>19 (44%)</td>
<td></td>
</tr>
<tr>
<td>Body-mass index (kg/m²)</td>
<td>24.1 ± 2.6</td>
<td>22.2 ± 3.1</td>
<td>0.39</td>
</tr>
<tr>
<td>Pre-op Diagnosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tobacco use</td>
<td>16</td>
<td>22</td>
<td>0.25</td>
</tr>
<tr>
<td>Pre-albumin (mg/dL)</td>
<td>18.1 ± 1.3</td>
<td>19.1 ± 0.9</td>
<td>0.347</td>
</tr>
<tr>
<td>Albumin (g/dL)</td>
<td>3.4 ± 0.4</td>
<td>3.2 ± 0.4</td>
<td>0.89</td>
</tr>
<tr>
<td>Absolute Lymphocyte Count (cells/μL)</td>
<td>1745 ± 468</td>
<td>1798 ± 401</td>
<td>0.20</td>
</tr>
<tr>
<td>Median Survival</td>
<td>26.8 months (18.3 – 29.1)</td>
<td>47.6 months (28.3 – 49.1)</td>
<td>0.0034</td>
</tr>
<tr>
<td>Pre-op Diagnosis</td>
<td></td>
<td></td>
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</tbody>
</table>

### Table 2: Demographic and perioperative data.

### Figure 1: Brain MRI Neuro- Oncologist and Radio-Oncologist Protocol. The maximal brain tumor was measured at 1 cm and 2 cm proximal to the most distal aspect of its midline. Primary outcomes was survival rate and secondary assessed by the sensibility and decrease of the brain tumor lesion as noted on MRI.
developed new lesions in other areas of the brain and repeated GKS was performed. No local recurrence was observed in any patient at a mean of 30.6 months post-GKS treatment. Our findings are in accordance with those of Kenia, and seems to be at a higher success rate that those described by others in treating patients with chemotherapy and/or radiotherapy alone. There appeared to be a predisposition in the direction of new tumor growth in individuals who did not undergo GK treatment. We establish no suggestion of a connection between recurrence and original appearance of multiple or solitary lesions but current systemic chemotherapy would be essential. The prognosis of untreated PCNSL remains grim with median survival time being low. GKR appears to be a valuable method for the management of primary and relapsed CNSL, since it enables exceptional local control in a short-term management period without severe complications.

Limitation
This study has a few limitations including: (i) it is a retrospective, comparative study and (ii) lack of prospective, randomized data collection confines us in assessing the full benefits and complications of GKRS.

Conclusion
Given the well tumor control, enhanced overall survival period and lesser number of complications, GKRS can be an ideal co-adjuvant treatment option for the patients with PCNS lymphoma adding of standard Methotrexate treatment. In addition, well and prompt tumor was reported using Gamma Knife Radiosurgery allowing the immediate improvement of neurological symptoms.

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References


