INTRODUCTION

Neoplastic meningitis (NM) is the leptomeningeal dissemination of metastatic tumors; a devastating complication from solid tumors. The incidence of NM has increased as patients are living longer due to significant improvements in treatment options in the form of large molecule target agents. There are case reports about cancers that don’t yet progress into NM, such as ovarian cancer,[1] prostate cancer,[2] and renal cancer.[3]

NM is clinically detected in 5–8% of the patients with cancers, while through autopsies NM detected in 19% of the cancerous patients.

Cerebrospinal fluid cytology (CSFC) is the gold standard for determining NM, with the reported sensitivity of CSFC being 71–94%. The survival of NM ranges from 8 to 16 weeks despite treatment.[10,11] Patients have a poor Karnofsky Performance Score (KPS), when diagnosed with either bulky central nervous system (CNS) disease, abnormal CSF-flow study, multiple serious neurological deficits, encephalopathy, and extensive systemic cancer without good treatment options have poorer prognosis and need palliative care instead of positive therapy.[9] For an improved outcome, most patients of NM need a combination of radiation therapy, systemic chemotherapy, and intrathecal chemotherapy.

Intrathecal chemotherapy is the main treatment of NM. Methotrexate, cytarabine, thiotepa, liposomal cytarabine are the traditional intrathecal chemotherapy regimens. Intrathecal methotrexate has a long history of treating NM.[12] Intrathecal methotrexate is now widely used to treat NM in the patients with those cancers with possible metastasis to the CNS, such as gastric cancer, breast cancer, lymphoma, non-small-cell lung cancer, multiple myeloma, as well as in the patients with cancers rarely spreading to CNS-atypical neurofibroma and with pancreatic cancer.[13–19] Though many physicians use intrathecal cytotoxic drugs in combination with system
chemotherapy or target agents,[15,16,20] it is irreplaceable in the treatment of NM, despite of some reported adverse reactions.[21,22] We acquire some clinical experience about how to minimize the side effect and how to institute the course of treatment.

**METHODS**

### Inclusion criteria

Subjects were required to present with the clinical signs and symptoms consistent with NM, including headache, confusion, cranial and spinal nerve involvement, nausea and vomit. CSF (200 μL) was collected from a lumbar puncture and it was centrifuged in (650 rpm) for 4 min using Slide Centrifuge (Shandon Cytospin 4, Thermo). Cell slides were May–Grunwald–Giemsa stained for 5 min then phosphate buffers were added and incubated for 10 min, followed by gentle rinsing with running water. The stained cell slides were observed under the microscope (Oil immersion lens ×1000, Olympus DP72). NM was diagnosed once tumor cells were found by experienced examiners as showed in Figure 1. Patients with cancer cells were allocated to intrathecal dexamethasone and methotrexate (IT DXM and MTX) group and palliative care group according to their families’ will.

Subjects were discontinuous cases from 2006 to 2014 who did CSF cytologic exams in the CSF cytological examination laboratory of the Second Hospital Affiliated to Hebei Medical University. Ethics approval was also given by the hospital review board.

### Treatment of neoplastic meningitis

After the NM diagnosis, the patients in IT DXM and MTX group received intrathecal dexamethasone 5 mg and methotrexate 10 mg, two doses a week as an inductive treatment of 4 weeks until the symptom was relieved or tumor cells reduced significantly in CSFC examination. Then the patients underwent treatments with a dehydrating agent, pain killer drugs, benzodiazepines, as well as other supportive treatments in the hospital. Then IT DXM and MTX was given one dose every 2 weeks in the outpatient department until the general condition severely deteriorated and could not sustain one’s life. Subjects in the palliative care group received supportive treatments in hospital or at home, according to the families’ determination based on the pain of lumbar puncture or economic reasons.

Intrathecal injections were conducted as follows: use of intravenous mannitol 250 mL was 20 min before lumbar puncture and remained throughout the process of lumbar puncture. The infusion apparatuses were readily available in case of use of emergency drugs. Ten milliliter CSF was slowly drained out of the subarachnoid space through a half-clogging needle for CSF examination. The needle was then returned into the cannulas. Dexamethasone sodium phosphate was diluted from 1 mL (5 mg) to 5 mL with physiological saline and then slowly injected into the subarachnoid space. During the injection, dexamethasone sodium phosphate was mixed with drawing back CSF repeatedly. Methotrexate was diluted to 5 mL and then injected the same way dexamethasone was treated.

### Data collection

The patients’ characteristics and treatment information at the diagnosis of NM were obtained in the medical record from the Second Hospital of Hebei Medical University. Survival data, subsequent therapeutic schedule, and side effects following discharge were obtained by making the phone calls to ask whether there is paralysis, severe vomiting, headache within 48 h after intrathecal injection, or the symptoms of bone marrow suppression, such as fever, infection, and low blood cell count. Overall survival was calculated from the diagnosis of NM.

### RESULTS

#### The patients’ characteristics

Twenty-three subjects were diagnosed as NM according to the positive CSF results as shown in Figure 1. Patient characteristics were summarized in Table 1. Eight patients received IT DXM and MTX treatment as IT DXM and MTX group, and 15 patients as the palliative care group was treated with palliative therapy, such as dehydrant drugs and painkillers.

Among them, 22 subjects showed high intracranial pressure (> 200 mmH2O), with other common presenting symptoms including inability to walk (n = 3), varying degrees of visual loss (n = 9), hearing damage (n = 4), sphincter disturbances (n = 4), seizure (n = 6), and confusion (n = 3). Furthermore, 1 patient received systemic chemotherapy, 2 received whole brain radiotherapy, and 1 received ventriculoperitoneal shunt (VP shunt) treatment. All patients showed positive results by CSFC exam.

#### Survival

Overall survival (OS) was assessed from the time of NM diagnosis to death and then Kaplan–Meier
analysis (group by treatment as IT DXM and MTX group/palliative care group; log-rank test) was conducted [Table 2 and Figure 2]. The OS of IT DXM and MTX group was significantly longer than that of the palliative care group \( (P = 0.01) \). The median survival of palliative care group is 7.53 weeks (5.5-9.57; \( n = 15 \)), and have the IT DXM and MTX group, 28.63 weeks (12.50-44.75; \( n = 8 \)), of the total patients, 14.87 weeks (7.93-21.81 weeks; \( n = 23 \)).

We collected the patients’ characteristics, symptoms, treatment method, and some laboratory examinations at the initial diagnosis of NM, including IT DXM and MTX, KPS, age, gender, primary tumor, cranial nerves damage, spinal nerves damage, seizure, confusion, and level of hemoglobin, albumin, and globulin. We analyzed the possible survival ratio of these factors using Cox’s proportional hazards regression model and the method of forward LR, by entering the factor when \( P < 0.05 \) and removing it when \( P > 0.06 \). IT DXM and MTX prolonged the OS of NM (regression coefficient = -2.923), odds ratio (OR) = 0.054 (0.09-0.323). Spinal nerves damage decreased the OS (regression coefficient = 1.595), OR = 4.928 (1.382-17.579). Other factors did not enter the Cox’s model (KPS, \( P = 0.935 \); age, \( P = 0.270 \); gender, \( P = 0.726 \); primary tumor, \( P = 0.220 \); cranial nerve damage, \( P = 0.564 \); seizure, \( P = 0.605 \); confusion, \( P = 0.485 \); hemoglobin level, \( P = 0.434 \); albumin level, \( P = 0.658 \); globulin level, \( P = 0.781 \)).

### Bias analysis

There are some innate biases in retrospective studies. Recall bias and confounding bias were the most important biases in our study. Recall bias is innate and uncontrollable, so the conclusion about IT treatment may be not well-grounded. For the latter bias, we analyzed some confounding factors between IT DXM and MTX group and palliative care group. We used KPS as a quantitative index of the subjects’ condition. There were no differences in KPS (2 independent samples \( t \)-test, \( P = 0.733 \)) and gender (Fisher exact \( t \)-test, \( P = 0.367 \)) between the groups, so we can exclude the imbalanced distribution of the KPS and gender and its effects on the different OS between the groups. Age of IT DXM and MTX group is higher than that of palliative group [Mann-Whitney \( U \), \( P = 0.043 \); 60.5 [56.5, 64.5] vs. 55 [44, 66]]. Aged patients present negative prognostic factors,[23,24] but IT DXM and MTX group had elder age and longer survival. This is possibly because of the different treatment methods. Given a small number of cases, we just compare the proportion of lung cancer and breast cancer, finding no difference between the groups (Fisher exact \( t \)-test, \( P = 0.685 \); \( P = 1.0 \), respectively), so we conclude that the primary cancer type was at an equilibrium distribution. Moreover, there were some biases coming from the researchers because this study didn’t involve blind method in experimental design.

### DISCUSSION

Neoplastic meningitis is a solid tumor at the advanced stage during which patients usually has severe pain and must administrate painkillers frequently. The diagnosis of NM often leads to palliative treatment that is intended to preclude the additional discomfort with aggressive treatment. Meningitis, seizure, vomit or sort of adverse effects were reported in the past studies. In our study, the patients in IT DXM and MTX group show no obvious side effects. This may be caused by a small number of cases, but we think that side effects could be decreased if the drugs were

### Table 1: The patients’ characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>IT DXM and MTX group/palliative care</th>
<th>Male/female</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>IT DXM and MTX group/palliative care</td>
<td>8/15</td>
<td>9/14</td>
<td>55 (21-67)</td>
</tr>
<tr>
<td>Presenting symptoms</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High intracranial pressure</td>
<td>22</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unable to walk</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visual loss</td>
<td>9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hearing damage</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sphincter disturbances</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Seizure</td>
<td>6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Confusion</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cancer type (IT DXM and MTX group/palliative care group)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lung cancer</td>
<td>4/6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breast cancer</td>
<td>1/2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gastric cancer</td>
<td>0/1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Malignant melanoma</td>
<td>0/1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>3/5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concurrent treatment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IT DXM and MTX</td>
<td>8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Systemic chemo</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WBRT</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VP shunt</td>
<td>1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**IT DXM and MTX:** intrathecal dexamethasone and methotrexate; **VP:** ventriculoperitoneal; **WBRT:** whole brain radiotherapy

### Table 2: Overall survival of different treatments

<table>
<thead>
<tr>
<th>Treatment</th>
<th>( n )</th>
<th>Median OS weeks</th>
<th>95% CI</th>
<th>( P ) (log-rank)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Palliative care group</td>
<td>15</td>
<td>7.53</td>
<td>5.50-9.57</td>
<td>0.01</td>
</tr>
<tr>
<td>IT DXM and MTX</td>
<td>8</td>
<td>28.63</td>
<td>12.50-44.75</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>23</td>
<td>14.87</td>
<td>7.93-21.81</td>
<td></td>
</tr>
</tbody>
</table>

**IT DXM and MTX:** intrathecal dexamethasone and methotrexate; **CI:** confidence interval; **OS:** overall survival
thoroughly diluted and slowly injected according to the course of treatment. Most of the patients’ discomfort was relieved in 3-4 weeks in IT DXM and MTX group with the decreased use of painkillers. IT DXM and MTX is well tolerant despite of the patients’ conditions. However, the patients with abnormal flow studies are associated with poor efficacy and intrathecal chemotherapy toxicity.\textsuperscript{[25]}

Apart from the low drug concentration, we conclude that good tolerance of IT DXM and MTX schedule is related to dexamethasone. Intrathecal steroid therapy can significantly reduce the IL-6 in CSF, a kind of inflammatory factor,\textsuperscript{[26] }so it may reduce nonspecific inflammatory reaction caused by tumor cells or chemotherapy agents. Dexamethasone has been reported as feasible and well tolerated with concomitant intrathecal liposomal cytarabine in patients with acute lymphoblastic leukemia.\textsuperscript{[27] }However, no prospective trials in adults with NM prove beneficial to use of intra-CSF glucocorticoids in combination with intra-CSF chemotherapy.

The natural processes of NM are disastrous if not well treated, for most patients will have a quickly deteriorated condition and die within 2 months. Intrathecal methotrexate is not a new therapy, but the random controlled trial is rare. IT DXM and MTX prolonged the patients’ survival significantly. The medium survival accords with William R and Theodore’s report,\textsuperscript{[4,11] }and is longer than that reported in Glantz’s study,\textsuperscript{[28] }in which most subjects were well treated, for most patients will have a quickly deteriorated condition and die within 2 months. Hence, flexible induction time should be discussed, we recommend two injections on a weekly basis for 4 weeks as induction treatment, and continue the treatment one more week if CSFC does not show a decrease in tumor cells. Should CSF relapse as a symbol of IT DXM and MTX termination? The answer is “no” by our experience. Restarting induction treatment could reduce the tumor cells in CSF with relieved symptoms, but randomized controlled trials with more clinical cases should be conducted to confirm this viewpoint. Moreover, new clinical trials of NM based on a tumor-specific histology are needed to establish the role of IT DXM and MTX treatment.

In conclusion, intrathecal dexamethasone and methotrexate are a safe and effective therapy. Although there are diversified intrathecal agents in recent years, other cytotoxic drugs and targeted agents such as trastuzumab\textsuperscript{[13] }and combined intrathecal chemotherapy\textsuperscript{[22] }prove efficient in treating NM. Thanks to the uncertain properties of new drugs, combined IT DXM and MTX as a basic treatment may be considered to ensure the therapeutic effect.

**REFERENCES**

3. Dalhaug A, Haukeland E, Nieder C. Leptomeningeal carcinomatosis from renal cell cancer: treatment attempt with radiation and


Received: 21-11-2014; Accepted: 08-06-2015