Safety and feasibility of switching from phenytoin to levetiracetam monotherapy for glioma-related seizure control following craniotomy: a randomized phase II pilot study

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Abstract Seizures are common in patients with gliomas, and phenytoin (PHT) is frequently used to control tumor-related seizures. PHT, however, has many undesirable side effects (SEs) and drug interactions with glioma chemotherapy. Levetiracetam (LEV) is a newer antiepileptic drug (AED) with fewer SEs and essentially no drug interactions. We performed a pilot study testing the safety and feasibility of switching patients from PHT to LEV monotherapy for postoperative control of glioma-related seizures. Over a 13-month period, 29 patients were randomized in a 2:1 ratio to initiate LEV therapy within 24 h of surgery or to continue PHT therapy. 6 month follow-up data were available for 15 patients taking LEV and for 8 patients taking PHT. In the LEV group, 13 patients (87%) were seizure-free. In the PHT group, 6 patients (75%) were seizure-free. Reported SEs at 6 months was as follows (%LEV/%PHT group): dizziness (0/14), difficulty with coordination (0/29), depression (7/14) lack of energy or strength (20/43), insomnia (40/43), mood instability (7/0).

Introduction

At diagnosis, ~20–40% of all brain tumor patients will have had a seizure [1] that may result in morbidity and decreased quality of life [2]. Thus, treatment with antiepileptic drugs (AEDs) is clearly indicated for brain tumor patients with preoperative tumor-related seizures [3–5]. Surgical treatment of brain tumors is also associated with a high risk of postoperative seizures. Approximately, one-third of patients have seizures after tumor resection [6–8], and in the case of supratentorial craniotomy, 20%–50% of patients have ≥1 seizure [9, 10].

Brain tumor patients who experience tumor-related seizures are often placed on phenytoin (PHT) therapy. Although there are no formal guidelines, it is common practice to continue PHT after craniotomy in these patients. However, PHT has many side effects (SEs) and undesirable interactions with drugs used to treat brain tumors [1, 11–14] which can complicate the pharmacological treatment of these tumors.

Levetiracetam (LEV) is a newer AED that possesses a different pharmacological profile from that of PHT, with no effects on liver enzymes and no known effect on the kinetics of other medications [15, 16]. Furthermore, other medications including enzyme-inducing AEDs have essentially no effect on LEV pharmacokinetics [15, 17]. In retrospective studies, LEV has been found to be effective and well tolerated as add-on or monotherapy in patients with either primary or metastatic brain tumors [18, 19]. Prospectively, LEV has been studied mostly as add-on therapy for persistent tumor-related seizures [20, 21], and
for this purpose, LEV appears to be safe and feasible and with few SEs.

Ideally, patients with brain tumor-related seizures would achieve seizure control with an AED that does not negatively interact with adjunctive chemotherapy and has few SEs. However, PHT is still commonly used as a first-line AED in brain tumor patients, and conversion of PHT therapy to LEV monotherapy has not been studied. Therefore, we prospectively studied the safety and feasibility of converting PHT to LEV monotherapy after craniotomy for glioma resection in patients with a history of tumor-related seizures.

Materials and methods

Key patient inclusion criteria for the study included: seizure history attributable to supratentorial glioma, PHT monotherapy for seizure prophylaxis, planned craniotomy for a history of epilepsy attributable to supratentorial glioma, PHT therapy, respectively (Fig. 1). Patients randomized to PHT therapy were followed postoperatively for 6 months, during which their medical charts were reviewed and two standardized telephone surveys were conducted at ~3-month intervals. Patients were provided with seizure diaries and a 24-h pager number to alert study personnel of any seizures or AED-related SEs. Seizure frequency data were based on patient-initiated contact of study personnel (through use of the 24-h pager), the telephone surveys, and review of medical charts. The primary end point was the proportion of patients who were seizure free 6 months after tumor resection. Data on AED-related SEs were collected with use of the telephone survey specifically addressing the SEs listed in Table 3.

Results

Characteristics of the study population are summarized in Table 1. Although there were some demographic differences between the two treatment groups, the majority of the study patients were male, suffered from generalized seizures, were diagnosed with glioblastoma, and all underwent craniotomy at the start of the study. One LEV subject died of the primary pathology before the first evaluation. Four other LEV subjects were withdrawn from the study by other physicians who preferred different AEDs; one patient was withdrawn by the neurosurgeon because the patient was a participant in another neuro-oncology trial, and PHT was the standard anticonvulsant in that study; the other three patients were withdrawn from this study by the neurosurgeon because of a personal preference for PHT as the anticonvulsant. One PHT subject could not be located after hospital discharge.

Fifteen patients in the LEV group and eight patients in the PHT group were assessed for the primary end point of seizure-freedom at 6 months (Table 2). Thirteen LEV patients (87%) were free from seizure compared with six PHT patients (75%). During the study period, all patients with follow-up were at least Engel Class 2 (rare seizures). Of the two patients who experienced rare seizures in the LEV group, one patient had stopped taking LEV, and this patient did not suffer additional seizures after resuming therapy; the other patient achieved seizure control with an increased dosage of LEV (1,500 mg twice a day). Most (11/15) LEV patients remained on the initial dose of 1,000 mg twice a day. Three LEV patients were able to lower their dosage to 500 mg twice a day without compromising their complete seizure control. PHT dosages were between 300 and 400 mg per day given as single dose. No patient required hospitalization for seizures or seizure control.

Table 3 shows the SEs reported in the telephone questionnaires administered ~3 and 6 months after surgery.
The most prominent difference between the LEV and PHT groups in terms of SEs was difficulty with coordination: no patient in the LEV group reported this difficulty, compared to 4 (57%) and 2 (29%) of the PHT group at 3 and 6 months, respectively. The one patient taking LEV 1,500 mg twice a day did not report more SEs than those taking 500 mg twice a day, suggesting that LEV dosage does not correlate with increased SEs. No SEs in any study patient resulted in a hospitalization or withdrawal from study.

Discussion

An ideal AED for brain tumor patients would be one that has few SEs and does not interfere with medications commonly used in brain tumor treatment. While PHT is very commonly used for seizure control in brain tumor patients, it has many undesirable pharmacological qualities. PHT induces the metabolism of the steroid dexamethasone [11, 13], and dexamethasone decreases PHT serum levels [22]. Certain chemotherapeutic agents
also lower PHT levels [12], and, vice versa, PHT can accelerate the metabolism of a variety of chemotherapy drugs including nitrosoureas, paclitaxel, 9-aminocamptothecin, thiopeta, topotecan, and irinotecan ([1, 14] and references therein). In terms of SEs, PHT adversely affects cognition [23, 24], which is often already depressed in brain tumor patients. Furthermore, brain tumor patients taking PHT are at increased risk of drug rashes [25], including Stevens-Johnson syndrome [26], an immune-mediated rash associated with severe morbidity and death.

Because LEV does not have any significant drug interactions, this AED poses fewer complications for the pharmacotherapy of brain tumors. A retrospective study of 34 patients with primary brain tumors suggested that LEV is effective and well tolerated for tumor-related seizure control, especially as add-on therapy to other AEDs [19]. Prospectively, LEV has been studied as add-on therapy for seizure control in brain tumor patients. In a cohort of 25 patients with primary brain tumors, Wagner et al. [21] found that add-on LEV reduced seizure frequency by more than 50% in 13 of the patients who experienced persistent seizures. Maschio et al. [20] prospectively followed 19 patients with primary brain tumors who had LEV added to existing AED therapy on account of persisting seizures; in that study, 14 patients had improvement in seizure control. There have been no reports of serious SEs of LEV therapy in any of the studies of brain tumor patients to date. For the reasons described above, safe conversion of PHT to LEV monotherapy may provide significant clinical advantages for the post-surgical medical treatment of primary brain tumors.

### Table 1 Characteristics of study population

|                         | Total patients n (%) | LEV patients n (% of arm) | PHT patients n (% of arm) |
|-------------------------|----------------------|---------------------------|---------------------------|
| Subjects enrolled       | 29 (100)             | 20 (69)                   | 9 (31)                    |
| Subjects with primary end-point follow-up | 23 (79)             | 15 (75)                   | 8 (89)                    |

#### Demographics

|                         |                       |                           |                           |
|-------------------------|-----------------------|---------------------------|---------------------------|
| Female                  | 6 (26)                | 6 (40)                    | 0                         |
| Median age (range)      | 45 (20–83)            | 46 (20–56)                | 39 (32–83)                |

#### Seizure type at presentation

|                         |                       |                           |                           |
|-------------------------|-----------------------|---------------------------|---------------------------|
| Simple partial          | 7 (30)                | 6 (40)                    | 1 (13)                    |
| Complex partial         | 5 (22)                | 1 (7)                     | 4 (50)                    |
| Generalized             | 11 (48)               | 8 (53)                    | 3 (38)                    |

#### Surgery

|                         |                       |                           |                           |
|-------------------------|-----------------------|---------------------------|---------------------------|
| Craniotomy before study surgery | 7 (30)           | 4 (27)                    | 3 (38)                    |
| Craniotomy for study    |                       |                           |                           |
| Gross total resection   | 13 (56)               | 10 (67)                   | 3 (38)                    |
| Subtotal resection      | 10 (43)               | 5 (33)                    | 5 (63)                    |

#### Final tumor pathology

|                         |                       |                           |                           |
|-------------------------|-----------------------|---------------------------|---------------------------|
| WHO grade 1             |                       |                           |                           |
| DNET                    | 1 (4)                 | 1 (7)                     | 0                         |
| WHO grade 2             |                       |                           |                           |
| Astrocytoma             | 3 (13)                | 2 (13)                    | 1 (13)                    |
| Oligodendroglioma       | 2 (9)                 | 1 (7)                     | 1 (13)                    |
| PXA                     | 1 (4)                 | 1 (7)                     | 0                         |
| WHO grade 3             |                       |                           |                           |
| Astrocytoma             | 4 (17)                | 4 (27)                    | 0                         |
| Oligodendroglioma       | 1 (4)                 | 0                         | 1 (13)                    |
| Oligoastrocytoma        | 2 (9)                 | 1 (7)                     | 1 (13)                    |

**WHO** World Health Organization (brain tumor grading scale)

**DNET** dysembryoplastic neuroepithelial tumor; **PXA** pleomorphic xanthoastrocytoma

### Table 2 Postoperative seizure status at 6 months

| Seizure status (Engel score) | LEV patients n (%) | PHT patients n (%) |
|-----------------------------|--------------------|--------------------|
| n at follow-up              | 15                 | 8                  |
| Class 1: seizure free       | 13 (87)            | 6 (75)             |
| Class 2: rare seizures      | 2 (12)             | 2 (25)             |
| Class 3: meaningful improvement | 0                 | 0                  |
| Class 4: no improvement     | 0                  | 0                  |

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In our study population, LEV and PHT groups had similar proportions of seizure-free patients (87% and 75%, respectively) at 6 months of follow-up. These seizure control rates are in line with those observed in a retrospective analysis of low grade glioma patients treated at this institution; of 253 patients who had tumor-related seizures, 169 (67%) were seizure-free 6 months after craniotomy for tumor resection [8]. Thus, LEV and PHT groups had expected levels of seizure control. Furthermore, no patient in our study had seizures refractory to either LEV or PHT therapy.

LEV and PHT were both well tolerated in our study population. Although difficulty with coordination is a potential SE for both LEV and PHT, no patient in the LEV study group (0/15) admitted to experiencing this difficulty while more than half of the PHT group (4/7) at 3 months reported this SE. Other known LEV SEs are somnolence and behavioral abnormalities. Both PHT and LEV groups had similar proportions of patients reporting somnolence. One subject in the LEV group reported feeling unusually hostile towards others after starting LEV therapy; however, this behavioral change did not require the discontinuation of LEV. No PHT study subject admitted to experiencing any emotional instability at any time point. In placebo-controlled trials of LEV for partial-onset epilepsy, LEV was associated with a 0.7% incidence of psychosis as compared to 0.2% for placebo patients. Although Maschio et al. reported that there were no LEV-related SEs observed in their study of primary brain tumor patients, Wagner et al. did report one patient in their study of 26 who developed a psychosis 4 weeks after initiating LEV therapy which resulted in discontinuation of this AED. Given the report of psychosis in Wagner et al. and our findings of emotional instability in up to 13% of our LEV group as compared to none in the PHT group (Table 2), we suggest that it would be important to evaluate brain tumor patients on LEV therapy for potential behavioral problems.

Our pilot data presented here suggest that it is safe and feasible to switch from PHT to LEV monotherapy after craniotomy for glioma-related seizures. Due to the small size of our study, the data presented here cannot serve as an official endorsement of LEV efficacy in this patient population. A large-scale, double-blinded, randomized controlled trial is warranted to compare PHT and LEV for seizure control equivalence and SE profiles.

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### Table 3 Results of patient questionnaire on SEs

|                      | 3 months postoperation | 6 months postoperation |
|----------------------|------------------------|------------------------|
|                      | LEV patients n (%)     | PHT patients n (%)     | LEV patients n (%) | PHT patients n (%) |
| **Number of respondents** | 15                    | 7                      | 15                  | 7                   |
| Questions regarding potential SEs |                       |                        |                     |
| Have you been experiencing any of the following symptoms? |                       |                        |                     |
| Dizziness          | 1 (7)                  | 0                      | 0                   | 1 (14)              |
| Headache           | 1 (7)                  | 1 (14)                 | 1 (7)               | 1 (14)              |
| Difficulty with coordination | 0                    | 4 (57)                 | 0                   | 2 (29)              |
| Excessive sleepiness | 3 (20)                | 2 (29)                 | 3 (20)              | 2 (29)              |
| Enlarged gums      | 0                      | 0                      | 0                   | 0                   |
| Depression         | 1 (7)                  | 1 (14)                 | 1 (7)               | 1 (14)              |
| Difficulty sleeping | 5 (33)                 | 1 (14)                 | 6 (40)              | 3 (43)              |
| Itchiness          | 0                      | 1 (14)                 | 0                   | 0                   |
| Nausea             | 0                      | 0                      | 0                   | 0                   |
| Vomiting           | 0                      | 0                      | 0                   | 0                   |
| Tingling, “pins and needles” sensation | 0                   | 1 (14)                 | 0                   | 0                   |
| Nervousness        | 1 (7)                  | 1 (14)                 | 0                   | 0                   |
| Emotional instability | 2 (13)                | 0                      | 1 (7)               | 0                   |
| Slurred speech     | 1 (7)                  | 0                      | 2 (13)              | 0                   |
| Lack of Energy or Strength | 4 (27)             | 4 (57)                 | 3 (20)              | 3 (43)              |
| Constipation       | 0                      | 1 (14)                 | 0                   | 0                   |
| Difficulty keeping balance | 0               | 0                      | 0                   | 0                   |
| Rash               | 0                      | 0                      | 0                   | 0                   |
| Have you been experiencing any other SEs that you attribute to your medication? | 3 (20) | 2 (29) | 2 (13) | 2 (29) |
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