TOPIRAMATE FOR THE TREATMENT OF JUVENILE MYOCLONIC EPILEPSY

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ABSTRACT - Objective: The aim of this study was to evaluate the efficacy and tolerability of topiramate (TPM) in juvenile myoclonic epilepsy (JME). Method: We assessed seizure control and adverse effects of TPM in 22 patients (18 females) aged 13 to 53 years. Target TPM dosage was up to 200 mg/day. The patients were subdivided into 3 groups: those treated with seizure control plus side effects (n=4); treated with non-controlled seizures (n=15) and with JME newly diagnosed (n=3). Results: Sixteen patients completed the first year of the follow-up. Generalized tonic-clonic seizures were completely controlled in 10 (62.5%); more than 50% of reduction in 4 (25.0%) and less than 50% in 2 (12.5%). Myoclonia were controlled in 11 (68.8%) and persisted in 5 (31.2%) patients. Absence seizures were present in 5 (22.7%) of whom 2 (9.0%) showed more than 50% of seizure reduction while 3 (13.6%) presented worsening. Discontinuations were due to inadequate seizure control and adverse events (N=4), low compliance and loss of follow-up (N=2) and subject choice (N=1). Conclusion: TPM showed to be an effective and well-tolerated drug in the treatment of JME. Although frequently observed, TPM side effects were tolerable and the drug could be maintained in the majority of patients.

KEY WORDS: topiramate, treatment, juvenile myoclonic epilepsy.

Tratamento da epilepsia mioclônica juvenil com topiramato

RESUMO - Objetivo: Avaliar a eficácia e tolerabilidade do topiramato (TPM) na epilepsia mioclônica juvenil (EMJ). Método: Avaliamos a resposta terapêutica e efeitos colaterais do TPM em 22 pacientes (18 mulheres) com idades entre 13 e 53 anos. A dose alvo utilizada foi até 200 mg/dia. Os pacientes foram divididos em 3 grupos no início do tratamento: aqueles com controle das crises mas que apresentavam efeitos colaterais (n=4); com crises não controladas (n=15) e com EMJ recém diagnosticada (n=3). Resultados: Dezessete pacientes completaram o primeiro ano de acompanhamento. Crises tônico-clônicas generalizadas foram completamente controladas em 10 (62.5%), tiveram redução maior de 50% em 4 (25,0%) e menor de 50% em 2 (12,5%). Mioclonias foram controladas em 11 (68,8%) e persistiram em 5 (31,2%) pacientes. As crises de ausências, presentes em 5 (22,7%) pacientes, tiveram redução maior do que 50% em 2 (9,0%) e agravamento em 3 (13,6%). A retirada do estudo foi devida principalmente ao controle inadequado das crises e efeitos colaterais indesejáveis (n=4), pouca adesão e perda do seguimento (n=2) e escolha do paciente (n=1). Conclusão: TPM foi considerada droga eficaz e bem tolerada no tratamento da EMJ. Apesar de frequentemente observados, os efeitos colaterais do TPM foram toleráveis e a medicação pode ser mantida na maioria dos pacientes.

PALAVRAS-CHAVE: topiramato, tratamento, epilepsia mioclônica juvenil.

Valproate (VPA) has been considered the drug of choice in the treatment of juvenile myoclonic epilepsy (JME). Different series reported good to excellent control of seizures in 70-85% of patients with VPA monotherapy or polytherapy. VPA is effective in the three types of seizures, generalized tonic-clonic seizures (GTCS), myoclonia and absences, which characterize this syndrome. Despite its satisfactory efficacy, side effects such as weight gain, polycystic ovary and hormonal disturbance have led to other therapeutic strategies.

Recently, topiramate (TPM), lamotrigine and zonisamide have also been shown to be effective in this syndrome. TPM may represent a therapeutic alternative in JME since it has an equally broad profile. In fact, some studies have demonstrated the efficacy of TPM in generalized seizures, including those of JME. Others have suggested that des-
Despite the effectiveness of TPM in the control of GTCS, its efficacy in the control of absences and myoclonic seizures is poorer\textsuperscript{4,6}.

This is a prospective study for evaluation of efficacy and tolerability of TPM in a series of patients with JME.

METHOD

In this study we included 22 JME patients followed up between March 1\textsuperscript{st}, 2003, and January 1\textsuperscript{st}, 2005 at Hospital São Paulo Epilepsy Outpatient Clinic, Universidade Federal de São Paulo. The Ethic Committee of the institution approved the protocol, and informed consent was obtained from all participants.

TPM was administered as a first drug or as a substitute drug in the treatment of the 22 patients (13-53 yr.; mean 23.2). Eight patients were being treated with VPA (500-2000 mg/day, in 2 of them associated with clonazepam 3 and 10 mg/day), 4 with CBZ (600-1400 mg/day), 2 PB (100 mg/day), one with oxcarbazepine (600 mg/day), one with lamotrigine (25 mg/day). Six patients were not being treated. These patients were divided into three groups. Groups 1 and 2 included patients already being treated with other AED and were constituted by: (a) patients with controlled seizures but presenting side effects (n= 4); (b) patients already receiving AED but with non-controlled seizures (n=15) and (c) patients with newly diagnosed JME (n=3). TPM was titrated according to patients' response over 12-14 weeks and maintained for an additional 32 weeks. Target TPM dosage was reached with the increase of 25 mg each 15 days up to 200 mg/day.

The doses of the initial therapy were reduced 25\% of initial doses per week. The exit criteria were occurrence of GTCS or intolerable side effects. The patients were evaluated in the 2\textsuperscript{nd}, 4\textsuperscript{th}, 6\textsuperscript{th}, 8\textsuperscript{th} and 12\textsuperscript{th} weeks and then in the 6\textsuperscript{th}, 9\textsuperscript{th} and 12\textsuperscript{th} months as to therapeutic effect and adverse events. Patients and family quantified the number of GTCS, myoclonia and absences per day in a diary. Psychiatric evaluation was performed in all patients and body mass index (BMI) was registered in each visit.

Statistical analysis – The small number of participants did not allow statistical comparison of TPM efficacy in previously treated and untreated groups. The comparison of those who presented seizure control, those with seizures not controlled and discontinued groups after TPM introduction was assessed by individual seizure type response and registered by frequency (%). TPM tolerability data was evaluated. Schedule Clinical Interview for DSM-IV (SCID I) was performed to measure depression and humor instability. BMI at the beginning and at the end of follow-up was compared using paired t-test of Student. We considered statistically significant p< 0.05.

RESULTS

Efficacy – Twenty-two patients (18 females) were enrolled, 16 of whom completed the first year of the follow-up (Figure). The mean dosage among study completers varied from 50 mg to 175 mg per day (mean 106.25± 55.43 mg/day). Suppression or reduction > 50\% of the frequency of seizures occurred in 15 patients (68.2\%). Sixteen patients com-

![Figure. Fluxogram of patient's follow-up.](image-url)
completed the first year of the follow-up. In relation to GTCS, present in 16 patients, there was complete seizure control in 10 (62.5%), more than 50% of GTCS reduction in 4 (25.0%) while 2 (12.5%) presented less than 50%. Myoclonia also present in 16 patients, were controlled in 11 (68.8%) and persisted in 5 (31.2%) patients. Absence seizures were present in 5 (27.7%) patients of whom 2 (9.0%) showed more than 50% of seizure reduction while 3 (13.6%) presented worsening of this seizure type. The frequency of myoclonia and GTCS is showed in Graphics 1 and 2. Most of the patients with controlled seizures plus side effects and those newly diagnosed had their seizures controlled. Among those who persisted having seizures despite the use of other AED before TPM introduction, myoclonia were improved in 46.6%, maintained the same frequency in 20% and showed aggravation in 33.3%, while GTCS were improved in 60% and aggravated in 40%.

Tolerability – The most prevalent side effect was weight loss. At the enrollment, 11 (50%) of the patients were over-weight (BMI > 25 kg/m²) while
7 (31.8%) were obese (BMI > 30 kg/m²). All overweight patients lost weight, 4 (36.4%) lost > 10% of weight, 6 (54.5%) lost 5-10% while 1 (9.1%) lost < 5%. Among those obese, 4 (57%) lost >10%. At the moment of this evaluation, the mean absolute variation of weight from the beginning to the end of the follow-up was -5.49±1.35 kg (p=0.001 paired T test) and the mean variation of BMI was -2.13±0.51 (p=0.001 paired t test).

The other side effects (Graphic 3) were represented by thirst/dry mouth (81.8%), paresthesias (68.2%), appetite decrease (40.9%), somnolence (31.8%), humor instability (27.3%), facial rubor (22.8%), headache/memor difficulty (18.2%), depression/language problems/attention difficulty/dizziness (13.7%), dry eyes/abdominal pain/upper respiratory infection (9.0%). Paresthesias and dry mouth were more common during the titration phase and tended to vanish after three months of therapy. These side effects were tolerable and the drug could be maintained in the majority of patients. Depression, language problems and attention difficulty were observed in 3 patients who were on TPM > 100 mg. Humor instability was registered in 6 patients who were on TPM 50-75 mg. These symptoms persisted with maintenance of the dosages; three of these patients had some benefit of fluoxetine 20 mg/day.

Discontinuations were either due to inadequate seizure control and adverse events (N=4) or low compliance and loss of follow-up (N=2) and subject choice (N=1).

**DISCUSSION**

TPM has proven to be effective in treatment of generalized epilepsy⁶. In patients with JME, it was initially shown that TPM was effective in cases refractory to other AED¹⁰, and later its efficacy was demonstrated in patients with recent onset epilepsy and intolerable side effects of other AED⁴,⁵,⁸.

In a previous study, Rosenfeld et al.⁴ evaluated the efficacy of TPM in 15 adults with JME. Fourteen of these patients had not achieved seizure control with VPA and/or presented excessive weight gain after institution of VPA therapy (5 subjects). TPM in an average dose of 663 mg/day improved seizure frequency in 11 patients, maintained control in two
and led to worsening in two. In the series of Prasad et al., TPM used in a minimum effective dose of 230 mg in monotherapy and 250 mg in polytherapy was considered to be an alternative to VPA. Tolerability to TPM was lower than to VPA, which the authors attributed to excessively rapid titration of the drug; in this series, one of the 19 patients showed worsening of the seizures. Levisohn et al. compared the efficacy of TPM vs. VPA in recently diagnosed JME patients. Twelve of 19 patients randomized to TPM concluded the study, and so did 7 out of the 9 patients in the VPA group. The average dose of TPM in the patients who concluded the study was 250 mg/day, in comparison to 750 mg/day of VPA.

In our series, TPM was administered in a lower dose (50 to 175 mg per day) than that of previous studies. This is a tendency in the most recent series presented that have shown minimization of side effects with maintained efficacy with low dose. In a series of drug naïve older patients with recently diagnosed partial-onset seizures Ramsay et al. showed that 50 mg of TPM as monotherapy, was similar to 200 mg. In the study of Edwards et al. TPM 100 mg/day was shown to be at least as effective as 600 mg/day of carbamazepine and 1250 mg/day of VPA in newly diagnosed generalized and focal epilepsies. These results agreed with previous observations such as that of Kwan and Brodie who mentioned the necessity of low-to-moderate dosages of AED in patients successfully treated with monotherapy.

In our series, TPM reduced the seizure frequency in approximately 2/3 of patients with JME in the first year of follow-up, seeming more effective in the control of GTCS and myoclonia than in absence seizures, despite the small number of patients with this last seizure type. However, this study shows that TPM may aggravate GTCS and myoclonia although this was observed only in the group of patients with more refractory seizures. Side effects were frequently observed in all patients. Some of them, such as thirst and paresthesias, tended to diminish with time, while other dose-dependent effects such as emotional instability, depression and language difficulties were persistent in patients with dosage over 100 mg/day.

Although frequently observed, TPM side effects were tolerable and the drug could be maintained in the majority of the patients.

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REFERENCES