Topiramate-Induced Acute Myopia, Diplopia, and Photosensitivity: A Case Report

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Introduction
Topiramate is primarily used as an antiepileptic drug, both as monotherapy and as an adjunct therapy in the control of partial and primary generalized epilepsy in children and adults (1). Effectiveness has also been demonstrated in migraine prophylaxis, depression, trigeminal neuralgia, bipolar disorders, eating disorders, and idiopathic intracranial hypertension (2). Potential ocular side effects of acute-onset angle closure glaucoma (ACG), acute myopia, nystagmus, diplopia, photosensitivity, suprachoroidal effusions, peri-orbital edema, and blepharospasm have been reported in recent articles (3). This report describes a rare case with acute myopia, diplopia, and photosensitivity associated with topiramate use.

Case Report
The patient was a 24-year-old woman who presented with a sudden loss of vision in both eyes. According to her medical history, there were no vision disturbances and she had not even used glasses. Her uncorrected visual acuity was 20/400 in both eyes and the best corrected visual acuity was determined to be 20/25 in the right eye with -5.50 spherical refractive correction and 20/20 in the left eye with -6.25 spherical refractive correction. On the second day of examination, diplopia developed. The patient’s clinical condition was considered to be related to the drug and topiramate was discontinued immediately. The clinical findings of the patient subsequently improved rapidly without treatment. On the fifth day of examination, her autorefractometry measurements were +0.25 -0.25 α 121° in her right eye and +0.25 in her left eye and her uncorrected visual acuity was 20/20 in both eyes with normal bilateral anterior chamber depth. She had no vision complaint or diplopia but she began to experience photosensitivity, which persisted for 4 months before regressing completely. When ophthalmologists encounter acute myopia and acute-onset ACG, especially in young patients, they should keep the use of topiramate in mind.

Keywords: Acute myopia, diplopia, photosensitivity, topiramate.
Scheimpflug image showed anterior chamber shallowness, lens thickening, and anterior transposition of the iris-lens diaphragm (Fig. 2a). Intraocular pressure assessed with a Goldmann applanation tonometer was 22 mmHg in her right eye and 19 mmHg in her left eye. A fundus examination was entirely normal; the cup-to-disc ratio in both eyes was 0.2. Gonioscopy revealed grade 2 occludable angles bilaterally. The results of orbital magnetic resonance imaging were normal. The visual field test results were normal bilaterally, with no scotoma/visual field defect. When she was questioned about diseases and medications, it was discovered that she had epilepsy and that a neurologist had recently initiated the use of topiramate (Topamax, 25 mg, 2x daily). Twelve days later the patient began to experience blurred vision. With the thought that the patient’s clinical status could be drug-related, her neurologist was consulted and topiramate was replaced with another medication. The patient was examined daily and no additional medication was required. On the second day of examination, however, she complained of diplopia; her autorefractometry measurements were -4.00 -0.25 α121° in her right eye and -3.50 -0.50 α130° in her left eye. On the fifth day of examination, her autorefractometry measurements were +0.25 -0.25 α121° in her right eye and +0.25 in her left eye. Her uncorrected visual acuity was 20/20 in both eyes and the anterior chamber depth was normal (Fig. 2b). Intraocular pressure assessed with a Goldmann applanation tonometer was 17 mmHg in the right eye and 15 mmHg in the left eye. She had no vision complaint but she began to experience photosensitivity (pupil sizes were normal; the right and left pupil diameter was 2.1 mm and 2.3 mm, respectively, in photopic conditions). One month after her first visit, the ophthalmologic examination
was completely normal, but the photosensitivity remained. The patient was advised to use polaroid glasses, and she was examined at monthly intervals. At the fourth month, the photosensitivity had receded completely.

**Discussion**

Topiramate is a sulfamate-substituted monosaccharide derived from D-fructose. It is used mainly as an antiepileptic drug for both monotherapy and adjunct therapy in the control of partial and primary generalized epilepsy in adults and children (1). Efficacy has also been reported in migraine prophylaxis, trigeminal neuralgia, bipolar disorders, depression, and eating disorders (2). Recently, it has been used in the treatment of idiopathic intracranial hypertension (4). It works via inactivation of sodium and/or calcium channels, hyperpolarization of K+ currents, inhibition of kainate receptor-mediated conductance, and activation of gamma-aminobutyric acid postsynaptic receptors (5). In addition, it also has weak anti-carbonic anhydrase activity. The mean plasma elimination half-life of the drug is about 21 hours. The half-life of the drug is closely related to the return of ocular side effects after discontinuation of the drug (5). Recent articles have described ocular side effects of topiramate, which include acute angle closure glaucoma (AACG), acute myopia, diplopia, blepharospasm, suprachoroidal effusions, peri-orbital edema, scleritis, oculogyric crisis, periorcular pain, and nystagmus (3). The exact mechanism by which topiramate initially triggers myopia is not completely understood. To date, a number of mechanisms have been suggested as possible triggers. One proposed mechanism is that fluid movements result in uveal and ciliary effusion because of the ability of topiramate to block sodium channels and change the membrane potential (6). Sen et al. (7) suggested that the entry of topiramate into the lens alters the osmotic status of the lens, causes the lens to swell and, consequently, results in AACG and myopia. In 2002, Ikeda (8) described side effects of topiramate therapy ranging from transient myopia to severe bilateral ACG. These complications attributed to ciliochoroidal effusion and swelling of the ciliary body can result in anterior rotation of the ciliary processes, narrowing of the ciliary sulcus, and forward displacement of the iris and lens. This has been demonstrated using standard or high frequency ultrasound in patients using topiramate (3). Another theory is that ciliochoroidal effusion syndrome may be due to an idiosyncratic reaction. This complication occurs at a low frequency, and some studies have reported no correlation between topiramate dosage and the level of intraocular pressure or myopia (3). Some studies have suggested that the onset of AACG with topiramate use is not dose-related and that AACG occurs with doses ranging between 50 mg to more than 100 mg topiramate (3). In our case, the patient’s complaints began on the 12th day of 50 mg/day topiramate use. Furthermore, pre-existing hypermetropia is not a prerequisite for development of AACG secondary to topiramate therapy (9). The incidence or prevalence of AACG in the population of patients treated with topiramate is not known. The condition has predominantly been reported in women (89%) (10). Some of these patients may be using selective serotonin reuptake inhibitors, which may aggravate the glaucoma by adding an element of pupillary block (11). Topiramate-induced AACG usually occurs within 2 weeks of the initiation of treatment (11). A number of patients taking topiramate have presented with characteristic findings of an acute attack of ACG, including blurred vision, headaches, nausea, and vomiting.

The acute myopia associated with using topiramate has been reported at between 2 and 8.75 diopters and causing sudden bilateral blurring of vision (12). Our case had -5.50 diopter myopia in her right eye and -6.25 diopter myopia in her left eye. The severity of ciliary body edema, ciliochoroidal detachment, and forward movement of the iris lens diaphragm causes myopia. Myopia on its own resolves following discontinuation of the drug, but occasionally myopia may persist after resolution of AACG. Drug-induced myopia has also been associated with sulfa drugs, such as acetazolamide, sulfamethoxazole/trimethoprim, indapamide, promethazine, spiranolate, isosorbide dinitrate, bromocriptine, as well as other drugs, including tetracycline, corticosteroids, hydrochlorothiazide, penicillamine, quinine, metronidazole, isotretinoin, and aspirin (13).

Diplopia and nystagmus have been reported in 14% to 15% of patients using high doses of topiramate (3). Although the dose of the drug was low (25 mg 2x daily) in our case, diplopia developed. Scleritis has been reported in 4 cases, oculogyric crisis in 2 cases, and single cases of blepharospasm, myokymia, periorcular edema, paresthesia, and periorcular pain have been described (3). A recent report demonstrated visual field defects with the usage of topiramate without the presence of elevated intraocular pressure (14).

A photosensitivity reaction following topiramate use has been more frequently observed in females (80%) and the condition is more common in patients between 30 and 59 years old (15). Photosensitivity is usually detected among people who have been taking the drug for 1 to 6 months (15). In the literature, the incidence of photosensitivity in patients using topiramate was found to be 0.12% (15). The mechanism of photosensitivity seen in patients using topiramate is not known (15). The photosensitivity in our patient began on the 10th day of usage and ended in the fourth month of discontinuation of the drug.

In the case of topiramate-associated AACG and acute myopia development, the drug should be discontinued immediately and an alternative drug should be prescribed by the primary physician. The initial treatment should include cyclo-
plegic agents to posteriorly displace the iris-lens diaphragm. Furthermore, topical and systemic ocular hypotensives and topical steroids can be used. Acetazolamide, a sulfonamide derivative drug, is not recommended for fear of further ciliary body edema. Also, topical miotics are contraindicated as a treatment option because they can cause further displacement of the iris-lens diagram, and they can lead to a relative pupillary block and worsen the ACG.

Peripheral iridotomy is not recommended in drug-induced secondary angle closure because the mechanism of this condition is not pupillary block; therefore, it has no therapeutic value in these cases. Laser peripheral iridotomy has been used in 23% of reported cases, but has not been sufficiently effective in relieving the secondary angle closure and should therefore be reserved for cases where the other treatments fail (3, 9). Topical steroids may be beneficial to reduce choroidal effusion by stabilizing cell membranes. Rapid resolution of an attack has been reported with the use of intravenous methylprednisolone and mannitol (16).

Parents of children and other patients who are to start topiramate treatment should be warned of the possible ocular side effects. Symptoms such as myopia and angle closure should be resolved with topical and oral intraocular pressure lowering medications, topical cyclopentolate, and discontinuation of topiramate. Physicians should inform patients to promptly report any symptoms of eye pain or blurred vision after commencing topiramate therapy, since if the symptoms remain unrecognized as a drug-related event, permanent ocular damage can occur. When ophthalmologists encounter acute myopia and acute angle closure glaucoma, especially in young female patients, they should closely examine the medical history of the patient and keep the use of topiramate in mind.

Disclosures
Informed consent: Written informed consent was obtained from the patient for the publication of the case report and the accompanying images.

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