Verapamil induced gingival enlargement in cluster headache

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Verapamil is an effective prophylactic treatment for cluster headaches and, therefore, is widely used. This report describes four patients with cluster headache who developed gingival enlargement after initiating treatment with verapamil. In two patients, it was possible to control this side effect adequately by optimising oral hygiene and dental plaque control. In the other two patients, lowering of the verapamil dose, in addition to optimal oral hygiene and dental plaque control, was necessary; in one patient verapamil had to be stopped completely to reverse the gingival enlargement. Doctors treating cluster headache with verapamil need to be aware of this side effect, especially as it may be preventable with good dental hygiene and dental plaque control, is reversible with reduction or cessation of verapamil, and can lead to dental loss.

CASE REPORTS

Case 1

A 32 year old woman developed intermittent daily headaches in 1997 for which she was referred the following year. She was diagnosed as having chronic cluster headache. Her medications included subcutaneous sumatriptan 6 mg prn (when required), which abbreviated the attack, and sodium valproate. In the past she had tried methysergide, sumatriptan tablets 100 mg prn and ergotamine tablets 2 mg prn, all of which were completely ineffective. Sodium valproate was discontinued and the patient started on verapamil, the dose of which was gradually increased to 760 mg daily under regular electrocardiogram (ECG) monitoring. There was a marked improvement with verapamil, resulting in a significant reduction in the frequency, duration, and severity of the cluster attacks. The only side effects she initially reported were mild constipation, lethargy, and ankle oedema.

About three months after starting verapamil she noticed that her interdental papillae had enlarged. A further three months later she saw a dentist who diagnosed gingival enlargement secondary to verapamil. However, as verapamil had dramatically improved the cluster headaches, the patient was reluctant to discontinue it. Dental plaque was removed by scaling and the patient was advised to maintain good oral hygiene. Over the next 18 months, despite regular professional dental care, the gingival enlargement continued to worsen gradually, with development of generalised nodular swelling and encroachment of the crowns of the adjacent teeth. In addition, the patient reported intermittent bleeding from the gums, especially after meals. In May 2000, she was persuaded to try other cluster headache preventive agents, while continuing verapamil. She had trials of lithium 400 mg twice daily and topiramate 200 mg twice daily, with which there was no improvement. During this period, the gingival enlargement continued to worsen and, in addition, the patient began to complain of discomfort from her teeth, especially when eating and looseness of the upper incisors. In January 2001, a course of intravenous dihydroergotamine 9 mg over three days was administered with dramatic improvement. The patient now had only two to four cluster attacks per week with each attack lasting only about 15–30 minutes. Verapamil and lithium were stopped without any deterioration in the cluster headaches. Over the next three months, the gingival enlargement, gum bleeding, and dental discomfort and looseness of teeth resolved completely.

Case 2

A 40 year old man developed cluster headaches in 1992 for which he was referred in June 2001. He was diagnosed as having chronic cluster headache. His medications included subcutaneous sumatriptan 6 mg prn, high dose and flow rate oxygen, verapamil 240 mg daily, sodium valproate 600 mg daily, lithium 300 mg daily, ergotamine tablets 2 mg daily,
methysergide 1 mg daily, and prednisolone 10 mg daily. Subcutaneous sumatriptan and oxygen were effective, abbreviating the attacks to 15–30 minutes. The combination of verapamil, sodium valproate, lithium, ergotamine, methysergide, and prednisolone was ineffective in suppressing the cluster attacks. The patient had started verapamil in December 2000. In the past he had tried indomethacin 150 mg daily and amitriptyline.

Sodium valproate, lithium, and prednisolone were discontinued. The verapamil dose was gradually increased to 600 mg daily under regular ECG monitoring over two months; there was mild improvement in the cluster headaches. He was then seen elsewhere and lithium 600 mg daily, methysergide 2 mg daily, and prednisolone 15 mg daily were added. In addition, the patient was administered intravenous dihydroergotamine (IV DHE) 17.5 mg over seven days; he was rendered pain free while receiving the IV DHE but the attacks recurred within a day of stopping the infusion. There was no added benefit with this combination of drugs.

He consulted again in September 2001, when he was advised to stop lithium, methysergide, and prednisolone. The verapamil dose was increased to 720 mg daily, which completely suppressed the cluster attacks. Unfortunately, two months later the patient noticed gingival enlargement with intermittent bleeding. We advised him to reduce the verapamil dose to 480 mg daily and to see a periodontist. The periodontist diagnosed gingival enlargement secondary to verapamil; the patient was advised to maintain good oral hygiene and given a course of antibiotics for possible infection, although there was no evidence for this. On reducing the verapamil dose, the cluster attacks recurred but, over the next two months, the gingival enlargement resolved completely. Interestingly, in May 2002, he took 1 g of “magic mushrooms” (containing psilocybin) which rendered him pain free for one month; since then he has been taking magic mushrooms 1 g once every one to two months, which renders him pain free for two to six weeks. He continues to take verapamil 480 mg daily and has no gingival problems.

**Case 3**

A 37 year old man developed intermittent daily headaches in 1996 for which he was referred to our clinic and was diagnosed as having chronic cluster headache. In the past he had taken sumatriptan tablets 100 mg prn, pizotifen, atenolol, amitriptyline, carbamazepine, indomethacin 50 mg three times daily, paroxetine, and diazepam, none of which produced any benefit. The patient was started on verapamil, the dose of which was increased to 400 mg daily under ECG monitoring over one month. Verapamil 400 mg daily completely suppressed the cluster headaches. He reported no side effects. Reduction of the verapamil dose led to recurrence of the headaches.

One year after starting verapamil, the patient noted the onset of gingival enlargement. Two months later, he developed bleeding from the gums, especially after meals and when brushing his teeth; in addition, the gingival enlargement continued to worsen gradually. A further two months later, he began to complain of discomfort from his teeth when eating. He then saw a dentist who diagnosed gingival enlargement secondary to verapamil. Dental plaque was removed and the patient was advised to maintain good oral hygiene by thoroughly brushing his teeth twice a day and rinsing his mouth with plain water after each meal. The possibility of substituting verapamil with another cluster headache preventive agent was raised but the patient declined the offer. Over the next six months all the dental symptoms gradually resolved.

**DISCUSSION**

Gingival enlargement or overgrowth is a side effect associated with the administration of several drugs. These drugs can be basically divided into three groups: anticonvulsants, the immunosuppressant ciclosporin, and calcium channel blockers. Recent therapeutic trends in the treatment of, particularly chronic, cluster headache have led to more widespread use of very high doses of the medicine, up to 960 mg daily, to be used more commonly. Our cases illustrate that verapamil can produce gum enlargement and given the exceedingly high doses of the medicine used in neurological practice it seems important to be aware of this problem and of its management.

Of the calcium channel blockers, the commonest agent associated with the development of gingival enlargement is nifedipine. Though similar problems have been associated with the administration of verapamil, felodipine, nitrendipine, diltiazem, and amlodipine. The incidence of
verapamil induced gingival hyperplasia is poorly defined. The only study that has addressed this issue identified one patient with gingival enlargement out of 24 dentate patients who used verapamil for more than one year, giving an incidence of 4.2%. This is lower than the 14–83% reported incidence of nifedipine induced gingival enlargement. Currently, the aetiology of drug induced gingival enlargement is not entirely understood, but it is clearly multifactorial. Some of the known risk factors include: presence of gingival inflammation (gingivitis due to poor oral hygiene); presence of dental plaque; and the dose and duration of drug therapy. 4.5–24

Gingival enlargement may cause significant morbidity because it poses an oral hygiene and dental plaque control problem; the tooth discomfort may affect mastication; it may alter tooth eruption; it may interfere with speech; and it may cause aesthetic concerns. 25

The term “gingival hyperplasia” is inappropriate because enlargement does not result from an increase in the number of cells but rather an increase in extracellular tissue volume with an inflammatory infiltrate of predominantly B lymphocytes. 26 Histologically, in verapamil induced gingival enlargement there is highly vascular connective tissue, acanthotic and thickened epithelium with long rete pegs containing dyskeratotic pearls, and varying amounts of subepithelial inflammatory infiltrate. 27 The histological picture is strikingly similar to that caused by phenytoin, ciclosporin, and other calcium channel antagonists. The mechanism by which drug induced gingival enlargement occurs is not well understood and may be distinct for each drug. Cell culture studies on gingival fibroblasts from a patient with verapamil induced gingival overgrowth and from control cells obtained from healthy gingiva suggest that verapamil affects the proliferation of selected fibroblasts subpopulations and alters the balance between regeneration and degeneration. 28

In the treatment of drug induced gingival enlargement, the first consideration should be given to the removal of local factors. The clinician should emphasise the importance of dental plaque control. Although the exact role played by dental plaque in drug induced gingival enlargement is unclear, there is evidence that good oral hygiene and frequent professional removal of plaque decreases the degree of gingival enlargement present and improves overall gingival health. 29 The possibility of discontinuing the offending drugs or of changing the medication should be raised. 30 Discontinuation of the drug usually results in complete regression of the gingival overgrowth. The options of discontinuing or substituting the medication should be examined in conjunction with the patient. Simple discontinuation of the offending agent is usually not a practical option but replacing it with another medication might be. If the non-surgical approach is not effective, periodontal surgery in the form of gingivectomy or periodontal flap procedures can effectively reduce the enlarged gingival tissues. 31

We have described four cases of patients with cluster headache who were treated with verapamil and subsequently developed gingival enlargement. The gingival enlargement was first noted at between three months and two years of starting verapamil. In cases 3 and 4, it was possible to reverse the gingival enlargement with optimum oral hygiene and dental plaque control, without altering the verapamil dose. In case 2, good oral hygiene and dental plaque control together with alterations in the verapamil dose were required to adequately control the gingival symptoms. In case 1, the symptoms were progressive despite good oral hygiene and dental plaque control; verapamil had to be stopped to reverse the symptoms.

These case reports highlight the importance of appreciating that verapamil can cause gingival overgrowth. Patients being considered for treatment with verapamil should be made aware of this potential side effect and encouraged to maintain meticulous oral hygiene. They should be advised to regularly consult an oral medicine specialist for control of dental plaque and monitoring for oral complications associated with gingival enlargement such as the gingivitis, bleeding gums, and loosening of teeth. In patients who develop gingival enlargement, it should be borne in mind that the gingival symptoms may be controlled successfully, even under the continuous administration of verapamil, by meticulous individual and professional oral hygiene. If this is not effective, then the verapamil dose can be reduced or the drug stopped completely. The surgical options should only be considered as a last resort in patients responsive to verapamil and unresponsive to other cluster headache preventive treatments in whom newer surgical approaches cannot be easily or appropriately considered. 32

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REFERENCES


ECHO

Dronabinol reduces central pain in MS

Patients with central neuropathic pain associated with multiple sclerosis (MS) should have better prospects of pain relief now that a randomised controlled trial in Denmark has shown that an oral synthetic derivative of cannabis is effective. This is a hopeful sign for those whose pain does not respond to current drugs.

The randomised double blind, placebo controlled crossover trial in 24 patients with MS and central pain established that dronabinol up to 10 mg daily for three weeks reduced pain intensity significantly by the end of treatment compared with placebo. The estimated relative difference in pain reduction from baseline between dronabinol and placebo was 20.5% (95% confidence interval −37.5 to −4.5). Median pain relief score was significantly raised for dronabinol (3 (0–6.7)) versus placebo (0 (0–2.3)) too, and mental health was better but functional ability was unchanged. Side effects were widespread, affecting the CNS and including musculoskeletal problems, though these were mostly tolerable once dosage was lowered. The patients were aged 23–55 and had had MS and pain for a median of seven and 4.5 years, respectively.

Cannabis derivatives seem to reduce allodynia or hyperalgesia in neuropathic pain and central pain in animals, and they might be helpful for pain suffered by up to 80% of MS patients. However, until now it was not known whether they would be effective specifically against central pain reported in about a third of MS patients.