Magnesium, Feverfew, and Riboflavin: Therapeutic Use in Migraine Prevention

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Case Reports on Patients Diagnosed with Migraine and Placed on A Nutraceutical Product Containing Riboflavin, Magnesium and Feverfew

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Approximately 28 to 32 million Americans suffer from migraine headaches. The majority of these are women between the ages of 22 and 35 years. Although the causes of migraine headaches are not clearly understood, it is believed to result from interplay between the brain tissue and the circulatory system supplying the central nervous system.

In a survey study of 20,000 individuals, about two-thirds of those who met the criteria for migraine, self-treated with over-the-counter (OTC) drugs to the exclusion of prescription medications.1 The desirability of prevention is clear to healthcare professionals and patients alike. In addition to obvious financial savings that effective prevention would generate, medical effects of medication leading to rebound headaches could potentially be reduced as well.

PATHOPHYSIOLOGY OF MIGRAINE

The induction of the migraine appears to be triggered by interactions between precipitating events and specific brain areas. These events appear to induce constriction of the cranial vessels that leads to a decrease in blood flow to the brain, which may induce periods of low oxygenation that can result in neurologic disjunction. Decreased oxygenation may be associated with the aura that is experienced in about 10% of migraine sufferers. Migraine auras manifest as perceptions of flashing lights, partial field of vision loss, unilateral numbness, weakness, or speech difficulty.

About 85% of patients will report migraine without aura. This most common type of migraine lasts about 4–72 hours, and is aggravated by physical activity, occurs unilaterally, and can be of moderate to severe intensity. Nausea, vomiting, photophobia, and or phonophobia may also occur.

The initial vasospasm of the cerebral event is believed to be mediated by the concentration of prostaglandins and other CNS neurotransmitters, including serotonin. In contrast, neurogenic inflammation is mediated by vasoactive neuroproteins. The pain associated with migraine headache usually occurs as a result of cerebral vasodilation and neurogenic inflammation that follows a period of decreased oxygenation. (Figure 1)

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Figure 1. Vasodilation of the cerebral vessels may result in migraine-related pain.
Psychological factors such as stress and depression have been associated with the onset of migraine, as have environmental factors like smoke, light, and weather changes. The consumption of large amounts of alcohol, citrus fruits, aspartame, chocolate, and caffeine may also trigger episodes. A number of medications have also been associated with migraine-related headaches. These include Tagamet, Pondium, Prozac, Premarin, Indocin, nicotine, nitroglycerin, oral contraceptives, reserpine, and ethinyl estradiol.4

FEVERFEW

Feverfew, Tanacetum parthenium, member of the Compositae or daisy family, is native to the Balkan mountains in Europe and can now be found in Australia, China, Japan, North and South American, and North America.2 Its use as an herbal remedy dates back 2,000 years.14 Feverfew currently is used for migraine headache prophylaxis and to treat rheumatoid arthritis.2

Chemistry

Some components in feverfew are:

- Sesquiterpene lactones: parthenolides, canin, artecanin, santamarin
- Flavonoid glycosides: luteolin, tanetin, apigenin, 6-hydroxy flavonols
- Sesquiterpenes and monoterpenes: camphor, borneol, germacrene, pimenes
- Other: polyacetylenes, pyrethrin, melatonin, tannins, essential oils, chrysanthemum acetate

The leaf is used for medicinal purposes. The most active component of feverfew is parthenolide, the most abundant sesquiterpene lactone.2

Mechanism of Action

The exact mechanism of action of feverfew is unknown; however, there are several proposed mechanisms: inhibition of serotonin release, inhibition of prostaglandin synthesis, inhibition of platelet aggregation and secretion, inhibition of polymorphonuclear leukocyte degranulation, inhibition of phagocytosis of human neutrophils, inhibition of mast-cell release of histamine, cytotoxic activity against human tumor cells, and antimicrobial activity; it also has antithrombotic potential.2

Inhibition of serotonin release

The active moiety in feverfew (parthenolide) inhibits the release of serotonin from blood platelets, similar to the action of methysergide, an ergot alkaloid.3 This mechanism explains the therapeutic benefit of feverfew for migraine.5,12

Inhibition of prostaglandin synthesis

The interference of phospholipase A by the plant causes inhibition of prostaglandin biosynthesis.10 In a study reported in 1985, feverfew decreased prostaglandin production by 86–88%, but cyclooxygenase inhibition was unaffected.4

Inhibition of platelet aggregation and secretion

Feverfew has been found to inhibit platelet aggregation by inhibiting thromboxane synthesis, which occurs via inactivation of cellular phospholipases.6,8 Numerous studies have shown that feverfew extract inhibits aggregation and secretion of intracellular granules caused by aggregating agents such as adrenalin, collagen, and adenosine diphosphate.8

Inhibition of polymorphonuclear leukocytes

Feverfew extract inhibits the secretory activity of polymorphonuclear leukocytes (PMN), showing a much greater inhibition of PMNs granule release than do high concentration NSAIDs.5

Smooth muscle

Sesquiterpene lactones, particularly parthenolide, have spasmolytic properties that cause smooth muscle to become less responsive to endogenous substances such as acetylcholine, noradrenaline, bradykinin, prostaglandins, histamine, and serotonin. These findings can be linked to an anti-migraine effect through inhibition of the influx of calcium in the vascular smooth muscle.13

Contraindication

Feverfew is contraindicated in people who may be allergic to other members of the Compositae or daisy family, such as chamomile, ragweed, or yarrow. It should not be used during pregnancy or by lactating mothers, or by children under two years old. Because parthenolide affects platelet aggregation in some in vitro studies, caution may be appropriate for patients with bleeding disorders or those anticipating surgery.2,3

Drug Interactions

NSAIDs may alter the efficacy of feverfew. The plants’ mechanism of action (mainly inhibition of platelet aggregation) may interact with anticoagulants and antiplatelets such as warfarin and aspirin.2,3

Adverse Effects

The most common reported side effects of feverfew consumption are GI disturbances after oral ingestion including diarrhea, heartburn, bloating and flatulence. Mouth ulcers, lip swelling, and tongue irritation can occur when one chews fresh leaves.2 When feverfew is discontinued after use of six months or more, people may experience rebound headache, stiffness in joints and muscles, nervousness, anxiety and insomnia, responses known as “post-feverfew syndrome.”10

Dosing

Currently there are no precise dosing recommendations, and doses vary. However, the Canadian Health Protection Branch recommends 125 mg of feverfew daily, containing at least 0.2% parthenolide (the active component) in each dosage unit.2,3,5 Feverfew dosage is generally based on the weight of the leaves. Commonly-used doses
in herbal practice are:
• Feverfew leaves: 2–3 daily
• Dried powdered leaves: 50–250 mg daily or 125–250 mg daily in two divided doses for migraine prophylaxis
• In the UK and Canada, feverfew products are standardized to contain at least 0.2% parthenolide

Clinical Evidence
Various studies have shown positive therapeutic outcomes of feverfew monotherapy for migraine prophylaxis; however, a few studies have shown no clinical benefit. In the table below are data collected from five different clinical trials obtained from a systematic review conducted in 1998.¹⁰

MAGNESIUM
Magnesium is the second most abundant intracellular cation in the body and the most common intracellular divalent cation and a cofactor in hundreds of enzymatic processes; its central role in smooth muscle activity and peripheral vascular resistance is well known.¹⁵ About 65–70% of serum magnesium is ionized, while the rest is protein-bound and complexed to small anion ligands.² Many studies of the role that magnesium plays in the pathogenesis of migraines have examined the total body supply of intracellular magnesium; these conflicting results may be due to the fact that even though total intracellular magnesium content is relatively stable, there are wide fluctuations in serum ionized magnesium. It is this ionized portion that affects the physiological component of a migraine. Migraine research found multiple relationships between magnesium deficit and migraine attacks.¹⁶

Pharmacology
Magnesium is an electrolyte necessary in a number of enzymatic processes, phosphate transfer, muscular contraction, and nerve conduction. Deficiencies have been documented in malabsorption syndromes, prolonged diarrhea, vomiting, pancreatitis, aldosteronism, kidney dysfunction, chronic alcoholism, and diuretic therapies.

<table>
<thead>
<tr>
<th>First author reference</th>
<th>Patients entered/ dropouts</th>
<th>Medication</th>
<th>Main outcome measures</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Johnson et al.</td>
<td>17/2</td>
<td>2 capsules (25 mg) powdered feverfew</td>
<td>Frequency of headache. Incidence of nausea and vomiting.</td>
<td>Frequency of headache increased significantly (p&lt;0.02) in patients receiving placebo compared to baseline values.</td>
</tr>
<tr>
<td>Murphy et al.</td>
<td>72/12</td>
<td>1 capsule (82 mg) powdered feverfew</td>
<td>Frequency, duration and severity of headache. Incidence of nausea and vomiting.</td>
<td>24% reduction (p&lt;0.005) in attack frequency, significant reduction (p&lt;0.002) of N/V. No change in duration and severity of headache.</td>
</tr>
<tr>
<td>Kuritzky et al.</td>
<td>20/nr</td>
<td>100 mg feverfew</td>
<td>Effect of feverfew on 5-HT uptake and platelet activity.</td>
<td>No effect</td>
</tr>
<tr>
<td>Weerdt et al.</td>
<td>50/6</td>
<td>1 capsule (143 mg) granulated feverfew</td>
<td>Severity of headache attacks. # work days lost.</td>
<td>No significant effect in either outcome.</td>
</tr>
<tr>
<td>Palevitch et al.</td>
<td>57/nr</td>
<td>2 capsule (50 mg) powdered feverfew</td>
<td>Pain intensity. Severity of nausea and vomiting, sensitivity to noise &amp; light.</td>
<td>Significant reduction (p&lt;0.01) in each outcome measure.</td>
</tr>
</tbody>
</table>

nr: not reported
MECHANISMS OF ACTION

Inhibition of platelet aggregation / Serotonin levels
Platelet aggregation and serotonin release have been shown to be present during migraine attacks. Magnesium has been shown to cause a dose-dependent inhibition of platelet aggregation. Decreased magnesium levels may contribute to thrombin-induced platelet aggregation, which can lead to serotonin release from platelets.16

Vascular dilating effect
Magnesium has a strong vascular dilating effect lending support to the vascular theory of migraine.2 Ionized magnesium levels are known to affect entry of calcium and release of intracellular calcium from the sacroplasmic and endoplasmic reticulum in vascular smooth muscle and vascular endothelial cells, and to control vascular tone and reactivity to endogenous hormones and neurotransmitters. Cerebral blood vessel muscle cells are particularly sensitive to ionized magnesium; magnesium deficiency results in contraction and potentiation of vasoconstrictors and excess magnesium results in vasodilation and inhibition of vasoconstrictors.16

NMDA receptor antagonist
Magnesium is intimately involved in the control of N-methyl-D-aspartate (NMDA) glutamate receptors, which play an important role in pain transmission in the nervous system and in the regulation of cerebral blood flow. Magnesium ions plug the NMDA receptors and prevent calcium ions from entering the cell. Lowering magnesium concentration facilitates activation of the NMDA receptor, which allows calcium to enter the cell and exert its effects on both neurons and cerebral vascular muscle.16 Blocking the receptor renders calcium unable to exert its vasodilatory effects.

Results of clinical studies
• Many clinical studies have researched the use of magnesium in migraine treatment and prophylaxis. The following data is from selected published clinical studies.
• Bigal et al. found that the relief of pain with IV magnesium sulfate (MgSO₄) was not different from treatment with placebo in the migraine without aura (MO) group, and was better than placebo in the migraine with aura group (MA). Magnesium sulfate was highly effective in relieving photophobia and phonophobia in both MO and MA groups. There was a greater response observed, in all symptoms, in the MA group than in the MO group.17
• In a study by Peikert et al., high oral doses of magnesium lowered the frequency of migraine attacks within 12 weeks of therapy. Compared to placebo, the therapeutic effects were already significant by the second therapy phase (weeks 5–8) and were confirmed by a significant reduction in the number of migraine days as well as the per patient consumption of acute medication. The duration and intensity of the attacks also declined without being statistically significant in comparison with the placebo group.18
• In a study by Mauskop et al., of the 40 patients to whom IV MgSO₄ was administered, 32 (80%) had at least a 50% initial reduction of pain intensity. In most patients, headaches began to improve before the end of infusion. Complete elimination of pain was observed in 80% of the 32 patients within 15 minutes of infusion. Of these 32 patients, 18 had persistent headache relief beyond 24 hours. Long-term responses to MgSO₄ varied in the different diagnostic categories.19
• Findings from a study by Mauskop et al. indicate that serum ionized magnesium levels can be used as a marker for detection of patients with migraine and cluster headaches who can benefit from magnesium infusions.20
• In a double-blind, placebo-controlled trial of oral magnesium supplementation in 24 women with menstrual migraine, positive results were noted. Taken at a dose of 360 mg/day taken in 3 divided doses for 4 months, there was a 50% reduction in the number of days with headache. Patients receiving active treatment also showed improvement according to the Menstrual Distress Questionnaire score. Four patients dropped out of the study, but only one did so because of adverse effects (magnesium-induced diarrhea). In a larger double-blind, placebo-controlled study involving 81 patients with migraine headaches, a significant improvement in patients receiving magnesium therapy was demonstrated. The frequency of migraine attacks was reduced by 41.6% in the magnesium group compared with only 15.8% in the placebo group; 3 patients receiving magnesium therapy dropped out of the study.21

Dosage
For migraine prophylaxis: oral intake of 300 to 600 mg/day.21,22

Adverse reactions
Diarrhea and gastric complaints are the most commonly reported adverse drug reactions. In one study, tolerability of magnesium was assessed. Sixteen (45.6%) of 35 patients in the magnesium group reported 35 adverse events during the course of treatment, mainly soft stools (5 patients), diarrhea (5 patients), and heart palpitations (3 patients). Altogether 17 episodes of adverse events occurred in 8 (23.5%) of the 34 patients on placebo.22
At higher doses of magnesium, low blood pressure, nausea, vomiting, urinary retention, decreased heart rate, and dilation of blood vessels have been documented. Coma and cardiac arrest are known to occur with toxic doses of magnesium. Magnesium may accumulate in patients who have decreased renal function; therefore, one must be cautious when consuming magnesium as a dietary supplement.
In addition, use of magnesium in pregnant women should be approached with caution due to its dilating properties.23

Drug Interactions

Before beginning therapy with magnesium or any over-the-counter supplement, a complete medication review should be conducted by a pharmacist or physician since magnesium may interfere with the absorption of many other medications. Lanoxin,® Macrobid,® and Vibramycin® may be less well absorbed in the presence of magnesium, decreasing their effectiveness. This is especially important with Lanoxin since its therapeutic concentration must be monitored closely for clinical effectiveness. Antimicrobials like nitrofurantoin and tetracyclines may not be effective against bacterial pathogens due to less-than-anticipated serum concentrations. Concomitant use with excretion-reducing drugs can increase the effects of supplemental magnesium and magnesium serum levels. These drugs include calcitonin, glucagon, and potassium-sparing diuretics.23

Family History

A strong family history is present in up to 80% of patients with migraines.21 This fact along with the identification of a gene for familial hemiplegic migraine suggests that genetic factors are present in a majority of migraine patients. Cellular magnesium content and magnesium metabolism are also under genetic control. Possibly there is an overlap between these two genetic mechanisms.21

Migraine Management

Another important aspect of migraine management is identifying and avoiding triggers. Many migraine headaches can be triggered by certain foods, especially those containing tyramine. Caffeine or caffeine withdrawal and strong odors, such as perfume, are other common triggers. It is often helpful for patients to keep a headache diary to record possible triggers. Typically, removing identified triggers significantly reduces the frequency of headaches.22

Riboflavin

Riboflavin, also known as the water-soluble vitamin B2, is essential for the body’s conversion of food to energy. It enables carbohydrates, proteins and fats to release energy, and is also needed for normal reproduction, growth and repair of skin, hair, nails, and joints. The riboflavin requirement in humans is often related to energy intake, but it appears more closely related to resting metabolic requirements. Differing amounts are recommended for infants, children, and pregnant women based on differences in their caloric intakes. The recommended daily intake for adults is 1.1 mg for women and 1/3 mg/day for men. Rich sources of riboflavin are liver, kidney, eggs, milk, cheese, yeast, broccoli and spinach.24

Indications: Riboflavin deficiency prophylaxis.24 Possibly effective in migraine prophylaxis.24-27

Chemistry

The active, phosphorylated forms of riboflavin, flavine mononucleotide and flavine adenine dinucleotide, are involved as coenzymes in oxidative-reductive metabolic reactions. These two coenzymes are necessary for normal tissue respiration. Riboflavin is also necessary for the functioning of pyridoxine and nicotinic acid.

Pharmacokinetics

Riboflavin is readily absorbed in the proximal small intestine by a saturable transport mechanism. It is enzymatically metabolized to two active metabolites in the small intestine: flavin mononucleotide and flavin adenine dinucleotide. Bile salts enhance absorption of riboflavin and people with biliary obstruction have decreased absorption. Approximately 6 to 12% of a dietary dose is excreted in the urine. The amount excreted in the feces can exceed the amount ingested following dietary doses due to the synthesis of riboflavin by intestinal bacteria. The elimination half-life is 1.4 hours with a terminal half-life of 14 hours.

Safety

Problems in humans have not been documented with normal oral intake of daily recommended amounts. Toxicity of high doses of riboflavin has not been reported.24 Likely safe when used at the recommended dietary allowance (RDA) of 1.4 mg per day. There is insufficient reliable information for using larger amounts during pregnancy.24

Drug Interactions

Concomitant use with Probenecid (Benemid) inhibits supplemental riboflavin absorption. Proantheline Bromide (Pro-Banthine) concomitant use delays and increases supplemental riboflavin.24

Adverse Effects

Bright yellow-orange discoloration of urine.24

Migraine Considerations

A mitochondrial dysfunction resulting in impaired oxygen metabolism may play roles in migraine pathogenesis. Migraine headache can be a prominent feature in patients affected by the syndrome of mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes (MELAS). Riboflavin is the precursor of flavin mononucleotide and flavin adenine dinucleotide, which are required for the activity of flavoenzymes involved in the electron transport chain. Riboflavin, when given to patients with MELAS or mitochondrial myopathies on the assumption that at large doses it might augment activity of mitochondrial complexes I and II, and improve oxidative metabolism, was able to improve clinical and biochemical abnormalities.28

Controlled Trials

Two controlled trials exhibited the benefits of riboflavin in migraine prophylaxis. A randomized placebo-controlled, double-blind trial was conducted by Schoenen et
al. to compare the effects of high-dose riboflavin versus placebo on migraine frequency, number of migraine days, duration, and severity of headache. Fifty-five migraine patients were randomized to receive either riboflavin, 400 mg a day, or placebo for a 3-month period. Using an intention-to-treat analysis, riboflavin was superior to placebo in reducing attack frequency (p=0.005) and headache days (p=0.012). In the group of patients who improved by at least 50%, i.e. responders, the attack frequency improved 19% in the placebo group and 56% in the riboflavin group (p=0.01); in the number of migraine days, the placebo group improved 15% as compared to a 59% improvement in the riboflavin group (p=0.002). The authors of the study concluded that because of its high efficacy, excellent tolerability, and low cost, riboflavin is an interesting option for migraine prophylaxis and a candidate for a comparative trial with an established prophylactic drug. Three minor adverse events occurred, two in the riboflavin group (diarrhea and polyuria), and one in the placebo group (abdominal cramps). No serious side effects were reported and the riboflavin was well tolerated.28

An open pilot study was conducted and 49 patients with recurrent migraines were given riboflavin 400 mg/day with breakfast for at least 3 months. The mean number of migraines fell by 67% and mean migraine severity improved by 68%. One patient stopped treatment because of gastric intolerance, but that patient was also taking aspirin. No other side effects were reported and the riboflavin was well tolerated. The study suggests that riboflavin supplements may reduce the recurrence rate of migraines.29

Conclusion

The controlled trials are encouraging information for further research. Riboflavin is a promising alternative for preventing migraine headaches because it is safe, well tolerated and inexpensive. However, more research needs to be conducted to prove the efficacy and long-term safety.

COMBINATION PRODUCTS: MAGNESIUM, FEVERFEW, AND RIBOFLAVIN

Since moderate success has been suggested with individual preparations of magnesium, feverfew, and riboflavin, use of these agents in combination may exert a synergistic antagonistic affect on migraine prophylaxis and treatment. Several combination dietary supplement products are currently available, including MigreLief® With Puracol,™ Herbal Migraine Formula,® and MigraHealth,™ 30

While individual data on these products is limited, information from physicians who are using these combination products in their practices suggest that combination products may assist to maintain proper tone of cerebral blood vessels. By improving tone, blood may flow freely from vessel to vessel and prevent sudden spasms. It may also inhibit platelet aggregation, stabilize membranes, and decrease the inflammatory process.31 Two case reports from a neurology practice that used a combination product of magnesium, feverfew, and riboflavin to treat migraine follow this review.

Randomized, placebo-controlled studies of the combination products are required to fully evaluate and confirm their benefit in migraine treatment or prophylaxis.

REFERENCES:


Case Reports on Patients Diagnosed with Migraine and Placed on A Nutraceutical Product Containing Riboflavin, Magnesium and Feverfew

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While serving as assistant professor of Neurology at the University of Miami School of Medicine, Dr. Cohen was in the forefront of the development of neurorehabilitation as a speciality. Her work as Director of Outpatient Neurorehabilitation Services at the University of Miami culminated in the book Comprehensive Management of Parkinson’s Disease, one of the first to outline a holistic approach to treating this condition. Dr. Cohen has practiced holistic neurology for the past ten years and maintains a holistic neurology practice in Denver. These case reports come from her practice following patient consent.

CASE HISTORY #1:
A 50-year-old, right-handed woman reported to the neurologist's office with a history of severe headaches dating back to high school. Over time, the headaches increased in frequency and intensity. She first sought medical assistance in her early 30s but only received Fiorinal to be taken abortively. By her early 40s, she suffered from severe headaches four to six times a month, each lasting 8 to 12 hours. Amitriptyline at 25 mg decreased the frequency of headaches to four times a month. Identified triggers at that time included MSG, sulfates, nitrates, dehydration, changes in weather, and flying. Verapamil was added to the regimen approximately five years later, but there was no clear change in the headache pattern.

At the time of her presentation, headaches were described as starting with a throbbing sensation in a headband distribution with pain intensity at 4 on a scale of 1 to 10. The pain then evolved to a severe left temporal throbbing considered "excruciating," sometimes associated with nausea. There were no visual changes. There was no numbness, tingling, or weakness. Abortive treatment with Maxalt or Imitrex nasal spray helped to decrease the pain, but two doses were required and the patient was left with a "significant hangover effect." Often headaches were so severe that they put the patient to bed.

A survey of pertinent lifestyle factors showed consistent sleep patterns with seven to eight hours of sleep a night without interruption. There was no clear source of toxin exposure in the work or home environment. Aerobic exercise was performed at least 20 minutes daily and she played tennis at least two times a week for two hours at a time.

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Given the prior identification of dietary triggers, diet consisted of unprocessed foods without soda, coffee, or chocolate. The patient described herself as "happy, anxious, and angry". Her most significant stressors were related to her family, particularly her teenage children. A high level of satisfaction was attained from her environment and financial status. Levels of fun and recreation, career, relationships, and level of spirituality engendered moderate feelings of satisfaction.

Medications were Verapamil 80 mg a day, Amitriptyline 25 mg qhs, either Rizatriptan 10 mg or Sumatriptan nasal spray 20 mg abortively for headache, and Temazepam 15 mg as needed for sleep with headache.

A complete neurology examination showed significant muscle spasm along the upper border of the trapezius muscles as well as in the occipital areas bilaterally, more so on the right side than the left. There were no abnormalities at the temporomandibular joint, including clicking or tenderness. Mental status, cranial nerve testing, and sensory examinations were within normal limits. Strength to confrontation was normal. There was persistent fixation about the left upper extremity but no drift or slowness of fine finger movements on that side. Reflexes were 2 throughout. The right toe was downgoing. The left toe was equivocal, a finding difficult to interpret given the history of prior surgery to the left toes.

Impression after the initial evaluation was migraine without aura possibly preceded by muscle contraction headache. Although the patient had identified triggers for her migraines and eliminated these from her lifestyle, she continued to have headaches at a significant rate of four to six per month. At the time of her visit, she was on subtherapeutic doses of Verapamil and Amitriptyline. She also waited to institute abortive treatment until the headache was well evolved. Findings on examination were discussed including the question of subtle weakness on the left side. Initial recommendations included an MRI of the brain as well as maintenance of a headache diary. The MRI showed a few punctate areas of increased signal bilaterally. Several changes were made to the medical regimen.

Verapamil was discontinued and Amitriptyline was slowly increased to 35 mg nightly. A combination product of magnesium, riboflavin, and feverfew (Migre Lief®) was initiated at one tablet twice a day. Recommendations were made to take abortive treatment at onset of migraine in order to optimize therapeutic benefit. Strategies for addressing muscle contraction and stress responses were also discussed.

Two and a half months later, there was a clear improvement in headache pattern. Headache frequency decreased from an average of five per month to approximately one per month. The patient did not have any headaches that forced her into bed. She noted improved response of the headaches to abortive management when taking the triptan at onset of migraine. She denied side effects of medication other than a dry mouth likely due to the Amitriptyline. Other than a transient increase in migraine frequency associated with significant stress, the patient's headache pattern has remained stable.

CASE HISTORY #2:

A 58-year-old, right-handed gentleman presented to the physician's office with an almost 20-year history of severe headaches. Headaches typically occurred two to three times a month, but in the winter occurred up to two times a week. Headaches started in a retro-orbital location as a moderate to severe throbbing pain either on the right or left sides. There was no aura. He experienced nausea but no photophobia. Headaches generally lasted 4 to 12 hours and were abortive successfully with Cafergot but usually required two doses. There were no clear triggers other than stress.

In looking at lifestyle factors, there was a high level of satisfaction with relationships and recreation with only moderate levels regarding career and financial issues. Stressors were most commonly related to career. Sleep was generally quite good. Dietary factors showed little intake of chocolate or caffeine. Alcohol intake was only occasional and there was no history of smoking. Aerobic exercise was performed four times weekly.

Medications were Propranolol at 80 mg a day and Niaspan at 1000 mg a day.

A neurology examination was pertinent only for a depression in the left skull secondary to placement of a radium plate to treat a scalp lesion many years before. Mental status, cranial nerve, motor, sensory, reflex, and coordination examinations were within normal limits.

Clinical impression was that of migraine without aura. In the past, there was no response to prophylactic treatment with Verapamil and Nifedipine. The patient had a partial response to Propranolol at 80 mg a day. In spite of this, he continued with three to four headaches per month and responses to Cafergot were diminishing. A recommendation was made to institute a combination product of magnesium, riboflavin, and feverfew (Migre Lief®) at one tablet twice a day. At a subsequent visit six months later, headache frequency had diminished to one headache every four to six weeks. Subsequent visits showed maintenance of this pattern over a year. The patient denied side effects from the medication.