N Alpha Methyl Histamine Versus Propranolol in Migraine Prophylaxis

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ABSTRACT: Objectives: To compare the efficacy and tolerability of the subcutaneous administration of N alpha methyl histamine versus oral propranolol in the treatment of migraine prophylaxis. Background: N alpha methyl histamine has a selective affinity for H3 receptors and could constitute a new therapeutic drug in migraine prophylaxis. Methods: Sixty patients with migraine were selected and enrolled in a 12-week double-blind controlled clinical trial to evaluate the efficacy of subcutaneous administration of N alpha methyl histamine (1 to 3 ug twice a week) n=30, compared to administration of 120 mg/day of oral propranolol n=30. The variables were: headache intensity, frequency of attacks, duration of migraine attacks and analgesic intake. Results: Fifty five patients completed the study. The data collected during the 4th week of treatment revealed that N alpha methyl histamine and propranolol caused a significantly (p<0.01) greater reduction between the basal values and final values of every variable studied. Conclusions: Both N alpha methyl histamine and propranolol are similarly effective in reducing or eliminating the headache in migraine prophylaxis. Low doses of N alpha methyl histamine injected subcutaneously may represent a novel and effective therapeutic alternative in migraine patients and may lay the clinical and pharmacological groundwork for the use of H3 receptor agonist in migraine prophylaxis.

RÉSUMÉ: N-alpha-méthyl-histamine comparée au propranolol dans la prophylaxie de la migraine. Objectifs : Le but de l’étude était de comparer l’efficacité et la tolérabilité de la N-alpha-méthyl-histamine par voie sous-cutanée à celle du propranolol par voie orale administrés en prophylaxie de la migraine. Contexte : La N-alpha-méthyl-histamine a une affinité sélective pour les récepteurs H3 et pourrait constituer un nouveau moyen de prévention de la migraine. Méthode : Soixante patients atteints de migraine ont été choisis et recrutés pour participer à un essai clinique contrôlé, à double insu, de 12 semaines afin d’évaluer l’efficacité de l’administration de N-alpha-méthyl-histamine par voie sous-cutanée (1 à 3 mg deux fois par semaine) n = 30, comparée à l’administration de 120 mg de propranolol par jour n = 30. Les variables étudiées étaient l’intensité de la céphalée, la fréquence des crises de migraine, leur durée et les analgésiques utilisés. Résultats : Cinquante-cinq patients ont complété l’étude. Les données recueillies au cours de la 4e semaine de traitement ont montré que la N-alpha-méthyl-histamine et le propranolol provoquaient une diminution significativement plus importante (p <0.01) par rapport aux valeurs de bases de toutes les variables étudiées. Conclusions : La N-alpha-méthyl-histamine et le propranolol sont aussi efficaces l’un que l’autre pour diminuer ou éliminer la céphalée en prophylaxie de la migraine. La N-alpha-méthyl-histamine par voie sous-cutanée à petites doses pourrait constituer un traitement alternatif nouveau et efficace chez les patients migraineux et établir les bases cliniques et pharmacologiques de l’utilisation d’un agoniste H3 en prophylaxie de la migraine.

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Migraine affects approximately 10% to 14.7% of the general population and 11% for lifetime migraine based on both the first edition of the International Classification of Headache Disorders (ICHD-I)1 and second edition of the International Classification of Headache Disorders (ICHD-II)2 definitions of migraine and it is two to three times more common in women than in men3,4. Many types of drugs are available for clinical use for the prevention of migraine, including β-blockers, antidepressants, antiepileptic agents, and calcium channel blockers, which have demonstrated efficacy in 70% of cases; nonetheless there is up to a 10% to 30% therapeutic failure rate, with persistent periodic migraine attacks, resulting in an elevated analgesic consumption and a significant deterioration in quality of life5-8. Currently, few of the drugs employed in migraine prophylaxis act on specific mechanisms of the migraine pathophysiology9. The cause and pathophysiology of migraine are not well understood. There seems to be a relationship between brain metabolism and cerebrovascular dysfunction, which activates pain pathways that cause a series of events ending in neurogenic inflammation10,11, an important component in migraine that is perhaps genetically determined. In 1991 we carried out the first study12 providing evidence of the beneficial effect of histamine in migraine prophylaxis and of the histamine metabolite, N alpha methyl histamine, which possesses a greater affinity for H3 receptors13,14. In our clinical open study, Phase II and Phase III, we demonstrated that N alpha methyl histamine, at a dose ranging from 1 to 3 ug, has therapeutic potential in migraine prophylaxis. It induced significant relief from migraine symptoms, with no secondary effects; interestingly15,16 N alpha histamine...
methyl histamine could constitute a new therapeutic drug in migraine prophylaxis, that improves quality of life for migraine patients who do not respond (30-40%) to the drugs used today; It could lay the clinical and pharmacological groundwork for the use of H3 agonists in migraine prophylaxis: and, finally; could help obtain a drug that acts upon specific mechanisms of pathophysiologic processes related to migraine. The aim of this study is to evaluate the therapeutic potential in migraine prophylaxis of subcutaneous administration of N alpha methyl histamine compared with oral administration of β-blockers (propranolol)\(^\text{17}\) in a clinical trial.

**METHODS**

Sixty patients with migraine with or without aura were selected using the criteria of the International Headache Society\(^\text{1,2}\) and enrolled in a 12-week double-blind controlled clinical trial to evaluate the efficacy of subcutaneous administration of N alpha methyl histamine (1 to 3 \text{ug} twice a week \(n=30\), compared to administration of 120 mg/day of oral propranolol \(n=30\). The selected patients hailed from multiple physicians and neurologists and their diagnoses were independently confirmed by a second member of the research team\(^\text{19}\). The trial procedure was explained to them and they were invited to take part in the study. All participants signed a letter of consent in accordance with the Helsinki statement and ethics committee of HGZ No 1 IMSS Colima, Colima, México. The patients were male or female adults between the ages of 18 and 65 years, all having a history of migraine for several years, with no additional neurological or cardiovascular pathologies; pregnant women, patients suffering from daily headaches and patients whose radiological tests, including computer-assisted tomography, revealed any pathology, were excluded from the study. Selected patients underwent a one-month period of prophylactic agent washout, during which headache frequency was monitored (Figure 1 flow diagram). They were then divided into two groups for treatment in randomized blocks of three\(^\text{19}\), double-blind fashion: the N alpha methyl histamine study group \((n=30)\) and the propranolol group \((n =30)\). This randomization was carried out by a research collaborator who throughout the duration of the study had no contact with the patients and prepared vials containing either 10 \text{ug/ml} subcutaneous N alpha methyl histamine or oral placebo; and 120 mg oral propranolol or subcutaneous placebo (Evan’s solution= phenol 0.4%, isotonic sodium chloride). The vials were numbered and were identical in appearance, which allowed the blinding to be effective since neither the patients nor the physicians were able to identify the vehicle or active drug. The treatment consisted of subcutaneous (posterior region of the upper arm) administration of N alpha methyl histamine (10 \text{ug/ml} in Evan’s solution) 1 to 3 \text{ug} twice a week. The regimen began with the administration of 0.1 \text{ml} volume of either subcutaneous N alpha methyl histamine or placebo, which was consecutively increased (by 0.1 \text{ml}) halting the increase at 0.3 \text{ml}. Continued repetition of this protocol (starting again with 0.1 \text{ml} volume administration) and 120 mg propranolol (40 mg/d x3) or subcutaneous placebo twice a week was carried out for a period of twelve weeks. During treatment, patients were allowed to take 500 mg acetaminophen tablets if they had moderate to severe headache with an intensity value of 2 on a scale of 1 to 3, and lasting for more than eight hours (hrs). The variables studied\(^\text{20}\) were: 1) headache frequency, measured by number of attacks per month; 2) intensity of pain (scale from 1 to 3); 3) duration of pain, measured by hours of headache per attack; 4) intake of rescue analgesics, measured by the number of acetaminophen tablets (500 mg) taken per month. Values for the parameters studied were collected over a period of four weeks before initiation of treatment (baseline), and efficacy and safety assessments were carried out every 30 days for a period of twelve weeks. Patients were instructed to keep a daily record of events. The relationship between an adverse event and the study treatment was assessed by the investigator as none, possible, probable, or definite. Patients who abandoned the study were still taken into account in the final analysis.

Statistical Analysis: Average descriptive statistics and standard deviations were applied to obtained data. The Wilcoxon rank-sum test was used to assess the statistical significance of differences between treatment groups in baseline characteristics (age, years since onset, age at onset), during the aforementioned weeks of treatment it was performed using a Mann-Whitney U rank sum test. In order to analyze the temporal course of each treatment (for each variable studied), a Friedman repeated measures analysis of variance (ANOVA) on ranks test was used to evaluate the statistical significance of differences between basal values and values found for the 4th and 12th weeks of treatment. With an alpha level of 0.05, the trial was designed to have a statistical power of 80 percent, \(p<0.05\) was considered significant\(^\text{21}\). Data statistical analysis was carried out using the Statistical Package for Social Sciences program (SPSS 10.0). The study was approved by the Ethical and Scientific Committee of our hospital and the Australian New Zealand Clinical Trials Registry (ANZCTR). ACTRN12610000126011.

**RESULTS**

From the total of 60 patients enrolled and randomized into the two groups (N alpha methyl histamine \(n=30\); propranolol \(n=30\), 27 in the N alpha methyl group and 28 in the propranolol group...
completed the study. A total of 90% of the patients were female (26 female; 4 male in the N alpha methyl histamine group and 28 female/ 2 male in the propranolol group). Mean age was of 35 years ± 11.7 (range 18 to 60). Headache duration was 12 ± 9.3 years and frequency of migraine attacks was 5.12 (95% IC 1.23 to 1.89) per month with or without aura. The treatment groups were similar at baseline based on demographic and clinical characteristics (Table) for all variables studied. The statistical analysis of data collected showed no significant differences between the baseline values obtained for the N alpha methyl histamine and the propranolol treatment groups (P>0.05).

We compared the response of one group to its baseline and the differential response of one group to the other. Analysis of the temporal course of events showed that by the beginning of

![Figure 2: Effects induced (in 60 patients with recurrent migraine) by the subcutaneous administration (twice a week, during 12 weeks) of N alpha methyl histamine (n=30) and oral propranolol (n=30) on the frequency. Data correspond to mean values (plus SEM) obtained during a 4-week period prior to initiation of treatment (Basal), and 12 weeks of treatment.](image)

![Figure 3: Effects induced (in 60 patients with recurrent migraine) by the subcutaneous administration (twice a week, during 12 weeks) of N alpha methyl histamine (n=30) and oral propranolol (n=30) on the intensity. Data correspond to mean values (plus SEM) obtained during a 4-week period prior to initiation of treatment (Basal), and 12 weeks of treatment.](image)

### Table: General and clinical characteristics of patients

<table>
<thead>
<tr>
<th>FEATURE</th>
<th>N-alpha methyl Histamine (n=30)</th>
<th>Propranolol (n=30)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age years (mean±)</td>
<td>35 (±15)</td>
<td>33 (±10)</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>4</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>26</td>
<td>28</td>
<td></td>
</tr>
<tr>
<td>Years of migraine (mean)</td>
<td>12(±9)</td>
<td>16 (± 10)</td>
<td></td>
</tr>
<tr>
<td>Migraine type</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>with aura</td>
<td>10</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>without aura</td>
<td>20</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>Age at onset (mean years)</td>
<td>22 (±9)</td>
<td>17 (± 8.8)</td>
<td></td>
</tr>
<tr>
<td>Frequency of headaches per 30-day periods at baseline. (mean score)</td>
<td>7.0 (±1.1)</td>
<td>6 (±1.1)</td>
<td>P&lt;0.86</td>
</tr>
<tr>
<td>Intensity of headache at baseline 1,2,3</td>
<td>2.9 (±2)</td>
<td>2.9 (±1.8)</td>
<td>p&lt;0.69</td>
</tr>
<tr>
<td>Duration of headache hours. (mean score)</td>
<td>46 (±12)</td>
<td>29 (±11)</td>
<td>P&lt;0.02</td>
</tr>
<tr>
<td>Tablets/mo of rescue. (mean score)</td>
<td>25(±9.4)</td>
<td>29 (±8)</td>
<td>P&lt;0.27</td>
</tr>
</tbody>
</table>

(by before undergoing period washout of prophylactic agents)
the 4th week, there was a significant decrease (with respect to basal values) in the magnitude of all parameters studied, as a result of the scheme followed for the administration of N alpha methyl histamine (P<0.001) or propranolol (P<0.001). The same was found for values at the 8th, and 12th weeks of treatment. (Figure 2-5)

The rate of response to N alpha methyl histamine (Figure 2) was 62% (17 of 27 patients) reported a 67% reduction in headache frequency (P<0.001) (mean, 7.0 95% CI 1.2 to 1.8 attacks per month before treatment vs 2.8 95% CI 1 to 1.2 attacks per month after treatment) number needed to treat (NNT) 4, and the rate of response to propranolol was 60 percent (17 of 28 patients), reported a 52% reduction (P< 0.001) (mean 6.0 95% CI 1.6 to 1.6 attacks per month before treatment vs 4.0 95% CI 1.6 to 1.1 attacks per month after treatment) NNT 4.

For pain intensity (Figure 3) 70% (19 of 27 patients) receiving N alpha methyl histamine reported a 47% reduction in headache intensity (P<0.001) (mean, 2.9 95% CI 1.2 to 2 before treatment vs 1.6 95% CI 1.5 to 2 after treatment) NNT 3; whereas 60 percent (17 of 28 patients) receiving propranolol reported a 52% reduction in intensity of pain (P<0.001) (mean 2.9 95% CI 1.8 to 5 before treatment vs 1.80 95% CI 1.4 to 3.4 after treatment) NNT 5.

Comparison between both groups (Figure 4) revealed a significant reduction in duration of migraine attacks: The rate of response to N alpha methyl histamine was 62% (17 of 27 patients), with a 67% reduction in duration of migraine attacks P< 0.001 (mean, 46 95% CI 12.5 to 6.4 hours of headache per attack before treatment vs 18.80 95% CI 9.6 to 9.2 hours of headache per attack after treatment) (NNT 4); and the rate of response to propranolol was 64% (18 of 28 patients), with a 71% reduction P<0.001 (mean 29 95% CI 11.1 to 6.5 hours of headache per attack before treatment vs 14.87 95% CI 8.2 to 9.4 hours of headache per attack after treatment (NNT 3.5).

In relation to rescue medication (Figure 5) 59% of patients receiving N alpha methyl histamine (16 of 27) reported a 47% reduction in the number of tablets ingested P<0.001 (mean, 25 95% CI 12.1 to 7.2 acetaminophen tablets per month before treatment vs 11.3 95% CI 3.2 to 10.1 acetaminophen tablets per month after treatment) (NNT5), whereas 64% of patients (18 of 28) receiving propranolol reported a 70% reduction (P< 0.001) (mean 29. 95% CI 7.5 to 4.5 acetaminophen tablets per month before treatment vs 15.47 95% CI 5.2 to 4 acetaminophen tablets per month after treatment (NNT 4).

Ten percent (3/30) of patients in the N alpha methyl histamine group withdrew without adverse events, because they were not satisfied with the speed of the results even though they did not present any side effects; some transitory burning and itching at the injection site was reported, but it was not significant enough to impede the blinding of the assay or the planned order of events. There were no modifications in arterial blood pressure or cardiac frequency in either group for the duration of the study, nor were there any alterations in the laboratory analyses performed at the beginning and end of the study. After 12 weeks of treatment, the effects of N alpha methyl histamine and propranolol remained identical to the values found at the 8th week.

**DISCUSSION**

It has been demonstrated that high concentrations of histamine activate H1 receptors, producing vasodilatation and release of nitric oxide which conduces to neurogenic edema and is responsible for the acute phase of migraine; however, the results obtained in this and in previous placebo-controlled studies show that histamine and the N alpha methyl histamine...
at low concentrations are safe drugs with therapeutic potential in migraine prophylaxis, exercising specific mechanisms on pathophysiologic processes involved in this disease. In migraine pathophysiology, the antidromic stimulation of trigeminal nerve endings induces the release of substance P and other neuromodulatory peptides, which in turn stimulate the release of histamine from mast cells. In meningeal blood vessels, activation of H1-receptors (H1-R) by high concentrations of histamine results in vasodilatation and plasma protein extravasation, causing vasogenic-neurogenic inflammation. Under conditions of homeostasis, low concentrations of histamine activate the H3 receptors of C-fiber nerve endings and mast cells and block the exit of neuropeptides, among them histamine itself, by negative feedback circuit between C-fiber nerve endings and mast cells, which control neurogenic inflammation. Histamine H3 receptors are autoreceptors that negatively regulate the release of histamine and other neurotransmitters and are believed to play a variety of physiological roles, including the regulation of feeding, arousal, cognition, pain, and endocrine systems.

Two histaminergic H3 receptors have been identified, which differ in their sensitivity to guanidino-nucleotide inhibition and are differentially activated by R alpha methyl histamine (Kd=12nM) and by N alpha methyl histamine (Kd=0.36nM); the first crosses the hematooencephalic barrier easily, while the second does not. The histamine metabolite N alpha methyl histamine possesses a greater affinity for H3 receptors. In our clinical study, Phase II and Phase III, we demonstrated that N alpha methyl histamine at doses from 1 to 3 ug, has therapeutic potential in migraine prophylaxis, inducing significant relief from migraine symptoms with no secondary effects; interestingly, we obtained an effective response at a low dose (1 to 3 ug), with a significant reduction in headache frequency, intensity, duration, and analgesic intake; however, when we employed doses greater than 4 ug, headache developed.

Our data reveal that the administration of N alpha methyl histamine, at low doses (1-3 ug), induces significant relief from migraine symptoms without complications. After the clinical trial was over, some of the patients treated with N alpha methyl histamine remained asymptomatic, without headache crises, while others showed significant relief from migraine symptoms for a period ranging from six to twelve months. A cross-over study was not carried out, due to the fact that the use of drugs having a prolonged therapeutic effect does not lend itself well to such a study. Altogether, the results obtained in this study show that N alpha methyl histamine is a safe drug with therapeutic potential in migraine prophylaxis.

This randomized study demonstrated that both N alpha methyl histamine and propranolol are similarly effective, and well tolerated, in reducing or eliminating the headache in migraine prophylaxis. Low doses of N alpha methyl histamine injected subcutaneously may represent a novel and effective therapeutic alternative in patients presenting with recurrent migraine who do not respond to beta adrenergic or calcium channel blockers or patients over 60 years-of-age who have hypotension or cardiac rhythm alterations and in whom the usual drugs are contraindicated, or in patients who have developed secondary gastritis and cannot tolerate further oral drug therapy. Twice-weekly, subcutaneous injection of histamine or N alpha methyl histamine in low concentrations has been accepted in our practice by patients who previously had not been satisfied with the daily administration of other medications, and may lay the clinical and pharmacological groundwork for the use of H3 agonist in migraine prophylaxis.

A better understanding of migraine pathophysiology along with the discovery of novel molecular targets has led to a growing number of upcoming therapeutic proposals. The possibility of better modulating the imbalance between central neurotransmitters that occurs with migraine has created an exciting search for new pharmacologic sites. Neuromodulators for the prevention of multiple mechanisms related to migraine are already available.

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REFERENCES


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