**AURASTOP® in the Treatment of Migraine Aura**

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**Introduction**

The phenomenon of aura is traditionally considered as a non-avoidable event occurring in about one quarter of migraine sufferers before the headache phase[1]. Although previous trials on migraine treatment have clearly demonstrated the benefit conferred by some agents, including triptans, on migraine headache, as well as their time-dependent effect, the therapeutic potential of modulating the phenomenon of migraine aura before pain occurrence has not been investigated so far. At least theoretically, interventions aimed at limiting migraine aura might also have an influence on pain control. Aura symptoms frequently persist beyond the 1-hour limit set by the International Headache Classification diagnostic criteria[2], with many accompanying manifestations (prostration, confusion, difficulty concentrating among the others) lasting up to 24 hours and whose complete resolution represents the end of the crisis. During this phase the patient experiences a high degree of disability by being forced to interrupt many activities of daily life as well as of psychological stress, though the symptoms of aura tend to recur with the same features over time.

To date, sparse studies have attempted to investigate the efficacy of selected molecules on the phenomenon of aura. Among these, the intranasal administration of ketamine proved effective in reducing the severity but not the duration of the aura[3]. Similarly, tonabersat as well as the intra-venous administration of prochlorperazine were tested in small studies or in anecdotal reports, but they did not receive approval for clinical use[4,5].

Over the last years, numerous nutraceutical compounds have been proposed for the preventive treatment of migraine with or without aura, leading many scientific societies to report ad hoc guidelines to support practitioners in their use[6]. In particular, the feverfew (Tanacetum Partenium) has been used for many years in the UK[7], and 5-hydroxy tryptophan for the prevention of migraine in Italy.

Based on what above, in the present study we tested the hypothesis that an association of molecules that has been proven to drastically reduce the duration and degree of disability of the aura symptoms can also influence the severity of pain that follows the aura itself. In particular, we investigated the therapeutic potentials of a new compound (Aurastop®), derived from the combination of tanacetum partenium (150 mg extracted at 0.8 % = 1,2 mg di of active partenolide), griffonia simplicifolia (20 mg of 5-HTP) and magnesium (185 mg of magnesium pidolatum).

**Keywords: Migraine aura; Tanacetum Partenium; Headache; Treatment**

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Patients and Methods

The present study is supported by the Società Italiana per lo Studio delle Cefalee (SISC), Lombardia section.

Participants

Patients with personal history of headache fulfilling ICHD-3 beta criteria for the diagnosis of migraine with aura were recruited at the Headache Centre of each participating hospital. Diagnosis of headache was made by experienced headache specialists, and each patient underwent a detailed clinical and neurological examination.

The following inclusion criteria were considered: 1) age between 18 and 60 years
2) diagnosis of migraine with aura, characterized by at least 3 episodes of aura/year with a minimum aura duration of 20 minutes.

The introduction of a preventive treatment during the observation period was considered an exclusion criterion. This study was an audit of outcome and, as such, according to the Italian guidelines, did not require ethics committee approval.

All participants received the combination of tanacetum parteninum (150 mg extracted at 0.8% = 1.2 mg of active parthenolide), griffonia simplicifoila (20 mg of 5-HTP) and magnesium (185 mg of magnesio pidolate) (Aurastop®).

Study design

At baseline (t0), the natural history of aura phenomenology was studied. To this purpose, each patient received a migraine headache diary, to keep track of aura and headache characteristics of the following 3 episodes. In particular, aura subtype (only visual, visual and somatosensory, visual, somatosensory and speech/language symptoms [here defined as complex]) aura duration, disability (on a scale ranging from 0 to 5), presence of concomitant/following headache characteristics (duration, intensity [Visual Analogic Scale, 0 to 10]), use of common pain medications (triptans, nonsteroidal anti-inflammatory drugs), and response to symptomatic treatment were considered.

After 3 episodes of aura (with or without migraine) migraine headache diary of each patient were re-evaluated (t1) considering inclusion/exclusion criteria and aura characteristics. Then, each patient received a blister with 6 tablets of Aurastop®, with the instruction to assume a tablet of Aurastop® at the beginning of the following 3 auras, recording aura characteristics on migraine headache diary as usual and a second tablet at the beginning of the pain (in case pain occurred). Patients were allowed to take the usual pain killer after 1 hour in case of persistent pain. Each patient and migraine headache diary data were further evaluated (t2) after these 3 aura episodes.

The primary end-point of the study was a reduction >50% of duration and disability of the aura phenomena. The secondary end-point was the modification of the headache features after the aura (in particular, duration, intensity, assumption of usual analgesics-triptans, and efficacy of the pain killer [on a 0 to 5 scale, with 0 indicating no pain and 5 indicating maximum pain severity]).

Statistical Analysis

Dependent variables were compared by McNemar’s χ2 analysis. Wilcoxon’s signed rank test was used to compare migraine aura and headache characteristics before and after treatment. Statistical analyses were performed using SPSS 21.0 (IBM SPSS Statistics 2013, Armonk, NY, USA).

Results

Two-hundred subjects with a diagnosis of migraine with aura (ICHD-3 beta criteria) qualified for the present study (mean age 33 ± 1.5 years [range, 18 – 54], males, 83 [33.2%]). As summarized in Table 1, Aurastop® determined a reduction of the duration of aura (t1= 34.2 ± 19.3 minutes vs t2 = 18.2 ± 10.3 minutes, p < 0.01), as well as of the degree of disability (t1 = 5 [4-5] vs t2 = 2 [1-2], p < 0.01). In particular, a 4-5 degree of disability was observed in more than 90% of patients before treatment versus a 1-2 degree in more than 90% of patients after treatment. Furthermore, the characteristics of migraine aura also changed after treatment with Aurastop®, with an obvious reduction of its complexity (p > 0.01), as well as of somatosensory manifestations (111 18.5% before treatment vs 20 3.3% after treatment). We also detected a significant reduction of headache crises (p < 0.01), of pain severity and duration (p < 0.01), of the number of analgesics presumed by each patient, and an increased level of efficacy determined by the same analgesics or triptans after treatment with Aurastop®.

### Table 1: Primary and secondary endpoints

<table>
<thead>
<tr>
<th>Migraine characteristics</th>
<th>Pre-treatment</th>
<th>Post-treatment</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Migraine aura</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Visual, n (%)</td>
<td>600 (100)</td>
<td>594 (99.0)</td>
<td>≤ 0.001</td>
</tr>
<tr>
<td>Sensory, n (%)</td>
<td>111 (18.5)</td>
<td>20 (3.3)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Complex, n (%)</td>
<td>29 (4.8)</td>
<td>1 (0.2)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Duration, minutes, mean ± SD</td>
<td>43.2 ± 19.3</td>
<td>18.2 ± 10.3</td>
<td>≤ 0.001</td>
</tr>
<tr>
<td>Disability (0-5),median (IQR)</td>
<td>5 (4-5)</td>
<td>2 (1-2)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Headache, n (%)</td>
<td>577 (96.2)</td>
<td>451 (76.7)</td>
<td>≤ 0.001</td>
</tr>
<tr>
<td>Duration, hours, mean ± SD</td>
<td>48.0 ± 13.3</td>
<td>3.8 ± 3.7</td>
<td>≤ 0.001</td>
</tr>
<tr>
<td>Pain intensity (O-10)</td>
<td>5 (4-7)</td>
<td>2 (1-2)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Benefit (0-5)</td>
<td>2 (1-2)</td>
<td>5 (4-5)</td>
<td>≤ 0.001</td>
</tr>
<tr>
<td>Use of analgesics, n (%)</td>
<td>576 (96.0)</td>
<td>372 (79.8)</td>
<td>≤ 0.001</td>
</tr>
</tbody>
</table>

**Figure 1**: Histogram of aura duration pre- and post-treatment with Aurastop®.
Discussion

The main finding of the present study is the observation of a ~96% reduction of self-reported aura episodes in patients who shifted from a standard treatment approach to the regimen including the new compound Aurastop®. Such a reduction concerned not only the duration of aura, but also the degree of disability determined by this phenomenon, which provides support to the hypothesis that Aurastop® may modulate cortical neuronal hyperexcitability and, consequently, modify the clinical features of aura itself. Furthermore, the observation that aura was no more followed by a headache phase in about 30% of patients, that pain severity and duration were significantly reduced, and that the efficacy of analgesics/triptans was increased in patients assuming Aurastop®, should be regarded as indirect demonstration that the new compound may interfere with the peripheral-central sensitization and with the TRPA1- and NMDA-dependent synaptic transmission as well.

A number of biologic effects may explain the clinical benefit conferred by Aurastop® in the treatment of migraine aura. First, Mg²⁺, a principal component of the new agent, has a well-known inhibitory effect on NMDA receptor activity which is likely to contrast the CSD phenomenon⁶⁹. Second, 5-HTP influences the kynurenine pathway, leading to an increase of kyna levels which, in turn, further inhibits the activity of NMDA receptors and the activation of the trigemino-vascular system, and also rapidly (kyna passes the blood-brain barrier [BBB]) interferes with CSD and BBB integrity[10-12]. Finally, partnolide, the active metabolite of tanacetum partenium, is an active agonist of TRPA1 receptors implicated in the release of CGRP from the perivascular terminals of neurons involved in neurogenic inflammation, the key mechanism leading to head pain[13]. The short delay between oral assumption of Aurastop® and the clinical effect on aura symptoms is well explained by the pharmacokinetic of tanacetum partenium, which, according to several results obtained in animal models[13], is characterized by a very short time for elution (1.3 minutes), absorption (~3 minutes), and passage through the BBB[14].

The results of this new analysis are well in line with those previously derived by the same Authors from a smaller independent cohort[15] and emphasize the therapeutic potentials of Aurastop®, though they were obtained in the setting of an observational study without a placebo group. Many molecules, however, have been widely used in medicine for years based only on clinical evidence of efficiency (i.e, beta-blockers for migraine prevention long before the pathogenesis of migraine was elucidated). On the other side, data from animal models provide strong support to the biologic effect of Aurastop®, in particular of one of its components, the 5-HTP and its metabolite Kynurenine acid, in modulating the cortical electric phenomenon underlying migraine aura[11].

Conclusion

In conclusion, the results of the present study strengthen the sparse anecdotal observations that Aurastop® may influence the early phases of migraine aura and cortical spreading depolarization resulting in pain and the accompanying symptoms. From a biological perspective, they might be regarded as a step forward in better understanding the basic mechanisms of the cortical hyperexcitability leading to headache.

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