The presence of patent foramen ovale (PFO) may allow venous blood to pass from the right atrium toward arterial blood in the left atrium through the interatrial septum. Although the incidence of any degree of PFO is high among the general population, the analysis of this trait in subjects with neurological diseases has raised interest in recent years [1]. The prevalence of migraine in Western countries is rather high, with some of these patients suffering migraine with aura (MA). The focal phenomenon that appears in MA is related to a phenomenon of propagated cortical depression and may be triggered by various stimuli, although the pathophysiology of aura and the mechanism by which the stimuli trigger migraine attacks are still unclear [2].

One of the neurological disorders in which an elevated prevalence of PFO has been reported is MA. Some studies have described an incidence of PFO in MA which is as high as among patients with stroke of undetermined origin [3–5]. Nevertheless, it is unknown if this association is incidental or if there is some connection or relationship. The interest in clarifying these relationships is further raised, as invasive techniques such as percutaneous closure of PFO is questioned as a feasible therapy in these cases [6].
Theoretically, some migraine triggers such as exercise or exertion can cause Valsalva manoeuvres, thus increasing the extent of right-to-left shunt (RLS) linked to PFO. If this shunt were indeed related to the genesis of migraine attacks, patients with MA and PFO might be able to recognise these activities as triggers of their migraines. The purpose of our study was to analyse a potential relationship between these triggers, the presence of RLS and MA attacks.

Patients and methods

The study population consisted of patients diagnosed with MA selected from specific outpatient visits for headaches, the first patients being recruited in late 2002. All patients met IHS criteria for migraine with typical aura [7]. Patients with a single episode of aura were expressly excluded, even when they had a history of migraine without aura. Less common forms of migraine or probable MA (acute onset or prolonged aura) were also excluded. As part of routine clinical practice, the patients were surveyed to determine medical history and triggers of migraine attacks. In particular, triggers included those commonly accepted, such as stress, sleep, changing weather, flashes of light, specific foods and hormonal factors, as well as circumstances that induce Valsalva manoeuvres such as strenuous exercise, exertion during bowel movements, weightlifting, sudden standing, coughing or laughing crises, and sexual activity. Informed consent was obtained from the patients as required by the local ethics committee to perform a transcranial Doppler.

The transcranial Doppler was used to study the vessels of Willis’ polygon through the temporal window and specifically determining the presence/absence of RLS as described in the method already published and used by other authors [8]. Briefly, agitated saline achieving a microbubble emulsion was injected through an antecubital venous access. One of the middle cerebral arteries was then simultaneously monitored by transcranial Doppler and the presence of RLS was determined by the appearance of transient high-intensity signals within the first 10 cardiac cycles after injection of saline solution. These measurements were repeated at baseline and after a controlled Valsalva manoeuvre. The examination allowed confirmation of the presence of RLS, as well as semiquantitative classification of the extent of the RLS. Patients with patterns of >25 or uncountable signals after the Valsalva’s manoeuvre (“shower” or “curtain” patterns) were defined as massive shunt [9].

Univariate analysis was used to determine the correlation between the presence of RLS and MA. In addition, univariate analysis was also employed in an attempt to identify the possible relationship between triggers (classic and those specifically investigated that produce Valsalva’s manoeuvre) and the presence or absence of RLS and the degree of shunt. We established a level of alpha error at 0.05 for single comparisons and for the subgroups analysis alpha was considered both at customary 0.05 and corrected for multiple comparisons at 0.025.

The same measurements were performed on a control series of patients with migraine without aura.

Results

We included a total of 72 patients diagnosed with MA and 31 with migraine without aura. A total of 103 patients were included with mean age 36±11.3 years; 36 men and 67 women. Mean age was 36±12 years (46 women, 26 men) among patients with MA and 37±9 years (21 women, 10 men) in those with migraine without aura. No differences were found in age or distribution by sex between patients with or without aura. The most relevant data on medical history are shown in Table 1.

In the survey on crisis triggers in patients with MA, stress was mentioned most often by 50 patients (69.4%), followed by changes in sleep patterns (n=37, 51.4%) and changing weather conditions (n=31, 43.1%). Strenuous physical exercise was the most common situation leading to Valsalva recognised as trigger, in 18 cases (25%). A total of 33 patients with MA (45.8%) reported at least one of these Valsalva-provoking activities as the trigger for their crisis.

For the migraine without aura controls, the trigger reported most often was stress by 29 patients (93.5%) followed by changing weather and variations in sleep patterns (n=18, 58.1%). Manoeuvres leading to Valsalva were reported by 12 patients (38.7%). There were no differences with respect to MA patients (Table 2).

RLS at baseline was found in 39 patients with MA, 54.1% of those analysed. After Valsalva’s manoeuvre, 44 (61.1%) patients showed positive RLS. In comparison with patients with migraine without aura, RLS was always more

Table 1 Medical history

|                  | Heart diseases | Respiratory diseases | Hypertension | Diabetes | DVT* | Smoker | Contraceptives |
|------------------|----------------|----------------------|--------------|----------|------|--------|----------------|
| MA†              | 3 (4.2%)       | 2 (2.8%)             | 10 (13.9%)   | 0 (0)    | 2 (2.8%) | 24 (33.3%) | 15 (20.8%)     |
| MwA‡             | 0 (0)          | 2 (6.5%)             | 4 (12.9%)    | 0 (0)    | 0 (0) | 6 (19.4%) | 6 (19.4%)      |

*Deep vein thrombosis
†Migraine with aura
‡Migraine without aura
common in patients with MA, whether at baseline (39 vs. 6, \(p<0.001\)) or after Valsalva’s manoeuvre (44 vs. 7, \(p<0.001\)). The semiquantitative assessment was 38.9% massive RLS in the MA group vs. 6.5% in the group without aura.

A meticulous analysis of patients who recognised situations potentially leading to a Valsalva manoeuvre as triggers indicated the following:

a) Among MA patients: 24 patients recognising these triggers showed positive RLS and 9 patients with these triggers showed negative RLS. Despite a trend, this difference was not statistically significant (\(p=0.063\)).

b) Among MA patients and considering the presence of massive shunt: patients reporting these triggers were more likely to have massive RLS (\(n=17, 51.5\%\)). In contrast, 28 patients (71%) did not recognise these triggers and did not have massive shunt vs. 11 patients (28.2%) without triggers and showing massive shunt (OR 2.7, 95% CI [1.02–7.17], \(p=0.043\)).

c) When considering the entire sample, including patients with and without aura, these differences also remained. Therefore, among those reporting these triggers massive shunt was found in 18 cases (40%) vs. 27 with lack of massive shunt (60%). Among patients without these triggers there was a lack of massive RLS in 46 cases (79.3%) vs. 12 cases (12.7%) lacking these triggers and with massive RLS, for a prevalence of massive shunt of 29.1% in all 103 patients (OR 2.5, 95% CI [1.01–6.11], \(p=0.032\)).

### Discussion

The results of this study provide some support for a potential correlation between triggers, the presence of RLS and migraine attacks. According to our results, patients with migraine who have RLS tend to recognise activities that can cause Valsalva’s manoeuvre, increasing the extent of shunt as a trigger of their migraine attacks.

PFO is a vestige of foetal circulation that results from a failure of the septum primum and secundum to fuse and is present in a high percentage of the unselected adult population (27%) [10]. Since the initial comments of Wilmshurst et al. [11, 12] on the elevated frequency of PFO among scuba divers with decompression illness and the improvement in migraine symptoms in those who had undergone closure of the foramen ovale, several series and studies on PFO and its relationship with neurological diseases have been published [1].

Perhaps the most relevant aspect has been the elevated frequency of PFO in stroke of undetermined cause [4, 13–15] and its recurrences [16, 17]. In fact, an association between stroke, PFO and migraine has been reported [5, 18, 19]. Along with stroke, however, migraine and particularly MA have been the focus of greater interest in studies on PFO.

The arguments put forth to support a connection between the presence of PFO and the genesis of MA are essentially the frequency of the association and the effects of foramen closure on headaches. The elevated frequency of PFO among patients with MA has been reported in various studies in recent years [4, 20] and interatrial septal aneurysm has also been reported more often [21]. In our study, RLS was found in 61% of the MA patients and 23% of the migraine without aura patients, figures comparable to those published in the literature. It is certainly striking in our study that many patients had RLS classified as massive, i.e., with “shower” or “curtain” patterns linked to significantly larger PFO. This finding has also been reported with very similar figures in both transoesophageal echocardiography studies [22], as well as transcranial Doppler studies [3]. Nevertheless, the numeric association does not imply causality.

According to most authors, the other basic reason for defending the causal role of PFO in migraine is the improvement in migraine after foramen ovale closure. This improvement resulted in a decrease of 14%–80% in the frequency of migraine attacks, depending on the series, a decrease in the severity measured by disability scales and even disappearance of the migraine attacks and/or aura [23–27]. Our study cannot contribute anything of interest in this regard as it is not an interventional study.

Many authors suggest that the association between PFO and migraine is incidental and linked to hereditary factors, with no direct link. A familial aggregation of PFO among the relatives of patients with cryptogenic stroke has been reported [28]. Several studies report a tendency

### Table 2 Migraine triggers

| Stress | Weather | Sleep | Foods | Valsalva | Exercise | >2 VPA*
|--------|---------|-------|-------|----------|----------|--------|
| MA†    | 50 (69.4%) | 31 (43.1%) | 37 (51.4%) | 19 (26.4%) | 33 (45.8%) | 18 (25%) | 10 (13.9%) |
| MwA‡   | 29 (93.5%) | 18 (58.1%) | 18 (58.1%) | 9 (29%) | 12 (38.7%) | 6 (19.4%) | 4 (12.9%) |

†Migraine with aura
‡Migraine without aura
*More than 2 Valsalva provoking activities
to inherit both migraines and an interatrial septum defect, and in fact the presence of septal defects appears to be transmitted in a dominant autonomic manner and significantly associated with MA [29], without this MA–PFO association being significantly linked to the female sex [30]. These findings sustain, in part, the hypothesis of a genetically determined association which would include a subgroup of MA associated to PFO, although no author has demonstrated a mechanism that could relate them.

Elegant hypotheses suggest that the presence of PFO may result in a shunt that sidesteps the pulmonary capillary filter, thus allowing some component capable of initiating the migraine attack to reach the central nervous system. Interestingly, a high prevalence of migraine has also been reported in patients with extracardiac shunts [31]. Wilmshurst and Nightingale recently provided a detailed description of this scenario [32]. The migraine triggers could be microembolisations from the venous territory [12] or, more likely, some activated molecule or particle that may elude the filtering or deactivation ability of the lung parenchyma. This role of the lungs in migraine as a filter is not new in the literature [33]. The candidate substance according to these authors would be the amines, in particular 5-hydroxytryptamine (5-HT), which is implicated in the genesis of migraine [34] and probably a platelet hyperactivation state already shown in the pathogenesis of the migraine [35]. To complete this hypothesis the capacity of the pulmonary vessels to inactivate 5-HT has also reported [36]. The series that report improvements in migraines with antiplatelet agents support this theory in part [37]. Whether or not the bypass resulting from PFO triggers the migraine when a specific threshold is exceeded or induces through chronic shunt a state of central hyperexcitability affected by other triggers is a matter of speculation. According to this scenario, patients with PFO and migraine may recognise that daily life situations that induce Valsalva manoeuvres do trigger crises by increasing the extent of RLS. This hypothesis has already been mentioned in the theoretical literature, although the question remains unanswered to our knowledge [38]. Our results would support the role of RLS as a trigger in at least some patients, due to the mechanisms mentioned. In addition, this statement would be true for both MA (where the association is particularly prevalent for reasons not yet understood) patients and for common migraine patients who have significant RLS. Nevertheless, the results only indicate a trend and the absence of a correlation would not invalidate the initial hypothesis, as it is known that patients with PFO experience shunt without pulmonary hypertension, both at rest and during activities as common as coughing [39–41]. We should again mention the high number of patients in our study who presented significant shunt at rest. In contrast to this theory, the preliminary results of the first randomised trial on PFO closure in migraine (MIST trial) did not achieve its primary endpoint for the complete elimination of headache [42].

As in similar case-control studies, a selection bias cannot be excluded and some patients with MA in particular, when triggers exist, are more likely to show up in headache clinics, although triggers were not the reason for consultation in this particular series. During data collection patients were included consecutively without considering any characteristics, except for MA according to the current diagnostic criteria. The control subjects were selected to match such characteristics as mean age and sex, and there were no significant differences in medical history between the two groups. However, in order to minimise type two error and confounding results, the control group should be increased and should be at least as large as the MA group. Another systematic error that cannot be ruled out in this design is recall bias when reporting triggers, although triggers were routinely collected in the medical interview prior to ultrasonographic study and determination of RLS, in order to minimise this bias. Finally, the type one level of error could be considered at the routine 0.05 value or could be corrected for subgroup analysis at 0.025 (Bonferroni method for multiple comparisons) in order to reduce spurious correlations. In our opinion this correction is too conservative when testing hypotheses using variables that are mutually correlated. In this case-control, exploratory study this correction (with correlated variables) might have increased the chance that true associations would not be discovered.

In conclusion, our study adds some information that supports the PFO–migraine connection, but this association remains, on the whole, unexplained. We consider it to be of maximum interest to answer a variety of questions: Is it an aetiological or accidental relation? What is the exact mechanism? and essentially, Could any new therapeutic procedure be justified?

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