Abstract

Background: The migraine attack is classically divided into the prodromal, aura, headache and postdromal phase. Previous studies have highlighted non-headache symptoms associated with migraine occurring during the prodromal or postdromal phase. This study aimed to track the evolution of non-headache symptoms throughout all phases of the migraine attack. We also wished to delineate the phenotype of patients with more symptomatic migraine episodes and explore the association between non-painful symptoms and migraine disease activity and patients’ disability.

Methods: Two-hundred and twenty-five migraine patients were enrolled and were asked to recall retrospectively whether non-headache symptoms occurred during the prodromal, headache and postdromal phase of their attacks. The occurrence of symptoms during the different migraine phases was tested using the Cochran’s Q tests, Cohen’s and Fleiss’ kappa. Differences between groups according to the presence of non-headache symptoms through the entire migraine attack and correlations between the frequency of non-headache symptoms experienced during all phases and patients’ disease activity and disability were also assessed.

Results: Ninety-nine percent of patients reported having at least one non-headache symptom in one phase of the migraine attack and 54% of patients had at least one non-headache symptom occurring during all phases of migraine. The occurrence of non-headache symptoms was different throughout the three phases of migraine, being higher during the headache phase than during the prodromal and postdromal phases. Symptoms with the highest co-occurrence throughout all migraine phases were neck stiffness, thirst and abdominal pain. Patients who experienced non-headache symptoms during all three phases of migraine were more frequently females, had a higher disability, were suffering from chronic migraine and had more frequently medication overuse headache.

Conclusion: Migraine is a complex neurological disorder with a wide constellation of non-headache symptoms that can affect the burden of the disease. A better characterization of the evolution of non-headache symptoms through the different phases of migraine can enrich our knowledge on migraine pathophysiology and improve the management of the disease.

Keywords: Migraine, Non-headache symptoms, Disability, Clinical phenotype
of non-headache symptoms starting before pain onset or persisting after headache resolution [3]. The migraine attack has been classically divided into the prodromal, aura, headache and postdromal phases. Head pain, by its presence or absence, marks the passage from one phase to another one [4]. Before the headache onset, numerous patients may experience premonitory symptoms, including mood changes, irritability, difficulty in concentration, food cravings, repetitive yawning, increased sensitivity to external stimuli, cranial autonomic symptoms and neck stiffness [5, 6]. During the headache phase, the migraine pain can be accompanied by nausea, vomiting, phonophobia and photophobia [2]. A variety of symptoms, including fatigue, sensory hypersensitivity, concentration difficulties, mood changes and neck stiffness, may persist for up to 24 h after headache pain resolution, thus characterizing the postdromal phase [7, 8]. Although migraine phases are usually described as consecutive stages, some non-headache symptoms may start during the earliest phase of the migraine attack and continue until the postdromal phase [9]. Previous studies investigated the prevalence and phenotype of non-headache symptoms occurring during the prodromal or postdromal phase [5, 6, 8, 10]. Only a few studies [5, 10] have explored the occurrence of the same non-headache symptom throughout all phases of the migraine attack. Recent magnetic resonance imaging studies showed that activation of brainstem, diencephalic and cortical areas could mediate non-painful symptoms associated with migraine headache [11–14]. Understanding the evolution of non-headache symptoms through the different phases of migraine could enrich our knowledge on migraine pathophysiology and improve the management of the disease.

The aim of this study was to characterize non-headache symptoms in a large sample of migraine patients and track the evolution of such symptomatology through the different phases of the migraine attack. We also sought to delineate the phenotype of patients experiencing non-headache symptoms during all phases of migraine and explore the association between non-painful symptoms and migraine disease activity and patients’ disability.

**Methods**

We enrolled 225 patients with migraine who have attended the Headache Clinic at San Raffaele Hospital for their first clinic appointment from February 2019 to July 2021. Inclusion criteria were: age $\geq$ 18 years; a diagnosis of migraine based on the International Classification of Headache Disorder criteria (third edition) [15]; evidence of a personally signed and dated informed consent document indicating that the subject had been informed of all pertinent aspects of the study. Exclusion criteria of the study included other primary headache disorders, other chronic pain conditions and the inability to provide a detailed clinical history. Patients taking migraine preventive treatments, with both episodic and chronic migraine, with a history of medication overuse and with comorbid depression or anxiety were included in the study. All patients underwent a neurological examination and the following clinical features of migraine were collected: age of migraine onset, frequency of monthly headache days (MHD), frequency of monthly migraine days (MMD), ongoing preventive treatments, migraine prevention used in the past and abortive treatments currently used to stop migraine attacks. The severity of attacks was determined as regards the severity of pain according the Numerical Rating Scale (NRS) [16] and presence of cutaneous allodynia during the migraine attack, quantified using the Allodynia Symptom Checklist (ASC-12) [17]. Patients’ disability was also quantified using the Migraine Disability Assessment Test (MIDAS) [18] and Headache Impact Test (HIT-6) [19]. During a semi-structured clinical interview, non-headache symptoms were orally listed and patients were asked to recall retrospectively whether each symptom usually occurred during their migraine attacks and whether these symptoms occurred during the prodromal, headache and postdromal phase of migraine. It was specified to patients to report the occurrence of non-headache symptoms taking into account those periods when they were not taking any migraine prevention and regardless the use of acute treatments. Non-headache symptoms investigated in migraine patients during the clinical interview are reported in Table 1.

**Table 1** List of non-headache symptoms investigated in migraine patients

| Category                      | Symptoms                     |
|-------------------------------|------------------------------|
| Fatigue/Cognitive changes     | Irritability                 |
|                               | Difficulty in concentration  |
|                               | Fatigue                      |
|                               | Lethargy                     |
|                               | Mood disturbance             |
|                               | Speech difficulty            |
| Homeostatic alterations       | Yawning                      |
|                               | Food craving                 |
|                               | Food aversion                |
|                               | Thirst                       |
| Sensory sensitivities         | Frequent urination           |
|                               | Thirst                       |
| Gastrointestinal symptoms     | Nausea                       |
|                               | Abdominal pain               |

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Standard protocol approvals
The study was approved by the local ethical committee and written informed consent was obtained from all participants.

Statistical analysis
Based on previous studies [20, 21], we estimated that a sample of at least 100 patients would allow solid results. The Cochran’s Q test was performed to assess any significant differences in the frequency of non-headache symptoms across the three phases of the migraine attack. The co-occurrence of symptoms during the different phases of migraine was tested using Cohen’s and Fleiss’ kappa coefficients with 95% confidence intervals. For the Cohen’s and Fleiss’ kappa coefficients we considered significant a kappa value > 0.3 (fair-good agreement), with a \( p < 0.05 \). Patients were divided into two subgroups according to the presence of non-headache symptoms through the entire migraine attack. Between-group differences in demographic and clinical characteristics were assessed using the independent \( t \) test or Mann-Whitney test for continuous variables and the Fisher exact test for categorical variables (version 26.0; SPSS software, IBM, Armonk, NY). Correlations between the frequency of non-headache symptoms experienced during all phases of migraine and patients’ disease activity and disability were assessed using the Spearman’s correlation test.

Results
The main demographic and clinical characteristics of migraine patients are summarized in Table 2. The mean age of subjects was 45 years (range: 19–76), 174 (77%) were women and 48 (21%) patients suffered from migraine with aura. One hundred and twenty-five (56%) patients had chronic migraine and medication overuse was present in 112 (49%) cases. Eighty-one (36%) patients were taking preventive treatments including beta-blockers, antiepileptics, tricyclic antidepressants and Onabotulinumtoxin A. Enrolled patients could have failed preventives in the past with a median of three treatments. Seventeen (7%) patients suffered from depression and 7 (3%) patients had a history of anxiety. On average, patients reported having migraine attacks with a moderate pain intensity and mild cutaneous allodynia. Patients were severely disabled by migraine and reported a severe impact of migraine on their life (Table 2).

Non‑headache symptoms during the prodromal, headache and postdromal phase
Two hundred twenty-three (99%) patients reported having at least one non-headache symptom in one phase of their migraine attacks. One hundred and ninety five (87%) patients reported experiencing at least one non-headache symptom in two phases of their attacks and 122 (54%) patients had at least one non-headache symptom throughout all three phases of migraine. The mean number of non-headache symptoms occurring throughout the entire migraine attack was 1.24 (range: 0–10).

Apart from food craving, the occurrence of non‑headache symptoms was different throughout the three phases of migraine (Cochran’s Q test: \( p < 0.001 \)), being higher during the headache phase than during the prodromal and postdromal phase. The occurrence of food craving did not significantly change over the migraine attack.

| Table 2 | Main demographic and clinical characteristics of migraine patients enrolled in the study |
|-----------------------------------------------|------------------------------------------------------------------|
| Median age [IQ range] (years)                | 46 [34–54]                                                      |
| Women/Men                                     | 174 (77%) - 51 (23%)                                           |
| Median disease duration [IQ range] (years)   | 30 [18–39]                                                      |
| Migraine with aura/Migraine without aura     | 48 (21%) – 177 (79%)                                            |
| Median monthly headache days [IQ range]      | 15 [8–26]                                                       |
| Median monthly migraine days [IQ range]      | 12 [8–18]                                                       |
| Episodic/Chronic migraine                    | 100 (44%) – 125 (56%)                                           |
| Median of preventive treatments used in the past [IQ range] | 3 [0.5–5]                                                      |
| Migraine with medication overuse/ Migraine without medication overuse | 112 (49%) – 113 (51%)                                           |
| Median of monthly tablets of acute treatments [IQ range] | 15 [8–26]                                                      |
| Median MIDAS score [IQ range]                | 36 [15–89]                                                      |
| Median HIT-6 score [IQ range]                | 65 [61–68]                                                      |
| Median NRS score [IQ range]                  | 8 [7–8.5]                                                       |
| Median ASC-12 score [IQ range]               | 4 [1–7]                                                         |

Measures are reported as median and interquartile range (25th – 75th percentiles). Sex, type of migraine, presence of aura and medication overuse are reported as frequencies
Abbreviations: ASC-12 Allodynia Symptom Checklist, HIT-6 Headache Impact Test, IQ interquartile range, MIDAS Migraine Disability Assessment Test, NRS Numeric Rating Scale (NRS)
(Cochran’s Q test: \( p = 0.341 \)) (Fig. 1). The most common reported non-headache symptom in the prodromal phase was neck stiffness \((n = 88, 39\% \) of patients), while difficulty in concentration was most frequently reported during the headache phase \((n = 196, 87\%)\) and fatigue \((n = 89, 40\%)\) during the postdromal phase (Fig. 1). Symptoms with the highest co-occurrence through all three phases were neck stiffness, thirst and abdominal pain (Table 3). No significant differences in the occurrence of non-headache symptoms were found between patients with and without aura, patients on migraine prevention and patients who were not taking preventive treatments as well as between episodic and chronic patients.

Pairing each phase of migraine in couple of two, we found that if lethargy, thirst and food craving occur during the prodromal phase, it is high likely that they will persist during the headache phase, while if phonophobia and mood disturbances occur during the prodromal phase, they will probably occur also during the postdromal phase. If patients report thirst during the headache phase, it is probable that they will experience such symptomatology also during the postdromal phase. The abdominal pain and neck stiffness if present tend to occur during all the three phases of migraine (Table 4).

**Phenotype of patients experiencing non-headache symptoms throughout the entire migraine attack**

Sex, type of migraine and presence of medication overuse were significantly different between patients experiencing non-headache symptoms throughout all phases of migraine and those patients having non-headache symptoms in one or two phases of migraine, being the female gender, chronic migraine and presence of medication overuse more prevalent in patients with a higher number of migraine-associated symptoms. Compared to patients...
### Table 4  Co-occurrence of non-headache symptoms during paired migraine phases

| Subject-reported symptoms | Migraine phases       | Percentage agreement (%) | k value   | p value* | C.I. 95%    |
|---------------------------|-----------------------|--------------------------|-----------|----------|-------------|
| Irritability              | Prodrome - Headache   | 52                       | 0.128     | 0.020    | 0.024-0.0231|
|                           | Prodrome - Postdrome  | 60                       | 0.162     | 0.005    | 0.037-0.280 |
|                           | Headache - Postdrome  | 46                       | 0.113     | 0.005    | 0.042-0.183 |
| Difficulty in concentration| Prodrome - Headache  | 45                       | 0.072     | 0.047    | 0.005-0.138 |
|                           | Prodrome - Postdrome  | 64                       | 0.198     | 0.004    | 0.060-0.320 |
|                           | Headache - Postdrome  | 37                       | 0.055     | 0.068    | 0.004-0.105 |
| Fatigue                   | Prodrome - Headache   | 49                       | 0.112     | 0.021    | 0.021-0.202 |
|                           | Prodrome - Postdrome  | 57                       | 0.075     | 0.256    | 0.056-0.206 |
|                           | Headache - Postdrome  | 52                       | 0.125     | 0.017    | 0.025-0.224 |
| Lethargy                  | Prodrome - Headache   | 71                       | 0.397     | < 0.001  | 0.280-0.500 |
|                           | Prodrome - Postdrome  | 60                       | 0.173     | 0.007    | 0.048-0.298 |
|                           | Headache - Postdrome  | 68                       | 0.195     | 0.003    | 0.060-0.330 |
| Yawning                   | Prodrome – Headache   | 73                       | 0.297     | < 0.001  | 0.142-0.416 |
|                           | Headache - Postdrome  | 72                       | 0.192     | 0.001    | 0.061-0.323 |
|                           | Prodrome - Postdrome  | 78                       | 0.237     | < 0.001  | 0.084-0.389 |
| Difficulty in concentration| Prodrome - Headache  | 75                       | 0.505     | < 0.001  | 0.400-0.610 |
|                           | Prodrome - Postdrome  | 64                       | 0.317     | < 0.001  | 0.416-0.218 |
|                           | Headache - Postdrome  | 74                       | 0.424     | < 0.001  | 0.305-0.543 |
| Lethargy                  | Prodrome - Headache   | 63                       | 0.206     | < 0.001  | 0.108-0.304 |
|                           | Prodrome - Postdrome  | 61                       | 0.148     | < 0.001  | 0.064-0.232 |
|                           | Headache - Postdrome  | 83                       | 0.058     | < 0.001  | 0.110-0.226 |
| Neck stiffness            | Prodrome - Headache   | 91                       | 0.398     | < 0.001  | 0.187-0.609 |
|                           | Prodrome - Postdrome  | 88                       | 0.207     | 0.001    | 0.009-0.405 |
|                           | Headache - Postdrome  | 85                       | 0.152     | 0.023    | 0.026-0.330 |
| Frequent urination        | Prodrome - Headache   | 88                       | 0.291     | < 0.001  | 0.103-0.479 |
|                           | Prodrome - Postdrome  | 81                       | 0.060     | 0.358    | 0.087-0.207 |
|                           | Headache - Postdrome  | 88                       | 0.070     | 0.267    | 0.096-0.236 |
| Mood disturbance          | Prodrome - Headache   | 61                       | 0.293     | < 0.001  | 0.199-0.387 |
|                           | Prodrome – Postdrome  | 52                       | 0.139     | 0.008    | 0.042-0.235 |
|                           | Headache - Postdrome  | 75                       | 0.365     | < 0.001  | 0.228-0.502 |
| Phonophobia               | Prodrome - Headache   | 48                       | 0.157     | < 0.001  | 0.088-0.225 |
|                           | Prodrome - Postdrome  | 41                       | 0.093     | 0.011    | 0.032-0.153 |
|                           | Headache - Postdrome  | 64                       | 0.393     | < 0.001  | 0.258-0.528 |
| Photophobia               | Prodrome - Headache   | 47                       | 0.139     | 0.001    | 0.068-0.209 |
|                           | Prodrome - Postdrome  | 39                       | 0.073     | 0.032    | 0.016-0.129 |
|                           | Headache - Postdrome  | 72                       | 0.271     | < 0.001  | 0.134-0.408 |
| Thirst                    | Prodrome - Headache   | 88                       | 0.408     | < 0.001  | 0.228-0.588 |
|                           | Prodrome - Postdrome  | 86                       | 0.352     | < 0.001  | 0.172-0.532 |
|                           | Headache - Postdrome  | 87                       | 0.186     | 0.005    | 0.006-0.378 |
| Speech difficulty         | Prodrome - Headache   | 69                       | 0.255     | < 0.001  | 0.156-0.354 |
|                           | Prodrome - Postdrome  | 68                       | 0.237     | < 0.001  | 0.135-0.339 |
| Nausea                    | Prodrome - Headache   | 47                       | 0.171     | < 0.001  | 0.140-0.230 |
|                           | Prodrome - Postdrome  | 36                       | 0.080     | 0.005    | 0.069-0.229 |
| Abdominal pain            | Prodrome - Headache   | 95                       | 0.525     | < 0.001  | 0.271-0.779 |
|                           | Prodrome - Postdrome  | 95                       | 0.498     | < 0.001  | 0.752-0.244 |
|                           | Headache - Postdrome  | 98                       | 0.793     | < 0.001  | 0.566-1.020 |

Abbreviations: CI: Confidence interval
* Cohen’s kappa coefficients with 95% CIs. Kappa value > 0.3 (fair-good agreement) with a p < 0.05 was considered significant.
with less symptomatic migraine, patients having non-
headache symptoms throughout all phases of migraine
showed higher headache attack frequency, intake of acute
migraine treatments, HIT-6 and MIDAS scores (Table 5).

**Correlations between non-headache symptoms**
and **migraine disease activity and patients’ disability**
The higher the number of non-headache symptoms
experienced through all phases of migraine, the higher
the MHD \((r = 0.22, p < 0.001)\), MMD \((r = 0.12, p = 0.02)\),
intake of acute treatments \((r = 0.16, p = 0.007)\), MIDAS
\((r = 0.19, p = 0.001)\), HIT-6 \((r = 0.20, p = 0.001)\) and NRS
\((r = 0.09, p = 0.002)\) scores were.

**Discussion**
In this study, we collected data about the occurrence of
non-headache symptoms before, during and after the
headache phase of the migraine attack in a large sample
of migraine patients. In recent years, we have witnessed
an increasing recognition of non-headache symptoms
in association with migraine pain and higher interest in
understanding their pathophysiology. Here, we showed
that almost all patients experienced at least one non-
headache symptom in one phase of the migraine attack
and more than half of our patients experienced at least
one non-headache symptom throughout all phases of
migraine, suggesting that such symptoms are important
migrainous features that need to be investigated.

Previous studies demonstrated that non-headache
symptomatology may precede the headache phase by
up to 72 hours and may persist for 1 day after the head-
ache resolution in the postdromal phase [6, 10]. In line
with previous studies, we found that the most com-
mon reported non-headache symptom in the prodro-
mal phase was neck stiffness while fatigue was mostly
reported during the postdromal phase [5, 8].

Although cognitive, mood and homeostatic changes
are usually described as manifestations of the pro-
drome and postdrome phase [5, 6, 8, 10], our data
showed that, when interrogated, migraine patients
most often recall having non-headache symptoms in
association to the headache pain. During the headache

| Table 5 | Phenotype of patients according to the presence of non-headache symptoms throughout all three phases of migraine |
|---------|--------------------------------------------------------------------------------------|
| **N° of subjects** | 122 (54%) | 103 (46%) | – |
| **Median age [IQ range] [years]** | 45 [34–51] | 47 [34–57] | 0.099 |
| **Women/Men** | 104 (85%) | 18 (15%) | – |
| **Median disease duration [IQ range]** | 30 [16–39] | 30 [20–40] | 0.914 |
| **Migraine with aura/Migraine without aura** | 30 (25%) – 92 (75%) | 18 (17%) – 85 (83%) | 0.253 |
| **Median monthly headache days [IQ range]** | 20 [9–30] | 12 [7–23] | 0.001 |
| **Median monthly migraine days [IQ range]** | 15 [8–20] | 11 [6–17] | 0.056 |
| **Episodic/Chronic migraine** | 40 (33%) – 82 (67%) | 57 (55%) – 46 (45%) | 0.001 |
| **Patients taking/not taking migraine prevention** | 47 (39%)/ 75 (61%) | 34 (33%)/ 69 (67%) | 0.4 |
| **Median of preventive treatments used** | 3 [1–5] | 3 [0–5] | 0.153 |
| **Migraine with medication overuse/ Migraine without medication overuse** | 70 (57%) – 52 (43%) | 40 (39%) – 63 (61%) | 0.004 |
| **Median of monthly tablets of acute treatments [IQ range]** | 16 [8–30] | 12 [5–20] | 0.015 |
| **Median MIDAS score [IQ range]** | 55 [20–103] | 34 [17–69] | 0.004 |
| **Median HIT-6 score [IQ range]** | 66 [63–69] | 64 [61–67] | 0.002 |
| **Median NRS score [IQ range]** | 8 [7–9] | 8 [7–9] | 0.156 |
| **Median ASC score [IQ range]** | 5 [2–8] | 4 [1–7] | 0.130 |

* Independent t test or Mann-Whitney test for continuous variables and the Fisher exact test for categorical variables
phase, beside the canonical migraine-specific symptoms such as nausea, photophobia and phonophobia, we found that difficulty in concentration was the most frequently reported symptom. This result is corroborated by previous magnetic resonance imaging studies showing an altered functional activity of brain networks involved in cognition in migraine patients during the headache phase of spontaneous and induced migraine attacks [22, 23].

Another interesting finding was that non-headache symptoms could start during any phase of the migraine attack and be present during all phases. These findings suggest that the notion of migraine phases coming in a linear and discrete succession may be too simplistic to explain the migraine attack. While migraine should be considered as a continuum with phases occurring in a sequential but overlapping manner.

Interestingly, patients with a higher number of migraine-associated symptoms were females, had a higher attack frequency, disability, intake of acute medications and a higher risk of medication overuse. These findings are consistent with previous data showing that female migraine patients experience a larger number of migraine symptoms, are more disabled by the disease and use greater healthcare resources compared to male patients [24, 25]. Hormonal fluctuations, different biology, genetic features and social environment may explain sex differences in the migraine phenotype [25]. Patients with more non-headache symptoms are those with a higher attack frequency and who require a higher intake of acute treatments, thus increasing the risk of medication overuse.

Our findings suggest that non-headache symptoms may play a role in determining the impact of migraine and patients’ disability. This speculation has important clinical, social and economic implications since patients with frequent symptomatic migraine attacks can have a poor quality of life and experience a large functional impact on their work, school and family life. Future studies assessing the migraine burden should therefore also take into account the repercussion of non-headache symptoms.

Our results provide further insights into the pathophysiology of the migraine attack. We found that neck stiffness, thirst, abdominal pain and food craving can occur throughout the prodromal, headache and postdromal phases of migraine. Of note, these symptoms can be mediated by the hypothalamus, via the orexinergic and dopaminergic systems [3, 4, 11, 12] suggesting its constant involvement during the migraine attack. In accordance with our findings, previous imaging studies have shown an early activation of the hypothalamic, trigeminal, pontine and thalamo-cortical pathways during the prodromal phase, which can persist during the headache phase [11, 22, 26, 27]. An early and persistent activation of the hypothalamic dopaminergic neurons may also explain the co-occurrence of lethargy during the prodromal and headache phase we have found in our sample of patients [11]. Pairing migraine phases in couple of two, we demonstrated that if phonophobia and mood disturbances occur during the prodromal phase, it is high likely that they will be present also during the postdromal phase. Based on these findings, it could be hypothesized an abnormal activity of the auditory, limbic and thalamic systems, which are respectively involved in sound and emotional processing, during the early and late stages of the attack.

The potential effect of acute and preventive treatments on non-headache symptoms has been poorly investigated. Only one study evaluated whether acute migraine treatments could ameliorate non-headache symptoms, showing no effects of triptans on such symptomatology during the postdromal phase [8]. Another recent study showed that patients treated with monoclonal antibodies targeting the calcitonin gene-related peptide (CGRP) pathway reported improvement of non-headache symptoms, like irritability and mood changes. Moreover, this study showed that anti-CGRP monoclonal antibodies could prevent not only the pain but also the onset of non-headache symptoms [28].

Our study is not without limitations. First, patients retrospectively recalled having non-headache symptoms during the different phases of migraine. When asked retrospectively, patients may fail to remember if non-headache symptoms occur during their attacks, especially chronic migraine patients who may lose the original migraine features of their attacks when evolving in a chronic form. Second, we cannot exclude that patients could have misinterpreted postdromal symptoms with acute treatments side effects, thus under- or overestimating the occurrence of postdromal symptomatology. In addition, differences between treated and untreated attacks were not evaluated. Third, the presence of comorbid psychiatric disorders in 24 patients may have influenced patients’ reporting of mood disturbances during the different phases of migraine. Fourth, the sample size of subgroups of patients (e.g. patients with and without aura) was relatively small. Future studies using headache diaries to prospectively collect data regarding the occurrence of non-headache symptoms are needed. Moreover the occurrence of non-headache symptoms during the aura phase should be explored in the future.
Conclusions
Overall, our results confirm the notion that migraine is not simply characterized by recurring pain episodes but is a far more complex neurological disorder with a wide constellation of non-headache symptoms that can play a role in increasing the burden of the disease. Although the identification of distinct migraine phases may ease the study of their underlying mechanisms, it should be kept in mind that most non-headache symptoms may occur throughout all phases. A better characterization of the evolution of non-headache symptoms through the different phases of migraine can improve our understanding of its pathophysiology. Non-headache symptoms should be investigated in routine clinical practice and the effect of acute and preventive treatments on such symptomatology should be further explored in future studies.

Abbreviations
ASC-12: Allodynia Symptom Checklist; CGRP: calcitonin gene-related peptide; HIT-6: Headache Impact Test; MIDAS: Migraine Disability Assessment Test; MHD: Monthly Headache Days; MMD: Monthly Migraine Days; NRS: Numerical Rating Scale.

Authors’ contributions
RM: design of the work, patient enrolment, data analysis, data interpretation and drafting/revising the work. IC: patient enrolment, data analysis, data interpretation and drafting/revising the work. MF: design of the work and drafting/revising the work. He also acted as study supervisor. All the authors approved the current version of the manuscript.

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Availability of data and materials
Data supporting the results of this study are available from the corresponding author, upon reasonable request.

Declarations
Ethics approval and consent to participate
The study was approved by the local ethical committee and written informed consent was obtained from all participants.

Competing interests
The authors declare that they have no conflict of interest related to the publication of this manuscript.

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References
1. GBD 2019 Diseases and Injuries Collaborators (2020) Global burden of 369 diseases and injuries in 204 countries and territories, 1990-2019: a systematic analysis for the global burden of disease study 2019. Lancet Lond Engl 396:1204–1222. https://doi.org/10.1016/S0140-6736(20)30925-9
2. Ashina M (2020) Migraine. N Engl J Med 383:1866–1876. https://doi.org/10.1056/NEJMra1915537
3. Ferrari MD, Goadsby PJ, Burstein R et al (2022) Migraine. Nat Rev Dis Primer 8:1–20. https://doi.org/10.1038/s41572-021-00328-4
4. Goadsby PJ, Holland PR, Martins-Oliveira M et al (2017) Pathophysiology of migraine: a disorder of sensory processing. Physiol Rev 97:553–622. https://doi.org/10.1152/physrev.00034.2015
5. Giffin NJ, Ruggerio L, Lipton RB et al (2003) Premonitory symptoms in migraine: an electronic diary study. Neurology 60:935–940. https://doi.org/10.1212/01.WNL.0000052998.58526.A9
6. Karsan N, Bose PR, Thompson C et al (2020) Headache and non-headache symptoms provoked by nitroglycerin in migraineurs: a human pharmacological triggering study. Cephalalgia 40:828–841. https://doi.org/10.1177/033310241910114
7. Bose P, Goadsby PJ (2016) The migraine postdrome. Curr Opin Neurol 29:299–301. https://doi.org/10.1097/WCO.0000000000000310
8. Giffin NJ, Lipton RB, Silberstein SD et al (2016) The migraine postdrome. Neurology 87:309–313. https://doi.org/10.1212/WNL.0000000000027289
9. Ong JJY, De Felice M (2018) Migraine treatment: current acute medications and their potential mechanisms of action. Neurother J Am Soc Exp Neurother 15:274–290. https://doi.org/10.1177/1129201618810530
10. Karsan N, Perez-Rodriguez A, Nagaraj K et al (2021) The migraine postdrome: spontaneous and triggered phenotypes. Cephalalgia Int J Headache 41:721–730. https://doi.org/10.1017/s333102420975401
11. Karsan N, Goadsby PJ (2018) Biological insights from the premonitory symptoms of migraine. Nat Rev Neurol 14:699–710. https://doi.org/10.1038/s41588-018-0098-4
12. Messina R, Gollion C, Christensen RH, Amin FM (2022) Functional MRI in migraine population. Ann Neurol 63:148–158. https://doi.org/10.1002/ana.21451
13. Maniyar FH, Sprenger T, Schankin C, Goadsby PJ (2014) The origin of nausea in migraine—a PET study. J Headache Pain 15:84. https://doi.org/10.1186/s12320-013-0125-4
14. Maniyar FH, Sprenger T, Schankin C, Goadsby PJ (2014) Photic hypersensitivity in the premonitory phase of migraine—a positron emission tomography study. Eur J Neurol 21:1178–1183. https://doi.org/10.1111/ene.12451
15. (2018) Headache Classification Committee of the International Headache Society (IHS) The International Classification of Headache Disorders, 3rd edition. Cephalalgia 38:1–211. https://doi.org/10.1177/0333102417738202
16. Aicher B, Peil H, Peil B, Diener H-C (2012) Pain measurement: visual analogue scale (VAS) and verbal rating scale (VRS) in clinical trials with OTC analgesics in headache. Cephalalgia Int J Headache 32:185–197. https://doi.org/10.1177/0333102411430866
17. Lipton RB, Bigal ME, Ashina S et al (2008) Cutaneous allodynia in the migraine population. Ann Neurol 63:148–158. https://doi.org/10.1002/ana.21211
18. Stewart WF, Lipton RB, Dowson AJ, Sawyer J (2001) Development and testing of the migraine disability assessment (MIDAS) questionnaire to assess headache-related disability. Neurology 56:520–528. https://doi.org/10.1212/01.wnl.0000065667.67763.6d
19. Kosinski M, Bayliss MS, Bjorner JB et al (2003) A six-item short-form survey for measuring headache impact: the HIT-6. Qual Life Res Int J Qual Life Anal 12:963–974. https://doi.org/10.1023/A: 1026119331193
20. Sim J, Wright CC (2005) The kappa statistic in reliability studies: use, interpretation, and sample size requirements. Phys Ther 85:257–268
21. Hong H, Choi Y, Hahn S et al (2014) Normogram for sample size calculation on a straightforward basis for the kappa statistic. Ann Epidemiol 24:673–680. https://doi.org/10.1016/j.annepidem.2014.06.097
22. Karsan N, Bose PR, O’Daly O et al (2020) Alterations in functional connectivity during different phases of the triggered migraine attack. Headache J Head Face Pain 60:1244–1258. https://doi.org/10.1111/head.13865
23. Coppola G, Di Renzo A, Tinelli E et al (2018) Resting state connectivity between default mode network and insula encodes acute migraine headache. Cephalalgia Int J Headache 38:846–854. https://doi.org/10.1177/0333102417715230
24. Buse DC, Loder EW, Gorman JA et al (2013) Sex differences in the prevalence, symptoms, and associated features of migraine, probable migraine and other severe headache: results of the American Migraine Prevalence and Prevention (AMPP) study. Headache J Head Face Pain 53:1278–1299. https://doi.org/10.1111/head.12150

25. Todd C, Lagman-Bartolome AM, Lay C (2018) Women and migraine: the role of hormones. Curr Neurol Neurosci Rep 18:42. https://doi.org/10.1007/s11910-018-0845-3

26. Schulte LH, May A (2016) The migraine generator revisited: continuous scanning of the migraine cycle over 30 days and three spontaneous attacks. Brain J Neurol 139:1987–1993. https://doi.org/10.1093/brain/aww097

27. Messina R, Filippi M, Goadsby PJ (2018) Recent advances in headache neuroimaging. Curr Opin Neurol 31:379–385. https://doi.org/10.1097/WCO.0000000000000573

28. Iannone LF, De Cesaris F, Ferrari A et al (2022) Effectiveness of anti-CGRP monoclonal antibodies on central symptoms of migraine. Cephalalgia Int J Headache 42:1323–1330. https://doi.org/10.1177/0333102422111526

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