Evidence of an increased neuronal activation-to-resting glucose uptake ratio in the visual cortex of migraine patients: a study comparing $^{18}$FDG-PET and visual evoked potentials

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Abstract

Background: Migraine attacks might be triggered by a disruption of cerebral homeostasis. During the interictal period migraine patients are characterized by abnormal sensory information processing, but this functional abnormality may not be sufficient to disrupt the physiological equilibrium of the cortex unless it is accompanied by additional pathological mechanisms, like a reduction in energetic reserves. The aim of this study was to compare resting cerebral glucose uptake (using positron emission tomography ($^{18}$fluorodeoxyglucose-PET)), and visual cortex activation (using visual evoked potentials (VEP)), between episodic migraine without aura patients in the interictal period and healthy volunteers.

Methods: Twenty episodic migraine without aura patients and twenty healthy volunteers were studied. $^{18}$FDG-PET and VEP recordings were performed on separate days. The overall glucose uptake in the visual cortex-to-VEP response ratio was calculated and compared between the groups. Additionally, PET scan comparisons adding area under the VEP curve as a covariate were performed. For case-wise analysis, eigenvalues from a specific region exhibiting significantly different FDG-PET signal in the visual cortex were extracted. Standardized glucose uptake values from this region and VEP values from each subject were then coupled and compared between the groups.

Results: The mean area under the curve of VEP was greater in migraine patients compared to healthy controls. In the same line, patients had an increased neuronal activation-to-resting glucose uptake ratio in the visual cortex. Statistical parametric mapping analysis revealed that cortical FDG-PET signal in relation to VEP area under the curve was significantly reduced in migraineurs in a cluster extending throughout the left visual cortex, from Brodmann’s areas 19 and 18 to area 7. Within this region, case-wise analyses showed that a visual neuronal activation exceeding glucose uptake was present in 90% of migraine patients, but in only 15% of healthy volunteers.

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In migraine, metabolism has been suggested to play a major role in the development of the disease [3]. The fact that similar cortical reactivity profiles can be found in both migraine patients and their asymptomatic first degree relatives suggests the existence of additional pathologic mechanisms in migraine sufferers that, when associated to visual hyper-responsivity, lead to the development of the disease [3].

In parallel with cortical hyper-responsivity, cerebral metabolism has been suggested to play a major role in migraine [4–6]. Hypotheses argue that reduced mitochondrial energy and ATP levels observed in the cortex of patients between attacks [7–9] might make them unable to deal with an energetically more demanding neuronal activity [5, 10]. This imbalance would later translate into a disruption in cortical homeostasis, with subsequent activation of the trigeminovascular system. Such view is supported by the favourable clinical response to metabolic enhancers [11] and ketogenic diet [12, 13] observed in migraine patients, as well as the capability of metabolic challenges to trigger migraine attacks [6].

**Background**

Between attacks, migraine patients often exhibit abnormal sensory information processing [1]. Cortical hyper-responsivity during any kind of sensory stimulus repetition, including visual, is the most common electrophysiological feature found in episodic migraine patients during the interictal state [2]. Such alteration however, although highly prevalent among migraineurs, is by itself insufficient to entirely explain migraine’s pathophysiology [3]. The fact that similar cortical reactivity profiles can be found in both migraine patients and their asymptomatic first degree relatives suggests the existence of additional pathologic mechanisms in migraine sufferers that, when associated to visual hyper-responsivity, lead to the development of the disease [3].

In the present study we sought to determine if interictal glucose metabolism in the visual cortex is proportional to visually-induced neuronal activation. For this purpose, we assessed cerebral glucose uptake, and recorded visual evoked potentials (VEP) in healthy volunteers and migraine patients. Our main hypothesis was that the interictal responsivity of the visual cortex in migraine patients would exceed the resting glucose uptake, rendering migraineurs vulnerable to a disruption of metabolic homeostasis in times of increased neuroenergetic demands.

**Methods**

**Study participants**

The study involved twenty episodic migraine without aura patients (MO; mean age (SD): 31.1 (±12.6), 85% fem) diagnosed in accordance with The International Classification of Headache Disorders 3rd edition (Beta version) [17] and twenty healthy volunteers (HV; mean age (SD): 36.1 (±11.4), 75% fem) who did not report having first degree relatives suffering from recurrent headaches of any type. There were no significant differences in age or gender proportions between groups. Participants were recruited among University students or their families or via our headache clinic. None of them took any medication on a daily basis (other than the contraceptive pill), and they were all free of any systemic or neurological disease other than migraine. Patients were not under any prophylactic treatment at the time of recordings, nor had they been for at least 30 preceding days. The mean number of monthly migraine days determined by headache diary inspection (during the month of the recordings) was 4.3 ± 2.5. All patients were recorded at an interval of at least 72 h before and after an attack. The study was approved by the Institution’s ethics committee (Centre Hospitalier Régional de la Citadelle, Liège, Belgium – protocol n°1422) and conducted following the principles of the Declaration of Helsinki. All participants gave their written informed consent.
Visual evoked potentials recordings and analysis

Visual evoked potentials (VEP) were used as a marker of visual cortex responsiveness. Recordings were performed in the electrophysiology laboratory of the Headache Research Unit (Neurology Department, Centre Hospitalier Régional de la Citadelle, Liège, Belgium). Subjects sat on a comfortable armchair, in a quiet room with dimmed light. Needle electrodes were placed at Oz (active) and Fz (reference) of the 10–20 EEG system. With the left eye patched, participants were instructed to fixate on a red dot in the centre of a screen displaying a black and white reversing checkerboard pattern (contrast of 80%, mean luminance 50 cd/m²) at temporal and spatial stimulating frequencies of 1.55 Hz (3.1 reversals/second) and 68° respectively. Six hundred epochs, each lasting 250 ms, were uninterruptedly recorded at a sampling rate of 5.000 Hz using a CED™ power 1401 device (Cambridge Electronic Design Ltd., Cambridge, UK). After DC subtraction, recordings were exported to EEGLAB [18] (an open-source MATLAB (The MathWorks Inc.) toolbox for electrophysiological signal processing), where they were band-pass filtered (low pass 100 Hz, high pass 1 Hz). Artifacted epochs exceeding two standard deviations of the channel mean limit were rejected (< 7% of epochs). We extracted the area under the curve (AUC) of each single trial, and thereafter averaged these individual values. This method is deemed to be less affected by phase synchronization and thus to provide a more straightforward measure of neuronal activation [19].

FDG-PET acquisition and analysis

PET acquisitions were made in the Radiodiagnostics Department of the Centre Hospitalier Universitaire (CHU) Sart Tilman, Liège, Belgium (Prof. R. Hustinx) using a Gemini TF PET/computed tomography (CT) scanner (Philips, Eindhoven, The Netherlands). Resting cerebral metabolism was studied 30 min after intravenous injection of 150 MBq FDG. Mean blood glucose level was 89.5 (range:110–77). Subjects were injected and scanned in a dark room with minimal environmental noise. They were instructed to maintain their eyes closed during the scan. Images were reconstructed using an iterative list mode time-of-flight algorithm. Corrections for attenuation, dead-time, random and scatter events were applied. PET acquisitions were analysed using Statistical Parametric Mapping version 12 (SPM12, Wellcome Trust Centre for Neuroimaging, http://www.fil.ion.ucl.ac.uk/spm) implemented in MATLAB 7.4.0 (MathWorks Inc., Sherborn, MA, USA). Images were first manually reoriented and then spatially normalised into a standard stereotactic space using an MNI PET template (Montreal Neurological Institute) and smoothed using an 8 mm full-width-half-maximum (FWHM) isotropic kernel. We performed global normalisation by applying proportional scaling. Cerebral regions identification and masks were established using the WFU PickAtlas toolbox (Wake Forest University School of Medicine, Advanced NeuroScience Imaging Research lab (ANSIR), Winston-Salem, NC, U.S.A.). PET scans were compared between groups (1) without adding any covariates, and (2) using the interaction with the averaged area under the VEP as regressor. An exploratory (p < 0.001 uncorrected) analysis followed by a more stringent comparison (p family-wise error[FWE] < 0.05 correction at a whole brain level) were performed. Only statistically significant clusters within regions of the visual cortex are reported in the results. To estimate the neuronal activation-to-overall resting glucose uptake ratio in the visual cortex, eigenvalues corresponding to Brodmann’s areas 17, 18, and 19 were extracted using a volumetric mask. The ratio between these values, and the mean area under the VEP curve was then calculated. In addition, based on the statistical parametric mapping results, for case-wise analysis a specific volumetric mask was generated upon the cluster in the visual cortex exhibiting a statistically significant difference in FDG-uptake between groups when including VEP-AUC as a covariate. By using this volumetric mask, eigenvalues corresponding to this specific region were extracted for each subject.

Statistical analysis

Statistical analyses were performed in Prism version 6.00 for Windows (GraphPad Software, La Jolla, California, USA). Continuous variables were compared using the t-test. The assumption of normality was evaluated using a Shapiro-Wilk normality test. For case-wise analyses, mean area under the VEP curve (accounting for visual activity) and eigenvalues corresponding to the specific cluster of the visual cortex exhibiting significant FDG-uptake differences (accounting for metabolic activity) were Z-transformed and paired for each subject. Proportions of participants with a visual > metabolic Z-score in each group were compared using Fisher’s exact test. The significance level was set at p < 0.05 for all statistical analyses.

Results

The mean area under the curve of VEP was significantly greater in migraine patients compared to healthy controls (1091 μV² ± 164.8 vs. 362.8 μV² ± 167.7; p < 0.01) (Fig. 1).

Four clusters of reduced metabolism where initially observed (uncorrected p < 0.001 level) in the visual cortex of migraineurs, but they did not withstand the correction for multiple comparisons. No clusters of increased PET signal were found in the visual cortex of migraineurs.
The mean ratio between the VEP-AUC and overall glucose uptake in the visual cortex (BA 17, 18 and 19) was three times higher in migraine patients (12.73 ± 1947 vs. 4.183 ± 1955; p < 0.01).

When adding VEP-AUC as a covariate, statistical parametric mapping comparisons revealed a cluster of significantly reduced uptake in patients, with three peaks extending over Brodmann’s areas 19, 18 and 7 on the left (− 20, − 78, 30 T = 5.66 peak-level p [FWE-corr] = 0.039; − 8, − 78, 20 T = 4.84; − 20, − 74, 42 T = 4.36, all cluster-level pFWE-corr = 0.021) (Fig. 2). Eigenvalues from this specific cluster were extracted and standardized for case-wise analysis, where a visual activation Z-score exceeding the glucose uptake Z-score was found in 90% of migraine patients, but in only 15% of healthy volunteers (p < 0.001, Fisher’s exact test) (Fig. 3).

Discussion
In this study, using 18fluorodeoxyglucose-positron emission tomography acquired at rest and pattern reversal-visual evoked potentials, we compared interictal glucose uptake and neuronal activation in the visual cortex of migraine patients and healthy volunteers. Our FDG-PET data confirm the finding of an interictally decreased metabolism in the occipital areas previously shown using phosphorus magnetic resonance spectroscopy [7–9]. An additional, novel finding, is that the ratio between stimulation-induced neuronal activation and resting glucose uptake differs between migraine patients and healthy controls. Using statistical parametric mapping, we identified the specific region of the visual cortex where this difference reaches its maximum. The potential pathophysiological implications of these findings are discussed below.

According to most neurophysiological studies, migraine patients exhibit cortical hyper-responsivity to repeated sensory stimuli between attacks [20]. Although this may constitute a favourable evolutionary trait [21], enhanced responsivity has a price to be paid in terms of cerebral metabolism. In the brain, the visual system ranks amongst the most energy-consuming systems [22]. This renders it more susceptible to metabolic imbalance when energetic demands increase, energetic supplies decrease, or in a combination of both.

The occurrence of a metabolic strain in the cortex is a plausible hypothesis for igniting a migraine attack [23]. The cornerstone of the metabolic strain model is the unbalanced relationship between enhanced sensory processing and reduced metabolic offers [5]. According to this model, unmet neuronal metabolic demands in the cortex would result in a focal disruption of homeostasis and release of molecules capable of activating the trigemino-vascular system [24–26]. This mechanism, although orchestrated by neurons [26], relies on astrocytes as concertmasters [27]. By determining an area of sensory-metabolic mismatch (suggestive of the presence of a higher cortical responsivity and relatively lower resting energy reserves) in the visual cortex of migraine patients, our results provide supportive experimental evidence to the initiatory phase of the metabolic strain model. Intuitively, energy reserves may be further decreased by daily life stressful and energy consuming events [28], or by the sensory hypersensitivity that patients present even between crises, which extends up to the premonitory phase [29] when, along with the energetic deficit, may worsen up to the tipping point of the attack. This view is supported by a previous study in which the authors simultaneously recorded VEP and cerebral blood flow velocity responses (VEFR) in a group of migraineurs. They observed an interictal increase of neurovascular coupling in patients interictally that, empirically, corresponds to higher neural activity, which
might further reduce energy reserves [30]. Hence, the presence of an increase of energetic requirements in the absence of adequate energetic supplies would be a plausible reason for the brain to activate its major alarm system, the trigeminovascular system, that might ignite a migraine attack in order to prevent the system from overloading, and thus re-establish cortical homeostasis [4]. This is supported by three out of four studies where authors positively triggered migraine using sensory overload [31–34]. On the other hand, metabolic challenges have also been shown to provoke migraine attacks. From an observational perspective, skipping meals is the third most common migraine trigger reported by patients [35]. In the laboratory, Schoonman et al. reported provoking headache in six out of fourteen migraine patients exposed to normobaric hypoxia [36], and more recently, in an elegantly designed study also addressing the effects of hypoxia, Arngrim et al. reported migraine-like attacks in eight out of fifteen migraine patients, accompanied by definite aura in three, and possible aura in another four [37].

Our results do not necessarily pull in one direction, favouring sensory over metabolic factors or vice versa, but they rather imply that the physiological sensory-metabolic equilibrium is abnormal in migraine between attacks. However, it is of interest that resting brain energy metabolism remains low even during migraine attacks [38–40] when sensory hyper-responsivity tends to normalize [29]. This suggests that the former might be a trait factor permanently predisposing to the next crisis, while the latter would be more of a state-dependent phenomenon.

As mentioned before, the sensory profile of migraine patients makes them susceptible of sensory overload [1], and therefore in the need of protective mechanisms. Compensatory metaplastic changes have been hypothesized to explain the between-subject variability of electrophysiological responses in migraine groups in previous studies [41]. Thus, it could be debated whether the reduced resting glucose uptake in migraine patients we observed constitutes compensatory mechanism aiming to decrease cortical activation. However, given the heightened visual responsiveness

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**Fig. 2** Areas of increased metabolism in healthy controls compared with migraine patients using the interaction with area under the VEP as regressor. Clockwise from top left: (a) coronal view, (b) sagittal view, (c) axial view, and (d) "glass brain" representation and design matrix.
we recorded in patients, this alternative hypothesis seems unlikely.

One cannot discuss cortical metabolism and homeostasis without considering the role of astrocytes (a deficient uptake of glucose by astrocytes could explain our FDG-PET findings in migraine patients based on current evidence [42–44]). Neurons have a very scarce, if any, energetic reserve. Because of that, their increased metabolic demands upon stimulation basically rely on the energetic supplies provided by astrocytes. Unlike neurons, during rest astrocytes store glucose in the form of glycogen. When the cortex is stimulated, glycogen reserves in astrocytes are rapidly transformed into less complex energetic substrates, which are then provided to neurons in order to fulfil their enhanced metabolic needs (Fig. 4) [42]. In exchange, astrocytes buffer excessive glutamate molecules released in the synaptic cleft [45]. Other functions of astrocytes, besides regulating neuro-metabolic coupling and glutamate concentrations, include controlling the amount of extracellular potassium [46], and acting as regulators of cerebral blood flow [47]. Indeed, the influence of astrocytes on synaptic transmission is such, that the concept of a tripartite synapse (presynaptic neuron, astrocyte, and postsynaptic neuron) is steadily solidifying [48]. In fact, when altered, deficient clearance of potassium and glutamate by astrocytes seems to explain the pathogenesis of specific forms of migraine with monogenetic inheritance [49]. For the case of more common types of migraine, the pathophysiological implication of astrocytes is less well elucidated, although genetic studies seem to also point in their direction [50, 51].
Interestingly, the visual cortex has the lowest astrocyte/neuron density in the brain, which renders it more susceptible to homeostatic disturbances [52, 53]. Prior clinical [54] and experimental [55] evidence has suggested that waves of cortical spreading depression, the culprit of migraine aura, originate in the visual cortex. Visual area V3A was specifically pointed to as the initial region in one report [53]. Another study described increased cortical thickness of this area in migraine without aura patients [56]. It might be worthwhile to mention that the cluster of mismatch we found in the present study extends over visual areas V3A and V7. Whether this observation is coincidental or entails clinical significance remains to be determined.

Our study has several limitations. First, PET scans and VEP where not recorded concomitantly and thus, the possibility of an important compensatory metabolic increase in migraine patients during stimulation remains open. Also, it is worth mentioning that a cycling pattern of increased susceptibility probably determined by the hypothalamus, or its limbic controls, is probably involved in migraine pathogenesis [57], and thus sensory-metabolic coupling might be preserved at one time point and impaired at another. Finally, even if there exists a metabolic imbalance during stimulation in the visual cortex of migraine patients, secondary activation of the trigeminovascular system would need to be further ascertained. In the future, studies addressing these issues would be necessary to clarify the possible role of a disruption in cortical homeostasis due to a sensory-metabolic disequilibrium in migraine pathophysiology.

Conclusion

Our findings indicate the presence of an area of increased neuronal activation-to-resting glucose uptake ratio in the visual cortex of migraine patients between attacks. Despite some methodological reservation, this observation supports the concept that an activity-induced rupture of cerebral metabolic homeostasis may be a cornerstone in migraine pathophysiology. The potential physiopathological implications of such finding should be explored in depth in future studies.

Abbreviations

- FDG-PET: Fluorodeoxyglucose-positron emission tomography; AUC: Area under the curve; VEP: Visual evoked potentials

Acknowledgements

This article has been selected as the winner of the 2018 Enrico Greppi Award. The Enrico Greppi Award is made to an unpublished paper dealing with clinical, epidemiological, genetic, pathophysiological or therapeutic aspects of headache. The Italian Society for the Study of Headaches (SSC) sponsors this award, and the award is supported through an educational grant from Teva Neuroscience. This article did not undergo the standard peer review process for The Journal of Headache and Pain. The members of the 2018 Enrico Greppi Award Selection Committee were: Francesco Pierelli, Paolo Martelletti, Lyn Griffiths, Simona Sacco, Andreas Straube and Cenk Ayata.

The authors thankfully acknowledge Prof. Roland Hustinx, Chief of the Nuclear Medicine Service at the Centre Hospitalier Universitaire de Liège, for his collaboration on this study (access to FDG-PET).

The contribution of the G.B. Bietti Foundation in this paper was supported by the Italian Ministry of Health and Fondazione Roma.

Funding

This project forms part of the EUROHEADPAIN project - FP7 n° 602633, and received support from the Fonds d’Investissements de Recherche Scientifique (FIRS) of the CHU de Liège.

Availability of data and materials

Further data from the underlying research material can be obtained upon request to the corresponding author.

Authors’ contributions

ML contributed in the study design, data acquisition, data processing (focus on electrophysiology), statistical analyses, and wrote the first draft; KDO contributed in the study design, data acquisition, data processing (focus on neuroimaging) and statistical analyses; KD recruited subjects with DM; DM wrote the initial Euroheadpain protocol with JS; GC, AM de N, JS, and DM contributed in the study design and revised the drafts of the manuscript; FS and VP revised the drafts of the manuscript. All authors read and approved the final version.

Ethics approval and consent to participate

The study was approved by the Institution’s ethics committee (EC number 412, Centre Hospitalier Régional de la Citadelle, Liège, Belgium – protocol n° 1422) and conducted following the principles of the Declaration of Helsinki. All participants gave their written informed consent.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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Received: 27 April 2018 Accepted: 21 June 2018

Published online: 05 July 2018

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