
Downloaded from:

Usage Guidelines:
Please refer to usage guidelines at contact lib-eprints@bbk.ac.uk.
An inability to exclude visual noise in migraine.

Marc S. Tibber\textsuperscript{a}, Maria Kelly\textsuperscript{b}, Ashok Jansari\textsuperscript{c}, Steven C. Dakin\textsuperscript{a,d}, Alex J. Shepherd\textsuperscript{b}


\textsuperscript{a}Institute of Ophthalmology, University College London, Bath Street, London, EC1V 9EL, UK.
\textsuperscript{b}Department of Psychological Sciences, Birkbeck College, Malet Street, London, WC1E 7HX.
\textsuperscript{c}School of Psychology, University of East London, London, E15 4LZ, UK.
\textsuperscript{d}NIHR Biomedical Research Centre at Moorfields Eye Hospital NHS Foundation Trust and UCL Institute of Ophthalmology, London, EC1V 9EL, UK.

Corresponding author:
Marc S. Tibber [mtibber@yahoo.com; Institute of Ophthalmology, University College London, Bath Street, London EC1V 9EL, UK; Tel: +44 (0)20 7608 6924].

Keywords: Migraine, vision, noise, coherence, motion.
1. Abstract:

Purpose: People with migraine are relatively poor at judging the direction of motion of coherently-moving signal-dots when interspersed with noise-dots drifting in random directions, a task known as motion coherence. Although this has been taken as evidence of impoverished global pooling of motion signals, it could also arise from unreliable coding of local direction (of each dot), or an inability to segment signal from noise (noise-exclusion). The aim of this study was to determine how these putative limits contribute to impoverished motion processing in migraine.

Methods: Twenty-two participants with migraine (mean age: 34.7±8.3 years; 16 female) and 22 age and sex matched controls (mean age: 34.4±6.2 years) performed a motion coherence task and a motion equivalent noise task, the latter quantifying local and global limits on motion processing. In addition, participants were tested on analogous equivalent noise paradigms involving judgements of orientation and size, so that the specificity of any findings (to visual dimension) could be ascertained.

Results: Participants with migraine exhibited higher motion coherence thresholds than controls (p=0.01, independent t-test). However, this difference could not be attributed to deficits in either local or global processing since they performed normally on all equivalent noise tasks (p>0.05, multivariate analyses of variance).

Conclusions: These findings indicate that motion perception in the participants with migraine was limited by an inability to exclude visual noise. We suggest that this is a defining characteristic of visual dysfunction in migraine, a theory that has the potential to integrate a wide range of findings in the literature.
Migraine is an episodic disorder characterised by throbbing (commonly unilateral) head pain, which may be accompanied by nausea, vomiting and an aversion to sound or light\(^1\). In approximately 30% of cases, a transient sensory and/or motor disturbance known as an aura is also experienced\(^2\). Certain visual stimuli can also trigger a migraine attack\(^3\) and numerous studies have shown that individuals with migraine exhibit subtle differences in visual psychophysical performance, both ictally and interictally (see reviews\(^4, 5\)). This is particularly the case for tasks involving judgements of visual motion\(^6\).

Processing of visual motion relies on at least two hierarchical processing stages. In the primary visual cortex (area V1), motion is processed \textit{locally}, i.e. cells are sensitive to the direction of motion within a small region of space\(^7\). This information is then relayed to the medial temporal (MT) and medial superior temporal (MST) areas, where it is integrated to form a global motion percept\(^8\). People with migraine seemingly process \textit{local} motion normally, since they perform as well as a control group when asked to discriminate or classify the direction of a stimulus containing a single direction of motion\(^6, 9-11\). However, people with migraine perform relatively poorly on motion coherence tasks where the participant must classify the direction of motion of a set of signal-dots moving coherently (in one direction) but interspersed with noise-dots drifting in random directions (Fig. 1A)\(^6, 9, 10, 12-14\).
Since the signal-direction in a coherence task cannot be determined from a single dot’s trajectory, the participant must make a judgement of global motion direction. As a result, high motion coherence thresholds are often taken as evidence of a selective deficit in global motion pooling. However, motion coherence judgements can be limited not only by global integration, but also, by unreliable local processing. This could be the case, for example, if higher cortical areas inherit input from V1 cells prone to high levels of random firing, i.e. elevated internal noise. A further limit on motion coherence performance is defined by an observer’s ability to segregate signal from noise dot directions. Thus, computational models show that human observers perform much better on coherence tasks than would be expected if they used a pure pooling strategy, suggesting that they are capable of selectively monitoring directions of interest.

To try and disentangle these putative limits to motion processing we used a technique known as equivalent noise (EN) analysis. This psychophysical paradigm allows performance to be parcellated into independent estimates of local and global processing. Similar to the motion coherence paradigm, EN analysis requires participants to classify the direction of motion of signal dots that are corrupted by noise. However, in EN analysis, noise is added by manipulating the standard deviation of the distribution of directions presented, rather than adding noise dots that drift in random directions (Fig. 1B). As a result, every dot contributes to the signal, and the optimum strategy is to integrate all directions of motion in the stimulus. Consequently, an estimate of global processing is obtained that does not rely on the participants’ ability to exclude noise. Further, by measuring performance in the
absence (as well as in the presence) of noise, an independent estimate of a participant’s ability to process information locally is also available.

We sought to determine if motion processing in migraine is (a) limited by local processing, global processing and/or noise exclusion, and (b) part of a more general integration deficit. To this end, participants with and without migraine were tested on a series of matched psychophysical tasks. A motion coherence paradigm was used to assess each participant’s ability to classify the direction of signal motion whilst excluding random noise. Independent estimates of local and global motion processing performance were obtained using a motion EN paradigm. Finally, to assess the specificity of any findings to motion processing participants undertook analogous EN tasks that probed local and global processing for judgements of orientation and size.

3. Materials and Methods:

Ethics approval was granted by the University of East London Psychology Research Ethics Committee and the Department of Psychological Sciences Ethics Committee at Birkbeck College. Informed written consent was obtained from each participant in accordance with the declaration of Helsinki.

Participants

Data were gathered from 22 participants with migraine (MG) and 22 migraine-free control participants (CON) (Table 1). The two groups were matched for sex (16
female) and did not differ significantly with respect to age [mean age: 34.7±8.3 (MG) and 34.4±6.2 years (CON); t(42)=0.04, p=0.97]. All participants with migraine fulfilled the International Headache Society (2004) diagnostic criteria for migraine without aura (MO) or migraine with visual aura (VA), and had been diagnosed previously by a general practitioner or neurologist. All participants had a minimum visual acuity of 20/20 binocularly (with or without optometric correction). No participant had a history of mental illness and none were taking daily medication at the time of testing.

**General procedure**

The experiment lasted 60-75 minutes and consisted of: (i) a brief test of visual acuity (assessed using a hand-held LogMar near visual acuity chart); (ii) a customised questionnaire about basic demographics and migraine history; (iii) a motion coherence paradigm; (iv) three EN paradigms, which probed local and global processing for judgements of visual orientation, motion and size (separately). Individual EN and coherence tasks were blocked and presented in a random order to avoid sequence effects. All responses were given verbally and relayed to the computer by the experimenter.

**Motion Coherence procedure**

Participants classified the direction of motion of a number of coherently moving dots (the signal) embedded in noise. All signal dots were restricted to motion in the horizontal plane (all left or all right on any given trial). Noise was added to the
stimulus by assigning a subset of dots directions of motion that were randomly
sampled from a flat distribution (Fig. 1A). Under the control of QUEST\textsuperscript{18}, an adaptive
staircase procedure manipulated the level of coherence on each trial, where coherence
was defined as the percentage of dots that constituted the signal. The staircase
converged on the level of coherence necessary for each participant to correctly
ascertain the direction of motion on 82% of trials: the motion coherence threshold (see
Supplementary Fig. 1A for further details). Lower coherence thresholds therefore
reflected superior performance, indicating that the participant needed fewer signal
dots to correctly identify the direction of signal motion. The staircase terminated after
75 trials and was preceded by 15 practice trials.
Table 1. Migraine group demographics and details of migraine history. Details are provided for:
(1) Type (MO: migraine without aura; VA: migraine with visual aura); (2) Sex (F: female; M: male);
(3) Age; (4) Onset (age of migraine onset); (5) Freq 1 (number of migraine attacks experienced within
the last three months); (6) Freq 2 (number of migraine attacks experienced within the last year); (7)
Last (time, in weeks, since last migraine attack); (8) Duration (average duration, in hours, of a migraine
attack when painkillers are administered); (9) Severity (index of migraine severity, derived from the
multiplication of average migraine duration by the number of years migraine has been experienced).

<table>
<thead>
<tr>
<th>Type</th>
<th>Sex</th>
<th>Age</th>
<th>Onset</th>
<th>Freq 1</th>
<th>Freq 2</th>
<th>Last</th>
<th>Duration</th>
<th>Severity</th>
</tr>
</thead>
<tbody>
<tr>
<td>MO</td>
<td>F</td>
<td>21</td>
<td>13.5</td>
<td>1</td>
<td>3.5</td>
<td>8</td>
<td>60</td>
<td>144</td>
</tr>
<tr>
<td>MO</td>
<td>F</td>
<td>25</td>
<td>23</td>
<td>4</td>
<td>16</td>
<td>2</td>
<td>6.5</td>
<td>192</td>
</tr>
<tr>
<td>MO</td>
<td>F</td>
<td>38</td>
<td>16</td>
<td>4</td>
<td>20</td>
<td>1</td>
<td>24</td>
<td>384</td>
</tr>
<tr>
<td>MO</td>
<td>F</td>
<td>39</td>
<td>30</td>
<td>3.5</td>
<td>12</td>
<td>1</td>
<td>24</td>
<td>144</td>
</tr>
<tr>
<td>MO</td>
<td>F</td>
<td>40</td>
<td>5</td>
<td>6</td>
<td>24</td>
<td>2</td>
<td>48</td>
<td>517.5</td>
</tr>
<tr>
<td>MO</td>
<td>F</td>
<td>43</td>
<td>32</td>
<td>3</td>
<td>10</td>
<td>4</td>
<td>60</td>
<td>108</td>
</tr>
<tr>
<td>MO</td>
<td>M</td>
<td>23</td>
<td>16</td>
<td>2</td>
<td>6.5</td>
<td>4</td>
<td>24</td>
<td>32</td>
</tr>
<tr>
<td>MO</td>
<td>M</td>
<td>34</td>
<td>11.5</td>
<td>2</td>
<td>3</td>
<td>5</td>
<td>96</td>
<td>26.25</td>
</tr>
<tr>
<td>MO</td>
<td>M</td>
<td>38</td>
<td>28</td>
<td>3</td>
<td>10</td>
<td>3</td>
<td>4</td>
<td>22.5</td>
</tr>
<tr>
<td>MO</td>
<td>M</td>
<td>40</td>
<td>5.5</td>
<td>3</td>
<td>15</td>
<td>2.5</td>
<td>60</td>
<td>80</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>21</td>
<td>10</td>
<td>3</td>
<td>12</td>
<td>2</td>
<td>6.5</td>
<td>100</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>24</td>
<td>19</td>
<td>12</td>
<td>182</td>
<td>0.29</td>
<td>4.5</td>
<td>1536</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>29</td>
<td>22</td>
<td>1</td>
<td>7.5</td>
<td>1.5</td>
<td>6</td>
<td>440</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>30</td>
<td>14</td>
<td>3</td>
<td>12</td>
<td>3</td>
<td>6.5</td>
<td>440</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>32</td>
<td>28</td>
<td>5</td>
<td>20</td>
<td>2</td>
<td>24</td>
<td>52.5</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>33</td>
<td>10.5</td>
<td>0</td>
<td>1</td>
<td>30</td>
<td>24</td>
<td>840</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>36</td>
<td>18</td>
<td>1.5</td>
<td>8</td>
<td>2</td>
<td>24</td>
<td>1575</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>40</td>
<td>32</td>
<td>8</td>
<td>18</td>
<td>1</td>
<td>72</td>
<td>67.5</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>44</td>
<td>28</td>
<td>5</td>
<td>24</td>
<td>1</td>
<td>10</td>
<td>45.5</td>
</tr>
<tr>
<td>VA</td>
<td>F</td>
<td>51</td>
<td>25</td>
<td>2</td>
<td>6.5</td>
<td>3</td>
<td>48</td>
<td>169</td>
</tr>
<tr>
<td>VA</td>
<td>M</td>
<td>38</td>
<td>6</td>
<td>12</td>
<td>48</td>
<td>0.29</td>
<td>12</td>
<td>110</td>
</tr>
<tr>
<td>VA</td>
<td>M</td>
<td>44</td>
<td>12.5</td>
<td>0</td>
<td>50</td>
<td>16</td>
<td>24</td>
<td>910</td>
</tr>
</tbody>
</table>

| Mean | 34.68 | 18.43 | 3.82 | 23.14 | 4.30 | 33.36 | 346.72 |
| Stdev| 8.25  | 8.81  | 3.26 | 37.65 | 6.66 | 25.89 | 463.62 |
Figure 1. Psychophysical procedures. (A) Example high (100%) and low (20%) coherence motion stimuli. Signal dots are shown in white and noise dots in black. Directions of motion are indicated by the orientation of the arrow-heads. (Note: in the actual experiment all dots were white). Below each example stimulus is shown the corresponding distribution of signal values (solid black line) and noise values (dark grey shaded region). In the coherence task, noise was increased by changing the proportion of signal to noise dots. (B) Zero and high noise motion stimuli, with corresponding distributions of motion directions. In the equivalent noise task, noise was added by increasing the standard deviation of motion directions in the stimuli. In the plots of signal and noise distributions, the reference direction is denoted by a vertical black dotted line; the (average) direction of signal motion is circled. (C) The equivalent noise function (solid black line) is constrained by 2 data-points: the ‘zero noise’ threshold, which represents the minimum directional offset that can be reliably discriminated, and the ‘high noise’ threshold, which represents the maximum level of noise that can be tolerated for a large directional offset. The function has two parameters (inset in C), providing estimates of internal noise and global sampling (see Supplementary Material).
Equivalent noise procedure

A fast, efficient version of the EN paradigm, adapted for use with clinical populations, was used to assess local and global processing limits. In the EN tasks, participants judged whether a number of signal elements, presented for a brief duration were, on average, drifting clockwise or anti-clockwise of vertical-upward motion (motion task; Fig. 1B), tilted to the left or right of vertical (orientation task; Supplementary Fig. 2A), or smaller or larger than a reference (size task; Supplementary Fig. 2B). The reference direction, orientation and size were defined by the fixation guide itself, which was comprised of a small white circle bisected by a vertical line (identical in all tasks).

Two independent staircases were randomly interleaved: a ‘zero noise’ and a ‘high noise’ condition (Fig. 1C). In the zero noise condition, external noise was set to zero and the staircase tracked the minimum orientation offset from vertical (orientation task), directional offset from vertical (motion task) or size offset from reference (size task) that could be reliably classified (Supplementary Fig. 1B). In the high noise condition, the staircase tracked the maximum level of external noise that could be tolerated for a large (fixed) signal offset (Supplementary Fig. 1C). In this condition, the signal level was fixed at ±22.5° for the orientation, ±45° for the motion and ±0.5 octaves for the size task. These values were selected on the basis of previous studies and pilot data\textsuperscript{15,19,20}. Both staircases terminated after 75 trials each. As per the coherence task, the staircases were under the control of QUEST and converged on 82% correct thresholds. For each participant and task a two-parameter EN function was fit to their data, providing estimates of internal noise (a measure of local...
processing) and sampling (global processing). (See Fig. 1C and Supplementary Materials). To accustom participants to the nature of the task, all test blocks were preceded by 15 practice trials. In addition, for a subset of observers (10 participants with migraine and 8 without), 15 catch trials were randomly interleaved into each EN paradigm. On each catch trial the stimulus was presented at a large signal level in the absence of external noise (±22.5°, ±45° and ±0.5 octaves for orientation, motion and size tasks).

Stimulus parameters

All stimuli were generated in Matlab (MathWorks, Cambridge, MA) using the Psychophysics Toolbox extensions and were presented on a MacBook Pro laptop computer that was connected to a luminance-calibrated LCD monitor at a spatial and temporal resolution of 1920x1080 pixels and 60Hz, respectively.

Test images were generated by randomly dropping 100 elements (disks) within a circular region with a diameter of 15°. For motion and size judgements, individual elements could overlap. In the motion task, overlapping elements led to occlusion. In the size task, the contrasts of overlapping elements were summed. For the orientation task, element overlap was avoided by ensuring that adjacent elements were separated by a minimum distance equal to twice their diameter. The resulting images were presented in the centre of the screen for 400 milliseconds against a background grey display. Stimuli were viewed in a dark room from a distance of 51cm. The fixation guide had a diameter of 0.44°.
For the orientation task, individual disks were comprised of random phase sine-wave
gratings with a spatial frequency of 3.4 cycles per degree presented at 50% contrast in
a circular hard-edged mask with a diameter of 0.44° (Supplementary Fig. 1A). For the
size task individual disks had the same characteristics as for orientation, but varied in
the size and were randomly oriented (Supplementary Fig. 1B). The spatial frequency of
the grating was scaled to the diameter of the disk such that the number of cycles
presented remained constant across changes in size. In addition, for the size task, the
contrast of individual disks was randomly jittered in the range of 25-75% (sampled
from a flat distribution) in order to minimise the availability of contrast cues. For the
motion tasks, white dots with a diameter of 0.44° were used instead of windowed
gratings (Fig. 1B). Individual dots had a lifetime of 300ms, were spatially updated
every 50ms, moved at 3°/sec and were presented at 50% contrast.

Data transformation and filtration

All variables, with the exception of age and age of migraine onset, were log
transformed as this typically reduced skew and kurtosis. Following this
transformation, the distribution of variables did not differ significantly from normal
(ps > 0.05; one-sample Kolmogorov-Smirnoff tests). Data were then filtered
(separately for CON, MO and VA groups) so that extreme outliers with respect to
parameter estimates and associated confidence intervals (>2.58 Z-scores from the
group mean) were excluded from analysis. This led to the exclusion of 5.42% of the
data, which represented outliers that were seemingly randomly distributed across the
different groups [migraine (1.75%); control (3.67%)], tasks [motion coherence
None of the variables of interest differed significantly between migraine sub-groups (MO and VA) (independent t-tests, \(p_s>0.05\)); consequently, MO and VA data were pooled for all subsequent analyses. The percentage of catch trials answered correctly was at ceiling, and did not differ between groups or across tasks (ANOVA, \(p_s>0.05\)).

**Motion coherence thresholds**

To determine whether performance on the motion coherence task differed between migraine and control groups (Fig. 2A), coherence thresholds were analysed using an independent t-test (Table 2). A one-tailed test was employed since there are multiple reports of elevated coherence thresholds in migraine (see Introduction). Motion coherence thresholds were elevated in the migraine group (32\(\pm\)3.3\%) relative to the control group (24\(\pm\)1.8\%) \((t_{37}=-2.37, p=0.01, \text{Cohen’s } d=0.78)\), requiring a higher proportion of signal to noise dots to reliably classify the direction of signal motion.
**Figure 2. Coherence and equivalent noise plots.** Group mean (A) coherence thresholds, (B) levels of internal noise and (C) sampling are shown for control and migraine participants. Scatter-plots show correlations between motion coherence thresholds and (D) motion internal noise and (E) motion sampling. Error bars denote the standard error of the mean. Deg. = degrees. Note: data have been log-transformed; however, for ease of interpretation, axis tick-marks denote equivalent untransformed values.
Table 2. Comparing group performance on motion coherence and equivalent noise tasks.

Migraine and control group performance were compared using independent t-tests. Appropriate corrections were made to the degrees of freedom (d.f.) where equal variances could not be assumed. \(P\) values reported are for two-tailed tests, with the exception of the analysis of motion coherence thresholds, for which a single-tailed test was used (corrected alpha=0.1) (see text for further details). Bonferroni corrections were made for three multiple comparisons in the analysis of equivalent noise measures, reflecting the three different visual dimensions tested (corrected alpha=0.0167). \(t=t\)-statistic; d.f.=degrees of freedom; \(p=\)significance level; Cohen’s d=effect size; Th=motion coherence threshold; \(\sigma_{\text{int}}=\)internal noise; \(n_{\text{samp}}=\)sampling. *significant effect at the stated alpha level.

<table>
<thead>
<tr>
<th></th>
<th>(t)</th>
<th>d.f.</th>
<th>(p)</th>
<th>Cohen’s d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coherence T(h)</td>
<td>-2.37</td>
<td>37</td>
<td>*0.01</td>
<td>0.78</td>
</tr>
<tr>
<td>Motion (\sigma_{\text{int}})</td>
<td>-2.33</td>
<td>33.02</td>
<td>0.03</td>
<td>0.71</td>
</tr>
<tr>
<td></td>
<td>(n_{\text{samp}})</td>
<td>-0.04</td>
<td>41</td>
<td>0.97</td>
</tr>
<tr>
<td>Orientation (\sigma_{\text{int}})</td>
<td>1.21</td>
<td>41</td>
<td>0.23</td>
<td>0.38</td>
</tr>
<tr>
<td></td>
<td>(n_{\text{samp}})</td>
<td>1.82</td>
<td>32.56</td>
<td>0.08</td>
</tr>
<tr>
<td>Size (\sigma_{\text{int}})</td>
<td>0.22</td>
<td>39</td>
<td>0.83</td>
<td>0.07</td>
</tr>
<tr>
<td></td>
<td>(n_{\text{samp}})</td>
<td>-0.67</td>
<td>38</td>
<td>0.51</td>
</tr>
</tbody>
</table>

Table 3. Predicting motion coherence thresholds. A regression analysis showing the prediction of motion coherence thresholds from variance in three predictor variables [motion internal noise, motion sampling and group (migraine or control)]. All variables were added to the model simultaneously (i.e. non-hierarchically). Beta=beta coefficient; Beta\(_{st}=\)standardized beta coefficient; \(t=t\)-statistic; \(p=\)significance level; \(\sigma_{\text{int}}=\)internal noise; \(n_{\text{samp}}=\)sampling. *predicts a significant proportion of unique variance in the outcome variable.

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Beta</th>
<th>Beta(_{st})</th>
<th>(t)</th>
<th>(p)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Motion (\sigma_{\text{int}})</td>
<td>0.15</td>
<td>0.17</td>
<td>1.34</td>
<td>0.19</td>
</tr>
<tr>
<td>Motion (n_{\text{samp}})</td>
<td>-0.36</td>
<td>-0.63</td>
<td>-5.40</td>
<td>*5.2x10(^{-6})</td>
</tr>
<tr>
<td>Group</td>
<td>0.1</td>
<td>0.28</td>
<td>2.20</td>
<td>*0.03</td>
</tr>
</tbody>
</table>
Internal noise and sampling

To determine whether there was a general trend for group differences in internal noise, a multivariate analyses of variance (MANOVA) was undertaken with one between-participants factor (group at 2 levels: migraine and control) and three dependent variables (orientation, motion and size internal noise) (Fig. 2B). This revealed no main effect of group for internal noise (Wilks’ $\lambda=0.85$, $F_{(3,34)}=2$, $p=0.14$, partial-$\eta^2=0.15$). A similar analysis revealed no effect of group on sampling (Wilks’ $\lambda=0.86$, $F_{(3,33)}=1.83$, $p=0.16$, partial-$\eta^2=0.14$; Fig. 2C).

To determine whether group differences existed on a subset of EN tasks, levels of internal noise and sampling were exposed to a series of post hoc independent t-tests comparing migraine and control group performances (Table 2). Since analyses were undertaken for all visual dimensions tested (orientation, motion and size), Bonferroni corrections were made for three multiple comparisons (corrected alpha level=0.0167). The analyses revealed no significant differences in levels of internal noise or sampling between migraine and control groups for any of the EN tasks.
Predicting coherence thresholds from internal noise and sampling

To determine how motion coherence thresholds related to EN performance, bi-variate correlations were undertaken (Fig. 2D&E). Motion sampling was found to be highly negatively correlated with motion coherence thresholds ($R=-0.63$, $p=1.8 \times 10^{-5}$). Participants who were good at global pooling of information in the EN task needed fewer signals dots in the coherence task to correctly classify the direction of signal motion (Fig. 2E). In contrast, motion internal noise did not correlate with motion coherence thresholds ($R=0.22$, $p=0.18$; Fig. 2D).

Next, a regression analysis was undertaken. This tested the extent to which the three predictor variables [group (migraine or control), motion internal noise, motion sampling] predicted variance in motion coherence thresholds (the outcome variable) (Table 3). The resulting model was highly significant ($F(3,34)=13.3$, $p=7 \times 10^{-6}$) and accounted for 54% of the variance in coherence thresholds ($R=0.74$). Both group (6.6%) and motion sampling (39.44%) variables were found to predict a significant proportion of unique variance in coherence thresholds, whereas internal noise did not (2.4%). These findings indicate that even when differences in levels of internal noise and sampling were factored out, group membership (migraine vs. control) accounted for a significant proportion of variance in coherence thresholds.

Finally, none of the psychophysical measures recorded (coherence thresholds, internal noise or sampling) correlated with migraine characteristics (Supplementary Table 1). However, we note that the migraine characteristics included were based on self-report (e.g. migraine frequency, duration and severity), and hence, were highly subjective.
and prone to recall bias. Nor do they capture the fact that the nature of participants’
migraines may have changed with time.

5. Discussion:

In support of previous findings, motion coherence thresholds were elevated in the
migraine group relative to the control group. However, this difference could not be
attributed to deficits in either local or global processing. EN analysis generated
statistically indistinguishable estimates of internal noise (local processing) and
sampling (global processing) for migraine and control groups across all three
judgements types (orientation, motion and size). Further, regression analysis indicated
that group membership (migraine or control) predicted a significant proportion of the
variance in coherence thresholds, even once levels of internal noise and sampling
were controlled for. As discussed below, these findings are consistent with a relative
inability to exclude visual noise in migraine.

The finding of elevated motion coherence thresholds in the migraine group is
consistent with a number of previous reports. Whilst basic judgements of local
position\textsuperscript{14} and motion\textsuperscript{11} do not differ between migraine and control groups, repeated
studies have shown impaired performance on global form and global motion
coherence tasks in which participants must detect global structure embedded in noise\textsuperscript{6,9,10,12-14}. However, it has been argued that so-called ‘global’ coherence paradigms of
this kind do not rely exclusively on global integration processes; instead, performance
may also be limited by local processing, i.e. internal noise\textsuperscript{15}, or the ability to exclude
external noise. Consequently, EN analysis was undertaken so that independent estimates of local and global processing limits could be obtained.

The EN analysis undertaken here showed that levels of internal noise did not differ between migraine and control groups across any of the dimensions tested (orientation, motion or size). This is consistent with a number of previous studies. For example, a technique known as the N-pass method, which measures the consistency in a participant’s responses to sequential presentations of identical signal plus noise stimuli, has been used to estimate levels of internal noise in migraine. The principle underlying the technique is that internal noise reflects the level of random firing in a cell population that is sensitive to the dimension of interest, e.g. the direction of motion. As a result, a participant that is characterised by high internal noise will show poor consistency in responses across sequential presentations, since intrinsic variability in cellular responses, which is independent of the stimulus, will limit performance and drive random responses. Studies using this technique have shown that for global motion and two out of three global form tasks tested, levels of internal noise in participants with migraine are indistinguishable from those of control participants.

The EN analyses undertaken here also indicated normal global integration in migraine: levels of sampling were indistinguishable from control participants’ for judgements of orientation, motion and size. Although EN analysis has been applied to the study of migraine previously, it has not been used to characterise visuospatial performance; instead, previous studies have incorporated judgements of visual contrast. Thus, the findings are not directly comparable to our own: contrast EN
analysis is different from spatial and motion versions of the task, most pertinently, with respect to the nature of the external noise added to the stimulus. Consequently, performance is captured by a more complex model that includes additional free parameters including a multiplicative noise term. Nonetheless, two independent studies using contrast EN analysis have reported indistinguishable levels of sampling in participants with and without migraine. Further, they showed that levels of additive internal noise (equivalent to the local noise parameter in the EN model used here) also did not differ between groups. This suggests that the findings we report (i.e. normal local and global processing in migraine) may extend to other (non-spatial) visual dimensions.

Taken together with previous studies, the data reported here can be reconciled with a simple model of visual processing in migraine that posits normal local and global processing, coupled with a low tolerance to external noise. Thus, performance is seemingly unaffected on tasks that only require integration of the signal (e.g. spatial and motion EN tasks), but is impaired on judgements that first require segregation of the signal from noise (e.g. form and motion coherence tasks). It is noteworthy that a selective deficit in the mechanisms of external noise exclusion has previously been demonstrated in another clinical group characterised by visuo-cortical dysfunction. Thus, in amblyopia, performance is reportedly normal on EN tasks that involve judgements of global form and motion, but impaired on related form coherence and motion coherence tasks. Although speculative, the similarity in the pattern of these findings in migraine and amblyopia, coupled with their widely differing aetiologies, raises the possibility that the mechanisms involved in external
noise exclusion are particularly vulnerable following cortical damage or cortical
reorganisation.

A number of cortical models of migraine have already been suggested in the
literature. The majority of these are based on the notion of abnormal levels of cortical
excitation⁴,¹¹, i.e. *hypo*-excitability (reduced neural activity), or more commonly,
*hyper*-excitability (elevated neural activity) relative to healthy controls (see review⁵).
Thus, strengthened excitatory connections⁴²,⁴³, impaired mechanisms of inhibition⁴⁴,
⁴⁵ and abnormal pre-activation levels⁴⁶ have all been posited in migraine. However,
these models are often poorly specified, such that precise behavioural predictions
cannot be made on their basis. For example, hyper-excitability could imply elevated
levels of stimulus-driven (i.e. spiking) activity, a specific elevation in base-line firing
rates, or else a *generalised* increase in activity, all of which would lead to different
predicted effects on the signal-to-noise ratio, and hence, visual psychophysical
performance⁵.

With respect to the current study, the data reported are clearly inconsistent with
versions of both the *hyper*- and *hypo*-excitability models that posit an abnormal level
of base-line firing rates, since these would predict an elevation or reduction
(respectively) in internal noise. Instead, we report normal levels of internal noise in
migraine across all three visual dimensions tested (coupled with a selective elevation
in motion coherence thresholds). An alternative version of the *hyper*-excitability
model, which *is* broadly consistent with these data, is one in which stimulus-driven
(spiking) activity is elevated, whilst base-line firing-rates are unaffected. Let us
assume that a predominant direction of motion is selected by the observer once a
threshold firing-rate is exceeded within a population of appropriately-tuned direction-
sensitive neurones: if a single direction of motion is presented, hyper-excitability will
increase the likelihood that activity associated with the target direction will reach
threshold, and hence be reported. However, for a noisy (e.g. motion coherence)
stimulus, a state of hyper-excitability will also increase the probability that activity
driven by the noise will reach threshold, and hence compete with representations of
the signal.

Consistent with this model of (stimulus-driven) cortical hyper-excitability, Antal et
al.\textsuperscript{9} demonstrated superior motion discrimination performance in migraine (relative to
controls) for a stimulus comprised of a single direction of motion (100% coherence),
coupled with impoverished (relative) performance once the coherence of the stimulus
was decreased (i.e. noise was increased). In an earlier study, Antal et al.\textsuperscript{47} showed that
a similar dissociation could also be induced in healthy control participants: following
an experimental reduction in the excitability of cortical area MT, the discrimination of
intermediate coherence motion was enhanced, whilst the discrimination of 100%
coherent motion was impaired. Although we did not find superior classification
performance in migraine for a stimulus comprised of a single direction of motion
(remember that these trials were interleaved with a high noise staircase in the EN task,
potentially making the task harder), we did find a selective impairment in the
processing of a noisy (motion coherence) stimulus. Taken together, these data suggest
that a dissociation in the processing of motion coherence stimuli and stimuli
comprised of a single direction of motion (as reported) may be a signature of cortical
(stimulus-driven) hyper-excitability.
In conclusion, the findings reported here are inconsistent with local or global processing deficits in migraine, but instead, implicate impaired mechanisms of visual noise exclusion. This hypothesis has the potential to integrate a wide range of findings from the existing literature and open up novel avenues for investigation. Specifically, it predicts that relative to control participants, people with migraine will be impaired on any visual discrimination or detection task for which signal and external noise must be segregated prior to an integration stage, provided that sufficient external noise is added to the stimulus. Future studies should focus on the mechanisms involved in visual noise exclusion, since little is known about this process. One possibility that has been raised is that impaired noise exclusion reflects a state of (stimulus-driven) cortical hyper-excitability, which increases competition between representations of the signal and the noise. An alternative possibility, which is equally speculative however, is that representations of the noise compete with the signal to a greater extent in migraine because of a failure in endogenous attentional control, i.e. an inability to selectively monitor channels of interest that are most likely to carry the signal\textsuperscript{48,49}. To begin to tease these possibilities apart, it is clear that sophisticated psychophysical techniques must be employed in conjunction with clearly specified models of cortical function, so that highly specific predictions can be tested. We believe that the efficient version of the EN paradigm, which can be adapted to test across multiple sensory dimensions \textit{and modalities}, represents an invaluable tool in this approach.

**Funding:** This work was supported by The Wellcome Trust and by the NIHR Biomedical Research Centre at Moorfields Eye Hospital and UCL Institute of Ophthalmology.
6. References:


26. Webster KE. Investigating internal noise in migraine: a possible mechanism underlying perceptual deficits University of Western Australia. Crawley: University of Western Australia; 2011.


Supplementary Material

Equivalent noise analysis

The standard equivalent noise (EN) function is of the form:

\[
\text{EN} = \sigma_{\text{obs}} + \sigma_{\text{int}} + \sigma_{\text{ext}} + n_{\text{samp}}
\]  

where (for motion) \(\sigma_{\text{obs}}\) is the participant’s offset threshold (i.e. the smallest directional offset from vertical that can be reliably classified), \(\sigma_{\text{int}}\) is the participant’s additive internal noise, \(\sigma_{\text{ext}}\) the external noise in the stimulus, and \(n_{\text{samp}}\) the effective number of samples that the participant pools to determine the average direction of motion.

The traditional method of EN analysis constrains (1) by measuring offset thresholds at multiple levels of external noise, typically 6 or more, thereby requiring several thousand trials. However, the novel, rapid method use here, provides reliable estimates of internal noise and sampling in fewer than 100 trials. This rapid EN approach constrains the EN function with just two data-points / staircases (Fig. 1C). The first (‘zero noise’ condition) involves a manipulation of the signal direction across trials in the absence of noise, such that a basic offset threshold is estimated; this constrains the fit along the ordinate axis. The second (‘high noise’) condition relies on an inverse manipulation: the mean of the signal is fixed at a high level whilst
the level of external noise is manipulated across trials, such that the maximum level of 
noise that can be tolerated for a given performance level is estimated. This constrains 
the fit of the model in the orthogonal dimension (along the abscissa), and avoids 
sampling uninformative regions of the curve.

Correction for stimulus wrapping

For circular dimensions, i.e. orientation and motion, the stimulus wraps (at π for 
orientation and 2π for motion). Thus, an orientation of 0° is the same as an orientation 
of 180°, whilst a direction of 0° is equivalent to one of 360°. Consequently, the 
standard deviation of a distribution that is sampled to generate noise underestimates 
the actual variance presented at high noise levels, such that the equivalent noise model 
predicts lower thresholds in this area of the curve than are actually recorded\textsuperscript{15}. To 
overcome this issue we ran Monte Carlo simulations of a model observer’s 
performance across a range of internal noise and sampling levels. These indicated that 
an observer’s sampling level (n\textsubscript{samp}) is a function of their high noise threshold [i.e. the 
maximum level of noise that can be tolerated (MTN)] and can be captured by the 
following equation:

\[
n\text{samp} = \exp(AMTN^2 + BMTN + C) \tag{2}
\]

where best fits are obtained with values for A, B and C of 0.0001, 0.0329 and -1.903 
for motion, 0.0006, 0.0681 and -1.95 for orientation, and -0.4228, 2.797 and -1.241 
for size judgements, respectively. Note that these values are specific to a defined 
threshold performance level (82% here). This simple association between MTN and
sampling holds true because at high levels of external noise the effect of internal noise is negligible.

Once an estimate of sampling has been derived from the MTN, internal noise can be calculated from the ‘zero noise’ threshold. Thus, when $\sigma_{\text{ext}}=0$, by re-arranging equation (1):

$$\sigma_{\text{int}}^2 = \sigma_{\text{obs, samp}}^2$$ (3)
**Supplementary Figure 1.** Example staircases. Example staircases are shown for one participant’s data for the (A) motion coherence task, (B) motion equivalent noise task (zero noise condition) and (C) motion equivalent noise task (high noise condition). Under the control of QUEST, the stimulus level was set (on each trial) to the most probable Bayesian estimate of the underlying threshold - in this case, the 82% correct threshold.
Supplementary Figure 2. Orientation and size equivalent noise tasks. Example stimuli are shown for (A) orientation and (B) size equivalent noise tasks with zero noise conditions and high noise conditions on the left and right, respectively. Underneath each is shown the corresponding distribution of directions or sizes present in the stimulus. The reference orientation / size is denoted by a vertical black dotted line; the average signal orientation / size is circled.
Supplementary Table 1. Correlations between psychophysical measures and migraine characteristics. Pearson’s correlation coefficients (R) and associated significance levels (p) are reported. Th=motion coherence threshold; σ_int=internal noise; n_samp=sampling.

<table>
<thead>
<tr>
<th>Measure</th>
<th>Th</th>
<th>Age</th>
<th>Onset</th>
<th>Freq1</th>
<th>Freq2</th>
<th>Last</th>
<th>Duration</th>
<th>Severity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coherence</td>
<td></td>
<td>R</td>
<td>0.36</td>
<td>-0.19</td>
<td>-0.17</td>
<td>-0.02</td>
<td>-0.13</td>
<td>-0.31</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.74</td>
<td>0.11</td>
<td>0.45</td>
<td>0.46</td>
<td>0.93</td>
<td>0.57</td>
<td>0.18</td>
</tr>
<tr>
<td>Orientation</td>
<td></td>
<td>R</td>
<td>-0.19</td>
<td>0.37</td>
<td>-0.11</td>
<td>-0.14</td>
<td>0.17</td>
<td>-0.09</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.23</td>
<td>0.09</td>
<td>0.63</td>
<td>0.54</td>
<td>0.45</td>
<td>0.67</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>n_samp</td>
<td>R</td>
<td>-0.07</td>
<td>-0.04</td>
<td>-0.28</td>
<td>-0.14</td>
<td>0.13</td>
<td>-0.06</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.66</td>
<td>0.86</td>
<td>0.25</td>
<td>0.56</td>
<td>0.56</td>
<td>0.78</td>
<td>0.96</td>
</tr>
<tr>
<td>Motion</td>
<td></td>
<td>R</td>
<td>0.08</td>
<td>0.00</td>
<td>0.12</td>
<td>0.15</td>
<td>-0.08</td>
<td>0.33</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.63</td>
<td>0.99</td>
<td>0.60</td>
<td>0.49</td>
<td>0.71</td>
<td>0.13</td>
<td>0.29</td>
</tr>
<tr>
<td></td>
<td>n_samp</td>
<td>R</td>
<td>0.09</td>
<td>-0.30</td>
<td>0.28</td>
<td>0.21</td>
<td>0.04</td>
<td>-0.12</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.57</td>
<td>0.18</td>
<td>0.24</td>
<td>0.35</td>
<td>0.87</td>
<td>0.58</td>
<td>0.15</td>
</tr>
<tr>
<td>Size</td>
<td></td>
<td>R</td>
<td>-0.17</td>
<td>-0.06</td>
<td>-0.17</td>
<td>-0.20</td>
<td>0.37</td>
<td>-0.37</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.30</td>
<td>0.79</td>
<td>0.49</td>
<td>0.38</td>
<td>0.38</td>
<td>0.09</td>
<td>0.10</td>
</tr>
<tr>
<td></td>
<td>n_samp</td>
<td>R</td>
<td>-0.14</td>
<td>-0.03</td>
<td>-0.05</td>
<td>-0.11</td>
<td>0.22</td>
<td>-0.18</td>
</tr>
<tr>
<td></td>
<td>p</td>
<td>0.40</td>
<td>0.91</td>
<td>0.84</td>
<td>0.65</td>
<td>0.36</td>
<td>0.46</td>
<td>0.84</td>
</tr>
</tbody>
</table>