Assessment of epilepsy using noninvasive visual psychophysics tests of surround suppression

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Abstract
Powerful endogenous inhibitory mechanisms are thought to restrict the spread of epileptic discharges in cortical networks. Similar inhibitory mechanisms also influence physiological processing. We reasoned, therefore, that useful information about the quality of inhibitory restraint in individuals with epilepsy may be gleaned from psychophysical assays of these physiological processes. We derived a psychophysical measure of cortical inhibition, the motion surround suppression index (SSI), in 54 patients with epilepsy and 146 control subjects. Multivariate regression analyses showed that SSI was predicted strongly by age and seizure type, but not by seizure frequency. Specifically, we found that patients with exclusively focal epilepsy, and no history of generalization, showed significantly stronger cortical inhibition as measured by the SSI compared to all other groups, including controls. In contrast, patients with focal seizures evolving into generalized seizures, and patients with generalized genetic epilepsy, showed similar levels of cortical inhibition to controls. The presumptive focus, when one could be identified, was rarely found in visual cortex, meaning that the relationship with the epilepsy subtype is likely to reflect some global difference in inhibition in these subjects. This is the first reported instance of raised SSI in any patient cohort, and appears to differentiate between patients with respect to the likelihood of their experiencing generalization of their seizures. These results suggest that such simple psychophysical assays may provide useful aids to clinical management, particularly at the time of diagnosis.

Introduction
Most epileptic seizures are thought to arise from impaired interactions between excitatory and inhibitory elements in the cerebral and hippocampal cortices. A key role appears to be played by an endogenous inhibitory restraint mechanism arising from the particular arrangement of inhibitory drives onto pyramidal cells, and which serves to oppose the spread of epileptic activity (Prince and Wilder 1967; Trevelyan et al. 2006; Trevelyan and Schevon 2013). The inhibitory effects provided by the cortical interneurons (Atallah et al. 2012; Wilson et al. 2012; Pouille et al. 2013) are apparent in recordings of primary visual cortex neurons during various forms of visual suppression (Sengpiel et al. 1998). The same inhibitory networks are also believed to underlie various perceptual phenomena, collectively known as psychophysical surround suppression (Tadin et al. 2003, 2006a; Betts et al. 2009; Golomb et al. 2009; Serrano-Pedraza et al. 2014; Tadin 2015; Yazdani et al. 2015), although this does not discount contributions from other non-GABAergic mechanisms (Tadin 2015). One such test is based on the paradoxical finding that one’s ability to perceive the direction of movement of a high-contrast sinusoidal grating is reduced, as the stimulus size is increased (Tadin et al. 2003). This is believed to arise from surround suppression in the motion visual area, MT (Tadin et al. 2003, 2006a; Tadin 2015). This psychophysical phenomenon can be represented as a single number, the surround suppression index (SSI), derived from the ratio of the duration...
thresholds of the large and small stimuli. Intriguingly, the SSI decreases with age (Betts et al. 2005; Yazdani et al. 2015), and is also significantly reduced in subjects with schizophrenia (Tadin et al. 2006a; Serrano-Pedraza et al. 2014) and depression (Golomb et al. 2009); in each case, this has been proposed to reflect deficits in cortical GABAergic inhibition. We therefore investigated whether changes in SSI are also found in people with epilepsy.

We hypothesized that patients with epilepsy would also show alterations in visual psychophysical performance, and that this may be a useful clinical indicator of seizure risk. We investigated whether the SSI correlated with clinical features such as seizure frequency and seizure type, in order to determine what, if any, prognostic value might be provided by this simple psychophysics assay. Specifically, we had hypothesized that people with epilepsy might show evidence of reduced cortical inhibition, but surprisingly, our data suggest otherwise. Contrary to our original hypothesis, we found that people with generalized epilepsy showed no difference in SSI from control groups; and patients with focal epilepsy that did not generalize, on the other hand, showed a higher SSI. This is the first identified clinical group to show higher values of this measure. We suggest that these patients have enhanced cortical inhibition, which may be a factor in their seizures being restrained to subregions of the cortex. We discuss possible clinical implications of these results.

Materials and Methods

Experimental procedures were approved by Newcastle and North Tyneside Research Ethics Committee (reference number 09/H0906/90). Participants gave written informed consent, and were paid a nominal fee for their participation. Fifty-four patients with epilepsy (mean age, 41.9 years; age range = 17.0–82.3 years; mean duration 16.8 years; duration range = 0–50 years; 30 male) were recruited via specialist tertiary epilepsy clinics in Newcastle upon Tyne, UK. Seizure types and presumed etiology were classified according to the recent ILAE guidelines (Berg et al. 2010). Seizure frequency was estimated from patient diaries or hospital records. Due to the well-recognized inaccuracies of patient self-reporting of seizures (Hoppe et al. 2007), we subdivided seizure frequency into five bins: <1/year, <1/month, <1/week, <1/day, and >1/day. A total of 146 control subjects (mean age, 36.6 years; age range = 17.3–69.1 years; 59 male) were recruited via the Newcastle University volunteer cohort. Patients filled out a questionnaire regarding concurrent health issues and current medication, and also completed an Addenbrooke’s Cognitive Examination (ACE).

All subjects performed a motion discrimination task as described previously (Tadin et al. 2003; Yazdani et al. 2015). Briefly, drifting sinusoidal grating patches were presented at two different contrasts, high (peak contrast 92%) and low (2.8%), either on a desktop computer (Dell) with a CRT monitor, or a tablet computer (Samsung 700T), both running custom written Matlab software, implemented using Psychtoolbox3 (Kleiner et al. 2007). The discrimination was a simple two-choice paradigm, with the gratings moving either left or right. The stimulus presentation duration was either shortened or lengthened depending on whether the previous response was correct or incorrect, resulting in a staircase which settled close to the duration threshold. Three staircases were run in parallel, with trials interleaved at random. The SSI was defined as the log ratio of the duration thresholds of the large and small stimuli.

There was no apparent difference in the SSI measured on the two different systems (13 subjects [9 patients, 4 controls]; SSI\textsubscript{system 1} = 0.56 [range −0.16 to 1.56]; SSI\textsubscript{system 2} = 0.57 [range: −0.26 to 1.23]), so all the data were pooled. Grating patches were either small (subtending 0.7° on the retina when the tablet was held at 50 cm [or 100 cm when using the desktop system], users were instructed to hold the tablet at about this distance) or large (5°), and moved either left or right at constant horizontal velocity of 2°/sec. We were able to train most patients to do these tests very easily, meaning that a data set could be attained within 10–15 min. Three patients were unable to do the test and were excluded from the analyses. The duration of stimulus presentation was varied according to an adaptive staircase, whereby correct answers led to shorter presentations of the gratings, while incorrect answers caused the stimulus duration to be increased. Three staircase runs were interleaved randomly, so that the consequences of a correct or incorrect answer were hidden from the subject during the test. Such staircases rapidly tend toward presentations close to the threshold duration at which subjects could reliably identify the direction of movement. The actual value for the threshold duration, defined as the value where performance reached 82% (following Tadin et al. 2003), was estimated by fitting a psychometric function (Watson and Pelli 1983) to all trial durations plotted against the binary answer (right = 1; wrong = 0). A bootstrap resampling technique was used to derive 95% confidence intervals for the fitted thresholds, as described previously (Read et al. 2015). The SSI, as defined by Tadin et al. (2003), is the log ratio of the threshold durations (TD) for the large and small, high-contrast stimuli, calculated as follows:

$$SSI = \log_{10}(TD_{\text{high large}}/TD_{\text{high small}})$$
Table 1. Patient data.

| Index | Gender | Age (years) | Age at onset (years) | Duration of epilepsy (years) | Presumed location | Seizure frequency | Antiepileptic drugs | SSI |
|-------|--------|-------------|----------------------|-----------------------------|------------------|-------------------|---------------------|-----|
| EP1   | M      | 49          | Not known            | Not known                   | Temporal         | 3                 | VPA/PHT/CLB/PGB     | 0.09|
| EP5   | M      | 35          | 34                   | 1                           | Frontal          | 2                 | CBZ                 | 0.28|
| EP6   | F      | 26          | 25                   | 1                           | Temporal         | 2                 | VAL/LTG/PGB/CLB     | 0.34|
| EP13  | F      | 61          | 34                   | Unknown                      | Temporal         | 3                 | LTG                 | 0.10|
| EP14  | F      | 27          | 26                   | 1                           | Temporal         | 1                 | None                | 0.40|
| EP18  | F      | 55          | 41                   | 14                          | Temporal         | 2                 | None                | 0.19|
| EP19  | M      | 33          | 3                    | 30                          | Possible frontal | 4                 | VPA/CBZ/PHT         | −0.08|
| EP24  | F      | 22          | 7                    | 15                          | Occipital (L)    | 3                 | TPM/ZN             | 0.19|
| Focal+|        |             |                      |                             |                  |                   |                     |     |
| EP27  | M      | 68          | 58                   | 10                          | Unknown          | 1                 | LTG                 | 0.05|
| EP30  | F      | 58          | 11                   | 47                          | Unknown          | 3                 | LEV/PER             | 0.21|
| EP31  | F      | 57          | 28                   | 29                          | Temporal         | 4                 | LTG/PGB             | 0.24|
| EP33  | M      | 82          | 51                   | 31                          | Temporal         | 3                 | LTG                 | −0.07|
| EP36  | M      | 59          | 47                   | 12                          | Parietal         | 2                 | PHT/LTG/LEV/MDZ     | 0.10|
| EP49  | F      | 33          | 4                    | 29                          | Temporal (L)     | 4                 | OXC                 | 0.30|
| EP51  | M      | 30          | 17                   | 13                          | Unknown          | 2                 | LTG/TPM             | 0.75|
| EP53  | M      | 33          | 18                   | 15                          | Frontal          | 3                 | VPA/LEV             | 0.65|
| EP56  | M      | 68          | 64                   | 4                           | Temporal         | 2                 | LEV                 | −0.12|
| EP57  | M      | 44          | 9                    | 35                          | Right hemisphere | 3                 | ZNS/LEV/VPA         | 0.79|
| Focal−|        |             |                      |                             |                  |                   |                     |     |
| EP3   | M      | 68          | 12                   | 56                          | Temporal         | 1                 | CBZ/LEV/LTG         | 0.21|
| EP4   | M      | 70          | 64                   | 6                           | Temporal         | 4                 | LTG                 | 0.11|
| EP7   | F      | 67          | 6                    | 61                          | Temporal         | 3                 | PHT/LTG/LEV         | −0.11|
| EP17  | M      | 42          | 7                    | 35                          | Temporal         | 3                 | CBZ/LEV             | 0.42|
| EP20  | F      | 28          | 11                   | 17                          | Frontal (L)      | 5                 | RTG/CLB/LEVCBZ/LEV  | 0.87|
| EP21  | M      | 52          | 21                   | 31                          | Frontotemporal (L)| 3                | CBZ/LEV             | 0.72|
| EP23  | F      | 62          | 13                   | 49                          | Temporal         | 4                 | CBZ/ZN             | 0.71|
| EP25  | F      | 43          | 7                    | 36                          | Temporal         | 2                 | ZNS                 | 1.01|
| EP26  | F      | 73          | 54                   | 19                          | Unknown          | 1                 | VPA                 | 0.19|
| EP32  | M      | 56          | 40                   | 16                          | Temporal         | 4                 | LEV/RIG             | 0.66|
| EP34  | F      | 34          | 21                   | 13                          | Temporal         | 4                 | PER/LEV/PGB         | 1.24|
| EP35  | M      | 22          | 16                   | 6                           | Temporal         | 3                 | CBZ/TPM/CLB         | 1.34|
| EP37  | F      | 27          | 23                   | 4                           | Temporal         | 4                 | PER                 | 1.00|
| EP38  | F      | 25          | 0                    | 25                          | Multifocal       | 4                 | LEV/CLB/CLB/LEV     | 0.88|
| EP39  | M      | 34          | 31                   | 3                           | Temporal         | 5                 | TPM/LEVCBZ/LEV      | 0.69|
| EP41  | M      | 26          | 16                   | 10                          | Temporal         | 5                 | CLB/LCM/LEV/ZN      | 0.52|
| EP42  | F      | 31          | 6                    | 25                          | Temporal         | 3                 | LTG / PGB           | 1.02|
| EP44  | M      | 42          | 14                   | 28                          | Frontotemporal   | 4                 | VPA/CLB/LGT         | 0.50|
| EP45  | M      | 35          | 11                   | 24                          | Frontal          | 4                 | VPA/PGB/ESL/PB      | 0.48|
| EP46  | F      | 31          | 5                    | 26                          | Temporal lobe    | 4                 | PGB/LEV/CBZ/PHT/CLB | 0.65|
| EP47  | M      | 50          | 45                   | 5                           | Anterior temporal (L)| 4              | ZNS                 | 0.87|
| EP48  | F      | 62          | 46                   | 16                          | Temporal         | 4                 | CBZ/LEV/VPA         | 0.03|
| EP50  | F      | 26          | 22                   | 4                           | Temporal         | 2                 | None                | 0.79|
| EP55  | M      | 51          | 36                   | 15                          | Temporal         | 3                 | LEV                 | 0.57|
| GGE   |        |             |                      |                             |                  |                   |                     |     |
| EP8   | F      | 41          | 5                    | 36                          | Generalized      | 5                 | None                | 0.48|
| EP9   | M      | 18          | 12                   | 6                           | Generalized      | 1                 | VPA                 | 0.43|
| EP10  | M      | 17          | 16                   | 1                           | Generalized      | 2                 | VPA                 | 0.31|
| EP11  | M      | 55          | 54                   | 1                           | Unknown          | 2                 | VPA                 | 0.42|
| EP12  | M      | 18          | 17                   | Only 1 seizure              | Occipital        | 1                 | None                | 0.22|
| EP15  | M      | 22          | 22                   | 0                           | Generalized      | 2                 | None                | 0.36|
| EP16  | M      | 18          | 17                   | 1                           | Generalized      | 1                 | None                | 0.29|
| EP22  | F      | 29          | 20                   | 9                           | Generalized      | 3                 | LEV                 | 0.82|
| EP28  | M      | 51          | 5                    | 46                          | Possible frontal | 3                 | VPA/LEV/CBZ         | 0.27|
| EP29  | F      | 23          | 11                   | 12                          | Generalized      | 4                 | ZNS                 | 1.19|
| EP40  | F      | 22          | 22                   | 0                           | Generalized      | 5                 | LEV                 | 0.28|
| EP43  | F      | 55          | 15                   | 40                          | Generalized      | 2                 | PRM/PEGB            | −0.26|
Although the results for low-contrast stimuli are not incorporated directly into this index, they provide an important control assay of whether the participants were doing the psychophysics test correctly and prove that the high-contrast stimulus was well above their contrast threshold. Since the threshold duration for a large high-contrast stimulus is typically longer than for a small stimulus (Tadin et al. 2003; Yazdani et al. 2015), the SSI tends to be positive, and increasingly positive values indicate stronger surround suppression.

All statistical analyses were performed using the Matlab statistical toolbox. Comparisons of two regressions were performed using ANCOVA (analysis of covariance, `aoctool` in Matlab). Multivariate linear regression used the `fitlm` tool in Matlab, treating the epilepsy subtypes as “categorical.” Model comparisons were made using the adjusted $R^2$ values, which takes into account the effect of adding predictors on $R^2$. Results were considered significant if $P < 0.05$.

**Results**

We present an analysis of the performance on a simple visual psychophysics test of 54 patients with a confirmed diagnosis of epilepsy, and 146 control subjects. Details of the individual patients are provided in Table 1. Results from a subset of the control group (36 of the 146) were published as part of a prior study (Yazdani et al. 2015); the extended dataset we show here confirm our previous reports that the SSI shows a highly significant negative correlation with age ($P < 0.001$; Fig. 1E [green diamonds]). The epilepsy cohort showed a similar, highly significant regression with age ($P < 0.001$), and furthermore, regression analysis showed that the epilepsy group...
was highly significantly different from the control group ($F_{1,196} = 7.15, P < 0.0001$), both with respect to the intercept ($P < 0.0001$) and the gradient ($P < 0.0001$) of the relationship with age. Analysis of the component tests (Fig. 1A–D) indicated that the epilepsy cohort differed only on the tests involving large high-contrast stimuli ($F_{1,196} = 10.08, P = 0.0012$), which importantly is the one in which surround inhibition is likely to be manifest (Barlow and Mollon 1982; Sengpiel et al. 1998) (other tests: small high contrast, $F_{1,196} = 1.33, P = 0.250$; small low contrast, $F_{1,102} = 0.02, P = 0.875$; large low contrast, $F_{1,104} = 3.51, P = 0.064$; all nonsignificant). These results suggest that grouped together, the epilepsy patients have a higher SSI, suggestive of enhanced cortical inhibitory mechanism, when compared with age-matched control subjects.

We next subgrouped the epilepsy cohort with respect to seizure type (Berg et al. 2010) and seizure frequency. The cohort was subclassified into three groups: those patients with focal epilepsy with a history of generalized seizures ($F^+, n = 19$), focal epilepsy without generalizing seizures ($F^-, n = 24$), and generalized genetic epilepsy (GGE, $n = 11$) (Fig. 2). There was no difference in performance on the Addenbrooke’s Cognitive Examination between the groups ($F^+, ACE = 90.5 \pm 6.2$ [mean $\pm$ SD], range: 72–96; $F^-, ACE = 88.5 \pm 6.3$, range: 73–99; GGE, $ACE = 92.0 \pm 4.1$, range: 85–100). Seizure frequency was binned into five groups (Fig. 3). Initial inspection of these plots suggested that, in addition to the effect of age, both seizure type and frequency might also influence the SSI.

We therefore examined the relative importance of these three potential predictors (age, seizure subtype, and seizure frequency) of SSI by performing multivariate regression analyses on progressively more complex models (Table 2).

We first considered the subclassification into seizure types, independent of the age and seizure frequency. The distribution of SSI values differed significantly between the four groups ($F^+, F^-, GGE$, and controls; ANOVA, $F_{3,196} = 11.66, P < 0.0001$; Fig. 2A); t-test analyses indicated that the $F^+$ group was the outlier. The previously noted regression with age was apparent for each subgroup individually (Fig. 2B), although this was only significant for the two larger sample groups, $F^+$ ($n = 19, R^2 = 0.259, P < 0.05$) and $F^-$ ($n = 24, R^2 = 0.527, P < 0.001$), but not for GGE ($n = 11, R^2 = 0.144, n.s.$). Next, when considering both age and epilepsy diagnosis together, we found marked increases in the adjusted $R^2$ values when first subdividing the complete dataset (age alone, $R^2 = 0.105$, Table 2) into controls and epilepsy subjects (adjusted $R^2 = 0.183$), and then further subclassifying into the $F^+$, $F^-$, and GGE subtypes (adjusted $R^2 = 0.318$). Importantly though, the age and subtype model was not further improved by adding the seizure frequency (adjusted $R^2 = 0.315$). This lack of effect of seizure frequency was better appreciated when this predictor was plotted for the three seizure subtypes individually (Fig. 3B–D). These plots also show that in our samples, the $F^+$ patients tended toward a higher seizure frequency (the median frequency bin for GGE was “<1 month,” for $F^+$ it was “<1/

**Figure 2.** Surround suppression is altered in patients with focal nongeneralizing seizures, but is not affected by seizure frequency. (A) Box plot of SSIs for the subjects grouped by seizure type. The box limits represent the first/third quartiles, with the median indicated by the middle line and the whiskers extending to data points that are <1.5 interquartile range beyond the box. The data for the group with focal seizures without generalization (red) were highly significantly different from all other groups (**$P < 0.01$). (B) Regression of SSI with respect to age for the same groups of subjects. SSI, surround suppression index.
Figure 3. Surround suppression is not affected by seizure frequency. (A) Box plots of the SSIs with respect to frequency of seizures for the pooled epilepsy cohorts, and for each of the three subgroups of epilepsy patients, plotted separately (B–D, all nonsignificant). SSI, surround suppression index.
The most commonly prescribed drugs were levetiracetam (19 patients), lamotrigine (14 patients), and sodium valproate (13 patients), but notably the pattern of drug prescriptions for the patients with generalized seizures (GGE and F+) and those without (F−) were broadly similar (Fig. 4B). Since the psychophysics test is presumed to reflect cortical GABAergic function, we subdivided the epilepsy cohort into two groups according to whether or not they were on drugs that are known to interact with GABA (Table 3; note that both groups contain people on polypharmacy). Notably, there was no difference in the SSI for these two groups (non-GABA drug group, n = 27, SSI = 0.40 ± 0.37; GABA group, n = 27, SSI = 0.49 ± 0.36). Furthermore, including the presence or absence of drugs with GABAergic effects as a predictor in the regression analyses did not explain any additional variance (adjusted $R^2 = 0.316$). This was also true when the regression analyses were restricted to the epilepsy subjects (age/epilepsy subtype, adjusted $R^2 = 0.475$; age/epilepsy subtype/GABA effect, adjusted $R^2 = 0.464$). Finally, we examined whether patients with low versus high SSI scores (subdivided at the median SSI) were predominantly within the GABAergic/non-GABAergic drug interactions groups (Fig. 4C). There was no significant difference between the low and high SSI patients (Fisher’s exact tests), either for all the patients pooled irrespective of seizure type or for the generalized and focal groups alone. We concluded, therefore, that drug interactions do not underlie the effects of seizure type and age on the SSI.

**Discussion**

**SSI may provide biomarkers of epilepsy**

The results of most general interest are the relationship between SSI with respect to the likelihood of seizure generalization, and the lack of a relationship with seizure frequency, especially since these are counter to what might have been anticipated. Our original hypothesis had been that people with epilepsy would have a reduced SSI, indicative of lowered inhibitory restraint. Instead, we found that as a group, patients with generalized seizures are no different from control subjects, but those with focal epilepsy that does not generalize (F−), have a raised SSI. This surprising finding contrasts with the reduced SSI in other groups: people with schizophrenia (Tadin et al. 2006a), depression (Golomb et al. 2009), low IQ (Melnick et al. 2013), and aged subjects (Betts et al. 2005; Yazdani et al. 2015). Notably, most of the previously noted associations with increased SSI are “good” factors (youth [Betts et al. 2005; Yazdani et al. 2015] and high IQ [Melnick et al. 2013]). The significantly raised SSI in the F− patient group, relative to the other epilepsy groups, could not be explained by differences in age or IQ (there was no difference in ACE scores between the epilepsy groups). And

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**Table 2. Model comparisons.**

| Models | $R^2$ | Adjusted $R^2$ |
|--------|-------|----------------|
| SSI versus Age | 0.105 | – |
| Age, epilepsy | 0.195 | 0.183 |
| Subtype, freq | 0.170 | 0.144 |
| Age, freq | 0.200 | 0.192 |
| Age, subtype | 0.342 | 0.318* |
| Age, subtype, freq | 0.342 | 0.315 |

Model parameters (age, subtype)

| Gradient (year$^{-1}$) | Intercept |
|-------------------------|-----------|
| Controls | −0.0041 ± 0.0011** | 0.484 ± 0.047** |
| F+ | −0.0071 ± 0.0031 | 0.566 ± 0.147 |
| F− | −0.0164 ± 0.0031** | 1.362 ± 0.141** |
| GGE | −0.0088 ± 0.0047 | 0.694 ± 0.167 |

Note that for the control group statistics, what is being tested is significant difference from zero, and for the other groups, it is the significant difference from the controls.

The optimal model is indicated by *, and the parameters for that model are indicated by **P < 0.001.
while we cannot fully discount a confounding effect of concurrent depression, this condition is not known to be differentially associated with the presence, or absence, of generalized seizures in patients with focal epilepsy.

There are parallels between our study and a previous study of patients with migraine, who also showed evidence of increased suppression in a closely related perceptual task measuring contrast perception (Battista et al. 2011). The intriguing possibility is that in these patients with focal (nongeneralizing) epilepsy, the pathological activity is kept focused by an enhanced inhibitory restraint. Furthermore, it may therefore be possible to assess the quality of this restraint in regions of the cortex far removed from the focal pathology, as we do here with an assay of visual cortical function that appears to have relevance to foci elsewhere in the cortex. This presents an interesting question concerning whether the enhanced surround inhibition is independent of the epilepsy, or has arisen in reaction to the pathology, which will be addressed in future studies requiring longitudinal, repeated testing of patients from the time of diagnosis.

A large body of evidence has linked suppression of motion perception to processing in the motion area of visual cortex (cortical area MT) (Tadin et al. 2006b, 2003), equivalent approximately to the border between Brodmann areas 19 and 37 (see Tadin 2015 for an extensive review of this literature, including discussion of the involvement also of other parts of the visual system). With one exception (Ep12), this cortical area was not considered to be the focus of pathology for any of our patients (Table 1), which begs the question then of why measuring inhibitory function in a specific location may...
be relevant to epilepsy with a focus in a different part of the brain. There are parallels here with previous studies showing how SSI correlates with the occurrence of other brain pathologies not necessarily linked to visual processing, including schizophrenia (Tadin et al. 2006a) and depression (Golomb et al. 2009). Another study also showed an increase in contrast suppression in patients with migraine (Battista et al. 2011). We speculate that the answer lies in how inhibition may be affected globally, for instance, arising during development, or reflecting certain brain states, or under the influence of neuromodulators. If this is so, then an assay of inhibitory function at a particular location may also reflect inhibition in other areas that are relevant to the pathological condition.

The seizure frequency data are also interesting, although it needs to be interpreted with some caution, because this can be very difficult to estimate accurately (Hoppe et al. 2007). For instance, ambulatory recordings have shown that there is under-reporting of many seizure events (Cook et al. 2013). With this caveat in mind, it is interesting to contrast the absence of any relationship between seizure frequency and SSI with that regarding the likelihood of seizure generalization: this difference suggests that seizure initiation and seizure generalization may occur through different mechanisms modulated by different factors.

For all groups, the association of SSI with age persisted, consistent with previous studies (Betts et al. 2005; Yazdani et al. 2015). It is noteworthy that the largest increases of SSI were found in young patients without a history of seizure generalization, and that this group showed a significantly steeper association. This may represent a progressive change in the risk of seizure generalization; undoubtedly some people in this group will at some stage in their life experience a generalized seizure, meaning that they would have moved epilepsy groups in our analysis. At an early age, then, these people might be considered “latent” with respect to seizure generalization. Furthermore, given the association between seizure generalization and sudden unexpected death in epilepsy (SUDEP), we speculate that having a relatively low SSI at the time of diagnosis, even without a history of seizure generalization, may be a poor prognostic indicator. Again, we will benefit from longitudinal studies of progression and variability in SSI in individuals with epilepsy.

Drug interactions were difficult to assess because the diverse drug regimes in our patient cohorts made it difficult to control for this variable. Since the SSI is considered to reflect cortical GABAergic interactions, we focused our attention on drugs that are known to modulate GABAergic activity. We performed several different analyses, showing that the different epilepsy cohorts had broadly similar pharma profiles, nor was there any apparent difference between patients with high and those with low SSIs. It remains a possibility that some drugs may interfere with performance on the test, but this is highly unlikely to explain the differences between the epilepsy groups.

### Usefulness for clinical practice

We have shown how a simple visual psychophysics test may provide a convenient and entirely noninvasive means of assessing the function of cortical networks in the clinical setting. These tests required only minimal training, and can provide a measure of SSI within 10 min. We adapted these to run on a tablet computer, thus providing a portable means of testing, which the patients could use either in the clinic, or in their own home. Importantly, the data collected on these tablet computers in the community and at clinics matched previous studies performed in laboratory conditions, in showing a progressive and highly significant decline in the SSI with increasing age.

The main clinical implication of our study relates to the association of SUDEP with generalized seizures. SUDEP affects approximately 1 in 1000 patients with epilepsy per year, and the single biggest risk factor is the presence of uncontrolled generalized tonic-clonic seizures, increasing the risk to 1 in 150 patients per year (Nashef et al. 2007; Duncan and Brodie 2011; Shorvon and Tomson 2011). Currently, there are no reliable biomarkers of SUDEP risk. Any biomarker that reliably predicts patients at risk of generalized seizures would therefore be hugely beneficial for risk stratification, counseling, and treatment strategy. To be useful, such a biomarker would ideally be present.
before the occurrence of a first generalized seizure. We suggest that the SSI may prove to be a promising candidate for such a biomarker: the raised SSI seen in patients who have never previously had a generalized seizure indicating a lower risk of SUDEP, whereas the normal SSI seen in patients with a history of generalized seizures indicating a higher risk. Since SSI also tends to decrease with age, this index will be most useful for patients who develop, or are diagnosed with epilepsy early in life.

Our groupings according to seizure types were based on seizures that had already occurred, and we studied patients at only a single time point. We therefore cannot know whether the patients with generalized seizures had a normal SSI initially, and nor can we know whether patients with an increased SSI will remain free of generalized seizures in the long term. It is noteworthy, however, that the decline of SSI with age is significantly more steep in the F− group than for other groups, which may mean that their risk of generalizing seizures, and therefore by extension, of SUDEP, may also change. These questions can only be addressed by further longitudinal studies. Nevertheless, given that visual psychophysical measures are simple, quick, and safe to administer, we feel that these further studies are justified.

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Conflict of Interest

None declared.

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