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Predictive processing of music in frontal and temporal cortices

Intracranial recordings and computational modelling of music reveal the time-course of prediction error signaling in frontal and temporal cortices.

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Running title: Predictive processing of music in frontal and temporal cortices
Abstract

Prediction is held to be a fundamental process underpinning perception, action and cognition. To examine the time-course of prediction error signaling, we recorded intracranial EEG activity from 9 pre-surgical epileptic patients while they listened to melodies whose information-theoretic predictability had been characterized using a computational model. We examined oscillatory activity in the superior temporal gyrus (STG), the middle temporal gyrus (MTG) and the pars orbitalis of the inferior frontal gyrus (IFG), lateral cortical areas previously implicated in auditory predictive processing. We also examined activity in anterior cingulate gyrus (ACG), the insula and amygdala, to determine whether signatures of prediction error signaling may also be observable in these subcortical areas. Our results demonstrate that the information content (a measure of unexpectedness) of musical notes modulates the amplitude of low-frequency oscillatory activity (theta to beta power) in bilateral STG and right MTG from within 100 and 200ms of note-onset respectively. Our results also show this cortical activity to be accompanied by low-frequency oscillatory modulation in ACG and insula - areas previously associated with mediating physiological arousal. Finally, we showed that modulation of low-frequency activity is followed by that of high-frequency (gamma) power from approximately 200ms in the STG, between 300ms and 400ms in the left insula and between 400 and 500ms in the ACG. We discuss these results with respect to models of neural processing that emphasize gamma activity as an index of prediction error signaling, and highlight the usefulness of musical stimuli in revealing the wide-reaching neural consequences of predictive processing.
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Introduction

In the field of music cognition, it is widely held that listeners’ expectations about how music will unfold provide a rich source of its aesthetic value and meaning (Huron, 2006; Meyer, 1956). According to this theory, predictive uncertainty and violations of expectation introduce tension, while events that confirm expectations convey resolution of tension. This theory of music processing is consistent with proposals that prediction is a fundamental process underpinning many aspects of perception and cognition (Clark, 2013; Friston, 2010), including decision making, motor action, visual perception and language comprehension (Gentsch, Weber, Synofzik, Vosgerau, & Schütz-Bosbach, 2016; Lewis, Schoffelen, Schriefers, & Bastiaansen, 2016; Rauss, Schwartz, & Pourtois, 2011; Summerfield & De Lange, 2014). For musical stimuli, the likelihood of different continuations, and therefore the strength of expectation violation, can be precisely specified using computational models. This is of great benefit in studying the neural mechanisms underlying expectations in music (Lehne & Koelsch, 2014; Omigie, 2015). However, it is also of great potential benefit in improving our understanding of neural mechanisms of predictive processing more generally.

Electrophysiological signatures of melodic and harmonic expectancy processing

Expectancy may be described as the anticipation of a future event based on its probability of occurrence. The concept of musical syntax (Pearce & Rohrmeier, 2018; Rohrmeier & Pearce, 2018) refers to the structural regularities of a musical style that govern these probabilities of occurrence (e.g. Koelsch & Siebel, 2005), whether in terms of melodic, rhythmic or harmonic structure (Large & Palmer, 2002). A learned, internal model of the syntactic structure of a musical style is thought to determine listeners’ musical expectations (Rohrmeier & Koelsch, 2012) and has been shown to influence not just auditory perception across expertise levels (Bharucha & Stoeckig, 1986; Margulis & Levine, 2006; Marmel, Tillmann, & Dowling, 2008) but also music production in expert musicians (e.g. Bianco et al., 2016).
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Over the decades, there has been a great deal of research on electroencephalographic (EEG) correlates of violations of melodic and harmonic structure. One early study, which compared differential event related potentials (ERPs) elicited by incongruous events in sentences to those in well-known melodies, revealed that incorrect musical events elicited an amplified N100, as well as a positivity around 300ms post note onset (Besson & Macar, 1987). Similar EEG signatures were found for terminal deviant notes in well-known melodies in a study incorporating a delay period to allow the build-up of expectancies (Paller, McCarthy & Wood, 1992). Since these pioneering works, the early negativity and the late positive potential (or P300) have been shown to be influenced by listeners’ musical expertise, the music’s familiarity, and the strength of the events’ incongruity (Besson & Faita, 1995; Brattico, Tervaniemi, Näätänen, & Peretz, 2006; Miranda & Ullmann, 2007; Omigie, Pearce, Williamson & Stewart, 2013; Peretz, Brattico, Järvenpää, & Tervaniemi, 2009).

Harmonically incongruent events have similarly been shown to elicit early anterior negativities, the amplitudes of which have also been shown to be sensitive to age and expertise, the degree of unexpectedness of the deviant, as well as the presence or absence of veridical knowledge regarding the occurrence of the deviant (e.g. Guo & Koelsch, 2016; Koelsch, Gunter, Friederici, Schröger, 2000; Koelsch, Schmidt, & Kansok, 2002).

Interestingly, however, while commonalities have been observed, previous work also reveals differences between melodic and harmonic expectancy processing. For instance, the late positivity seen for incongruous musical events has been shown to be more discriminating of harmonic than melodic stimuli (Hantz, Kreilick, Kananen & Swartz, 1997). Irregular melody endings have been shown to elicit an ERP effect peaking at 125ms after the onset of notes but not the later ERP signature observable for irregular final chords. Melodic and harmonic deviants also elicit slightly different scalp distributions of activity (Koelsch & Jentschke, 2010) and neuropsychological work points to differences in the neural substrates of melodic and harmonic syntax processing (Samson & Zatorre, 1988). Taken together, while the literature, as a whole, points to some common basis for
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harmonic and melodic structure processing, these results highlight the importance of studying each
domain of musical syntax alone in its own right. A previous study has used intracranial recording to
better localize processing of harmonic syntax in the temporal and frontal cortices (Sammler et al.,
2013) but intracranial recording has not been used with melodic stimuli.

**Oscillatory and anatomical correlates of musical expectancy**

Accompanying the characterization of ERPs to incongruent harmonic and melodic events, there is
increasing interest in characterizing the patterns of oscillatory activity that are associated with such
events. It is noteworthy that the degree of violation of musical expectancy has long been associated
with power variations in various frequency bands, primarily in right frontal brain regions (Janata &
Petsche, 1993). More recently, however, variations in beta power in particular (Pearce, Ruiz, Kapasi,
Wiggins & Bhattacharya, 2010) and long-range synchronization across frontal and temporal regions
(Herrojo-Ruiz, Koelsch, & Bhattacharya, 2009) have also been reported as a function of expectancy
violations.

When considering predictive processing during music listening, it is relevant to examine not just the
electrophysiological signatures of expectation violation but also the anatomical substrates involved in
these processes. Neuropsychological studies have shown that patients who have undergone right
temporal excisions show deficits in melodic discrimination (Samson & Zatorre, 1988). These studies
emphasize the importance of Superior Temporal Gyrus (STG) in musical pitch processing (Zatorre &
Samson, 1991). However, neuroimaging studies that have associated computational models of tonal
processing with neural processing have further implicated rostral frontal areas in the processing of
musical key (Alluri, Toiviainen, Jääskeläinen, Glerean, Sams & Brattico, 2012; Janata, Birk, Van
Horn, Leman, Tillmann & Bharucha, 2002). A recent meta-analysis confirms that, although regions
of the STG are heavily implicated, frontal areas especially are essential for the processing of higher-
order tonality (Janata, 2015). Some previous computational approaches used to examine tonality
Predictive processing of music in frontal and temporal cortices processing are based on low level models of time-varying acoustic features (Alluri et al., 2012). Others have been able to show the anatomical substrates of tonality processing by modelling a higher-level concept of tonality. For example, Janata and colleagues (2002) used domain general unsupervised machine learning methods (self-organizing maps), trained on a specially-constructed melody that rotates between closely aligned keys, to associate low level pitch percepts with key strengths. The model used in the present research uses domain general information-theoretic principles of statistical learning and probabilistic prediction to learn the syntax of a musical style through exposure. Combined with a high spatial resolution methodology like intracranial EEG, this modelling approach makes it possible to elucidate the anatomical substrates of violations expectations in a precise way.

While studies specifically examining tonal expectancies have primarily implicated the inferior frontal gyrus (IFG) (Maess, Koelsch, Gunter & Friederici, 2001; Kim, Kim & Chung, 2011; Koelsch, Fritz, Schulze, Alsop, & Schlaug, 2005; Musso et al., 2015, Sammler, Koelsch, & Friederici, 2011; Tillmann, Koelsch, Escoffier, Bigand, Lalitte, Friederici., & von Cramon, 2006), it is important to note the heterogeneity of this sub-region of the frontal lobe (Amunts, Lenzen, Friederici, Schleicher, Morosan, Palomero-Gallagher & Zilles, 2010; Hartwigsen, Neef, Camilleri, Margulies & Eickhoff, 2018). Indeed, it is worth noting that while previous studies have tended to report sensitivity to musical syntax in Brodmann area (BA) 44 and BA45 (Bianco et al., 2016; Cheung, Meyer, Friederici & Koelsch, 2018; Koelsch & Siebel, 2005), others have rather associated the subjective experience induced by incongruous musical events (i.e. the tension associated with expectation violations) with BA47 and more orbital frontal areas (Lehne, Rohrmeier, & Koelsch, 2014; Mikutta et al., 2015). In one single case-study, using depth-electrode recordings, higher theta power was associated with harmonically unexpected chords in left orbitofrontal cortex (OFC) (Mikutta et al., 2015). Furthermore, using fMRI, BOLD increases in the pars orbitalis of the left IFG have been associated with music-induced subjective tension (Lehne et al., 2014).
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Finally, while early work on expectancy processing has tended to focus on lateral cortical areas, it is interesting to note that recent studies have begun to highlight an important role of subcortical structures. Re-analysing fMRI data using a region of interest (ROI) analysis, Koelsch, Fritz and Schlaug (2008) reported greater BOLD activation in both left and right amygdala during the processing of irregular chord endings. Since then, evidence has suggested the involvement of left amygdala theta power in the processing of harmonically unexpected chords (Mikutta et al., 2015, but note data from only one patient was evaluated), and right amygdala activity in mediating subjective tension in response to longer musical pieces (Lehne et al., 2014). Thus, taken together, a growing literature suggest that not only cortical but also subcortical areas may be involved in processing violations of melodic expectation. However, very few studies have characterized the neural signatures of expectation violation using methods, like intracranial EEG, that possess both fine temporal and spectral resolution. Further, despite it being necessary in order to conclusively demonstrate the neural signatures of predictive error signaling, few studies have parametrically manipulated the strength of expectation violation induced by the musical stimuli in a highly refined way.

Computational approaches to melodic expectation modelling

Studies that examine the where and when of neural oscillatory activity in response to expectation violations point to the potential fruitfulness of using musical expectancy to examine the predictive coding framework (Clark, 2013; Friston, 2010; Heilbron & Chait, 2018). Essentially, contemporary models, within this framework, suggest that predictive processing may involve the coordination of slow and fast brain oscillations. In particular, it has been suggested that whereas a lower (e.g. beta) frequency channel from deep cortical layers propagates top-down information, bottom-up information from superficial cortical layers is propagated using the gamma frequency channel (Wang, 2010). Similar patterns have been reported on a large-scale cortical level by Arnal, Wyart & Giraud (2011) who exploited the specificity with which visual input predicts auditory input in
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Just as for audiovisual speech, musical stimuli contain probabilistic regularities that an enculturated listener can be expected to have internalized. Accordingly, musical stimuli whose probabilistic structure has been characterized using computational modelling have the potential to elucidate the neural correlates of predictive processing (e.g. Omigie et al., 2013). The Information Dynamics of Music model (or IDyOM: Pearce, 2005; 2018) implements the theory that listeners weigh the probability of different possible continuations to a musical excerpt based on the frequency with which different continuations followed similar contexts in their previous experience (Meyer, 1956; Pearce, 2005). While many studies rely on music-theoretic approaches to characterize musical syntax, IDyOM embodies the hypothesis that listeners internalize the syntactic structure of musical styles through domain-general processes of implicit statistical learning based on long-term exposure to music (Bigand & Poulin-Charronnat, 2006; Tillmann, Bharucha, & Bigand, 2000) and generate probabilistic expectations based on this internalized syntax. Accordingly, models like IDyOM offer the opportunity to consider musical processing in a more domain-general way.

Support for the notion that statistical learning influences expectancy formation comes from listeners’ subjective rating of expectedness of specific events in a musical context. For instance, listeners rate small intervals as more expected than large ones, reflecting the relative frequency with which such intervals occur in melodies (Huron, 2006). When required to give subjective ratings of how well each of a set of notes fits a musical pattern (Cuddy & Badertscher, 1987), listeners also produce rating profiles that reflect the tonal hierarchy present in western music. IDyOM has proven very reliable in predicting listeners’ explicit unexpectedness and uncertainty ratings (Hansen, Vuust & Pearce, 2016; Pearce et al., 2010; Hansen & Pearce, 2014), as well as their reaction times in implicit
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tasks that probe listeners’ melodic expectations (Omigie, Pearce & Stewart, 2012). IDyOM has also
demonstrated high discriminative power. Specifically, it has been able to predict listeners’ style-
specific expectations (Hansen et al., 2016) and successfully simulate other music cognitive processes
such as recognition memory (Agres, Abdallah & Pearce, 2018), perceptual similarity (Pearce &
Müllensiefen, 2017) and metrical inference (Van der Weij, Pearce & Honing, 2017).

In allowing precise characterization of all events in a melody, computational models of melodic
expectation like IDyOM allow the careful interrogation of neural responses to events with respect to
their information content (IC: an information-theoretic measure of unpredictability) in the context of
a naturalistic listening condition. Indeed, an earlier study combining scalp EEG and IDyOM output
demonstrated parametric sensitivity to the IC of melodic events of an observed early negativity and
later positivity (Omigie et al., 2013). However, that scalp EEG study did not and could not address
some of the claims put forward by a predictive coding framework with respect to the anatomical
specificity of underlying oscillatory mechanisms. Similarly, while both statistical learning and
music-theoretic methods (Egermann, Pearce, Wiggins & McAdams, 2013; Koelsch, Kilches,
Steinbeis, & Schelinski, 2008; Steinbeis, Koelsch & Sloboda, 2006) have emphasized the ability of
music’s probabilistic pitch structure to influence listeners’ physiological arousal; there is still only
sparse evidence for the involvement of the very subcortical regions known to mediate such
autonomic responses. Here, we aimed to identify any consistencies existing between cortical and
subcortical responses to parametric manipulations of expectation violation that may confirm a role of
the latter in predictive processing of music. In other words, we aimed to examine the possibility that
the network of regions involved in predictive processing include subcortical as well as cortical areas.

The current study
Our study capitalized on the excellent temporal and spatial resolution of intracranial depth electrode
recordings (e.g. Omigie et al., 2015 a, b) and state of the art computational modelling of music to
Predictive processing of music in frontal and temporal cortices shed light on predictive processing of melodic structure, both in terms of patterns of oscillatory activity and the underlying anatomical regions. To take advantage of the detailed stimulus characterization afforded by computational modelling, we analyzed oscillatory activity in response to all notes in all the melodic stimuli presented to participants. Specifically, we examined detailed timing of activity in the STG, Middle Temporal Gyrus (MTG) and pars orbitalis of IFG— all held to be involved in auditory deviant and musical syntax processing—as well as the amygdala, Anterior Cingulate Gyrus (ACG) and insula, three key structures of the limbic network previously implicated in error, syntax and salience processing as well as, most importantly, in mediating physiological arousal (Critchley, Mathias & Dolan, 2001; Lappe, Steinsträter, & Pantev, 2013; Sander, Grafman, & Zalla, 2003).

Melodic stimuli used in the current study contained between 32 and 64 notes, each with a duration of 600ms and separated from the next note by 100ms. To constrain the scope of our investigation, we use the notion of a prediction unit cycle beginning with prediction-related activity reflecting the likelihood of different note continuations and continuing with prediction error signals following the onset of a given note. Here it is important to note that we focus on the latter half of this cycle: namely the time-course of prediction error signaling following note onset, which is nevertheless dependent on the existence of predictions regarding what that note would be. We suggest that the feedback-related activity or prediction error signaling following a given note’s onset reflects the extent to which predictions mismatch with the actual incoming sensory information, and may be expected to occur from as early as within 300 ms of the occurrence of the given incoming note (Durschmid et al., 2016). Critically, we used IDyOM estimates of IC as our measure of the extent to which listeners’ predictions are violated, and as the variable with which to predict neural consequences of these violations. In Figure 1, we provide a schematic view of the oscillatory activity we expected to see in different regions of interest as a function of increasing IC during the prediction error signaling phase. Firstly, based on a range of EEG studies examining the neural time-course of
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Musical syntax processing in frontal and temporal cortices, we also anticipated early increases in low level frequency bands (theta-alpha power) as a function of increasing note unexpectedness (IC), and this not just in STG but also in syntax processing areas such as MTG and pars Orbitalis of the IFG (e.g., Sammler et al., 2011). Further, based on the results of studies examining novelty processing in auditory mismatch negativity paradigms in general (Haenschel, Baldeweg, Croft, Whittington, & Gruzelier, 2000; Todorovic, Schoffelen, van Ede, Maris, & de Lange, 2015), we anticipated a modulation of beta power by IC in the STG in particular. Finally, and most importantly, in line with the proposed role of higher frequency activity (gamma) in propagating prediction errors especially in low level sensory processing areas, we expected the strength of gamma activity to increase with increasing note unexpectedness, also specifically in the STG (Fuentemilla, Marco-Pallarés, Münte, & Grau, 2008; Mikutta et al., 2015). We anticipated that this prediction error-signaling gamma modulation would start at around 140 ms and peak at approximately 230 ms in line with previous studies using oddball paradigms (Durschmid et al., 2016; El Karoui et al., 2015). Thus, in summary, and based on the predictive coding and more general music and auditory deviance detection literature, we predicted a modulation of theta-alpha power in lateral frontal and temporal cortices more generally (STG, MTG and IFG), followed by a modulation of beta and gamma power in auditory cortices (STG) specifically.

Finally, a number of subcortical areas have been associated with salience detection and musical syntax processing, so we examined the possibility that these areas show graded responses to parametric manipulations of expectation violation that would confirm their role in predictive processing. First, in line with evidence of its involvement in the detection of salient events (Sander et al., 2003), the processing of unexpected musical chords (Koelsch et al., 2008a), as well as tension in real music (Lehne et al., 2014), we predicted we would observe sensitivity to IC in the amygdala (Mikutta et al., 2015). Further, we predicted that the insula, which has been associated with both musical syntax processing and the mediation of physiological arousal, would, along with the
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**Materials and Methods**

**Participants**

Data were collected from 12 patients implanted with depth intra-cerebral electrodes for pre-surgical evaluation at the Epilepsy unit in the Pitié-Salpêtrière Hospital. However, data from only 9 patients (M = 34.4 ± 12.65 years; 2 Males) were analyzed due to highly artifacted data in one patient and no electrodes in relevant regions in the other two (see Table 1). The study was approved by the ethical committee for Biomedical Research of Pitié-Salpêtrière Hospital in Paris (CPP Paris VI, INSERM C11-16). All patients gave informed written consent and clinical investigations were conducted according to the principles expressed in the Declaration of Helsinki.

**Stimuli**

Stimuli were the same as those used in previous behaviourial and scalp EEG studies (Omigie et al., 2012; Omigie et al., 2013) and were characterized using IDyOM (Pearce, 2005). The melodies varied
Predictive processing of music in frontal and temporal cortices in length from 32 to 64 notes and were in a range of major keys. Individual notes were created using the grand piano instrument of a Roland sound canvas MIDI synthesizer before being converted to audio files. In order to focus on pitch expectations in particular, the rhythmic structure of the melodies had been removed in a musically sensitive manner by a skilled musicologist so that each note had the same duration of 600ms and an equivalent inter-onset interval of 700ms (100ms between notes). Each note was rendered at the same sound intensity. Finally, in order to provide an incidental detection task for the patients to engage in, a single note in 6 out of the 56 melodies, was modified to play in a different timbre, specifically the marimba instrument.

**Information-theoretic characterization of stimuli**

The computational model IDyOM (Pearce, 2005, 2018) learns from a corpus of melodies and the current melody being processed in order to generate an Information Content (IC) value for each note in a given melody. The configuration of the model used here was exactly as described in previous studies whereby the model was trained on an independent corpus of western tonal melodies as well as the melody itself as it unfolded (Omigie et al., 2012; Omigie et al., 2013). The IC of a note is the negative log of the probability of the note’s occurrence and indicates the degree of unexpectedness of the given note in its given context (Pearce, 2005). We sought to verify whether a parametric relationship could be observed between note IC and oscillatory activity in particular ROIs encompassing lateral and medial frontal and temporal areas. Accordingly, all the notes of each melody were sorted by their IC, irrespective of which melody they belonged to, and assigned to ten bins of increasing IC by decile (following the main analysis in Omigie et al., 2013). Figure 2 shows the IC profile of a sample melody used in the experiment, the IC distribution for all notes in the 56 melodies and descriptive statistics for the IC of notes assigned to the 10 bins. Table 2 shows the mean tonal stability values (Krumhansl & Kessler, 1982), the size of the preceding interval and the mean position of the notes within the melody. In the western tonal system, the stability of a pitch within a key is related to its position in a hierarchy, and higher ranking/more stable pitches are often
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more frequent and more predictable than lower ranking ones (Krumhansl, 1990). Also, larger pitch
intervals between notes are less frequent than smaller ones in western melodies (Huron, 2001).
Accordingly, as can be seen in Table 2, the higher the mean IC of notes in a bin, the more tonally
unstable they tended to be, and the greater their tendency to follow larger intervals. Both mean tonal
stability per bin and mean interval size were shown to correlate strongly with mean IC per bin (IC
and tonal stability: \( r = -0.92, p<.001 \); IC and Interval Size, \( r = 0.9, p<.001 \)) confirming a contribution
of both to the model’s IC estimates.

**Task**

Participants were presented with the 56 melodies in 3 blocks. They were asked to listen to the
individual melodies and indicate after each whether they had heard a note played in a deviant timbre
using keys labelled YES or NO on the presentation laptop. Performance on this task was
implemented to keep participants’ attention high and notes from those melodies containing a deviant
timbre were not included in the analysis. With regard to performance, participants showed a high Hit
Rate (M=0.83, SD=0.12, range=0.67-1), although the False alarm (FA) rate showed a wider range
due to two participants who expressed great hesitance with the task and accordingly were encouraged
by the experimenter to respond based on their subjective feeling (M=0.14, SD=0.28; range=0-0.77).
When these two participants were excluded, a much smaller FA rate (M=0.04; SD=0.039), in line
with expectations, was obtained. Given the irrelevance of the task to the study, no participants were
excluded based on their performance.

**Intracranial EEG recordings**

Participating patients had been stereo-tactically implanted with depth-electrodes to localize the
epileptogenic focus for possible resection. Invasive exploration was planned based on brain locations
suspected to be at the origin of epileptic seizures. The hypotheses regarding localization resulted
Predictive processing of music in frontal and temporal cortices from analysis of data previously gathered in the clinic (examination and history, continuous surface EEG-video, MRI and, in some cases, PET and ictal SPECT). Local field potential data were acquired at a sampling-rate of 4Hz (ATLAS system, Neuralynx, Bozeman, MT, USA) and bandpass filtered between 0.05 and 1Hz. Electrodes were composed of 4–12 contacts, each 2.3 mm long, 5 to 10 mm apart, mounted on a 1.12 mm wide flexible plastic probe (Ad-Tech Medical Instruments, Racine, WI). These were stereotactically inserted using a Leksell frame (Adam et al., 1996). To determine localisation of contacts, the postoperative CT scans were coregistered with the preoperative 1.5 Tesla MRI. The EpiLoc toolbox (Stereotaxy: Techniques, Images, Models) from the Institut du Cerveau et de la Moelle Epinière was used to automatically establish the MNI coordinates and those coordinates which fell within the regions of the MNI template for each region of interest were adopted in a first step. In a second step, such contacts were examined on the postoperative MRI scans using Slicer (Fedorov et al., 2012) and, using further visual inspection, were either confirmed or not as belonging to the given ROIs. To allow conclusions to be drawn with anatomical specificity, only those contacts for which at least one adjacent contact was present in the (ROI) were considered for inclusion in the analysis, although this resulted in fewer contacts available for analysis.

**Data preprocessing and time-frequency analyses**

Contact and artifact rejection as well as data preprocessing were carried out using Fieldtrip (Oostenveld, Fries, Maris, & Schoffelen, 2011) and custom written MATLAB scripts. Data were epoched from 2000 ms before to 2000 ms after the onset of each note. Artifacts related to epileptic activity were automatically removed by excluding 1) trials whose maximum amplitude exceeded the mean amplitude of the trial by at least 5 standard deviations (SD) as well as 2) contacts in which more than 5% of trials were excluded (following the previous criterion). Trials were also visually checked for spikes and abnormal rhythmic activity. Signals were analyzed in a bipolar-montage to minimize the influence of distant sources (Bollimunta, Chen, Schroeder, & Ding, 2008), after subtraction of the signals of adjacent contacts. The Cartesian coordinates (x, y, z) corresponding to
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the new bipolar signals were calculated as the mid-distance location between two adjacent recording sites, where both sites were located in the region of interest. To examine oscillatory activity, data were high-pass filtered at 0.01 Hz and then bandpass filtered to allow the examination of 5 frequency bands of interest: Delta (1-3Hz), Theta (4 -7 Hz), Alpha (8-12Hz), Beta (12-30Hz) and Gamma (30-80 Hz). A Hilbert transform was then carried out on the bandpass filtered data to obtain the analytic signal and accordingly the oscillatory power time-course for each epoch. The time-course of all epochs was then baselined to the 100ms window directly preceding the onset of the sound by subtracting the mean power during this period from the entire epoch.

Statistical Analysis

Predictive processing was studied in 81 intracranial contacts across 9 epileptic patients. Activity was recorded from cortical (67 contacts) and subcortical structures (14 contacts). The distribution, number and location of contacts in each of the ROIs for all 9 participants are shown in Figure 3 and Table 3 respectively. In order to control for the variance associated with different contacts and patients, data were analyzed using linear mixed-effects models with the lme4 package in R (Bates, Maechler, Bolker, & Walker, 2013). Each trial, representing an individual note in an individual melody, was split into equal-sized time-windows (e.g. TW1:0-100ms, TW2:100-200ms, etc.) in order to determine the relative latency of significant modulations of neural activity by IC. Within these windows, mean spectral amplitude for each contact, for each patient, for each frequency band in a particular ROI was estimated. The amplitude of oscillatory power served as the dependent variable in the linear mixed-effect models (for each band, for each time window) with IC bin (IC) as fixed effect, and contacts and participants as random factors with intercept to account for any variation across individual contacts and participants. To boost statistical power (necessary given the limited number of contacts), we analyze contacts from the left and right hemispheres together initially (El Karoui et al., 2015). However, we examine the lobes separately in case of non-significant effects, in order to ensure that non-significant effects were not due to contacts showing
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contrasting effects. p values reported are obtained using the anova function from the car toolbox in R, and are FDR corrected for multiple comparisons (across time-windows and across frequency-bands) for each ROI separately. Finally, given the heterogeneous regions measured from, and to complement the previous analysis, further models examining modulation of activity in each contact individually were estimated. As using FDR to correct across time-windows, frequency bands and contacts resulted in no effects surviving this conservative approach, results using less conservative alpha values of p < 0.01 are presented.

Results

STG

Figure 4 shows the time-course of oscillatory power as a function of IC across all trials across the 21 contacts in bilateral STG (13 left and 8 right contacts) with grey-shading indicating the specific time-windows in which IC significantly predicted oscillatory power. Significant modulation was found from within 100ms and continued until 400ms after note onset in the theta band (TW1: β= 0.03, SE=0.005, p<0.001; TW2: β=0.05, SE=0.008, p<.001; TW3: β=0.05, SE=0.01, p<.001; TW4: β=0.03, SE=0.01 p=0.005), until 300ms after onset in the alpha band (TW1: β=0.03, SE=0.007, p<.001; TW2: β=0.07, SE=0.01, p<.001; TW3: β=0.05, SE=0.01, p<.001), and until 200ms after onset in the beta band (TW1: β=0.02, SE=0.009, p<.05; TW2: β=0.04, SE=0.009, p<.001). A medium latency modulation was found in the gamma band between 200ms and 400ms (TW3: β=0.01, SE=0.003, p<0.001; TW4: β=0.01, SE=0.003, p<0.001) while, finally, modulation by IC was found between 500 and 600ms after onset in both the alpha (TW6: β=0.03, SE=0.01, p<0.05) and in the gamma band (TW6: β=0.008, SE=0.003, p<.05).

Figure 5 shows the results of the linear regression modelling on the single contact level. Here, 50% (4 out of the 8) of contacts in the right temporal lobe and 31% (4 out of 13) of contacts in the left temporal lobe showed modulation at a significance level of p<.01. The previously observed pattern
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of lower frequency activity being followed by higher frequency activity was corroborated in these analyses as low frequency activity (theta to beta) for individual contacts was observed from within 100ms to 300ms (and then again at 500-600ms), while effects in the gamma band were only present from 200ms to 400ms. Finally, it is interesting to note that while both anterior and posterior STG contacts showed modulation in the theta and beta bands, only posterior STG contacts showed modulation in the alpha and gamma band.

**MTG**

Thirty-three contacts in the left (25 contacts) and right (8 contacts) MTG were examined. Linear mixed model analysis with all contacts in the MTG showed some evidence for modulation of activity in lower frequency bands, followed by modulation in gamma bands but these modulations did not survive correction for multiple comparisons. Running the analysis on the right temporal lobe contacts alone revealed a significant modulation in beta power between 100 and 200ms (Beta: TW2: $\beta=0.04$, SE=0.01, p=0.05) and a trend in the gamma band ($\beta=-0.007$, SE=0.002, p=0.08) whereby gamma power was reduced for higher IC notes. No significant modulation of activity in the left temporal lobe was observed. Finally, 37.5% (3 out of a possible 8) of contacts in the right and 24% (6 out of a possible 25) of contacts in the left showed modulation either within the first 200ms in lower frequency bands (predominantly in delta but also in theta and alpha bands- see Figure 5) or at later time windows in alpha and gamma (300ms to 600ms) whereby power in the alpha and gamma bands were reduced for high IC events. Visual inspection suggests that contacts showing modulation across the different frequency bands were distributed across both anterior and posterior MTG.

**IFG pars orbitalis.**

Linear mixed-effect model analysis of 13 contacts in both lobes (2 left and 11 right) suggested modulation in alpha (0 to 100ms) and delta (200 to 500ms) power but neither this grand analysis, nor analysis of the hemispheres separately, survived correction for multiple comparisons, although, as in
Predictive processing of music in frontal and temporal cortices

STG and MTG, the distribution and timing of activity on a single electrode level (p<.01) showed gamma frequency activity being the longest latency. Specifically, alpha-beta activity within the first 100ms and delta between 300 and 500ms was later followed by beta-gamma band activity between 500 and 600ms. Modulation was observed in 50% (1 out of a possible 2) of left contacts and 18.2% (2 out of the possible 11) of contacts in the right lobe.

Anterior Cingulate Gyrus, Insula and Amygdala.

Figure 6 shows the time-course of spectral power in bilateral ACG, with grey-shading indicating the specific time windows at which IC significantly predicted amplitude of oscillatory power. Five ACG contacts (1 left, 4 right) were examined. The ACG showed significant modulation by IC in the theta band between 200 and 500ms (TW3: β=0.03, SE=0.01, p<.05; TW4: β=0.04, SE=0.01, p<.05; TW5: β=0.03, SE=0.01, p=0.05), in the alpha band between 100 and 300ms (TW2: β=0.03, SE=0.01, p<.05; TW3: β=0.03, SE=0.01, p<.05), and in the gamma band between 400 and 500ms (TW5: β=0.01, SE=0.004, p<.05). Single electrode analysis showed two contacts (one in the left lobe (100%) and one of a possible four in the right (25%)) in the ventral anterior cingulate region showing significant modulation in the theta band (0 to 300ms) and then in the alpha band (200 to 400ms) (see Figure 7).

At the single contact level, 5 contacts in the left (2) and right (3) insula and four contacts in the left amygdala were examined. While a grand linear mixed effect analysis of insula contacts did not show any effects that survived correction for multiple comparison (see Figure 6), analysis of contacts in the left insula alone revealed significant modulation in the delta band (TW1: β =0.03, SE=0.007, p<.001; TW2: β = 0.06, SE=0.014, p<.001; TW3: β =0.07, SE=0.019, p<.001; TW4: β =0.08, SE=0.02, p=.001; TW5: β =0.07, SE=0.02, p<.05) and gamma band (TW4: B=0.017, SE=0.006, p<.05) but no significant modulation in the right hemisphere. Single electrode analysis confirmed that two insula contacts (out of a possible 2 in the left lobe, 100%) showed modulation from 0 to
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400ms in delta band followed by modulation in gamma band from 300 to 400ms. Finally, analysis of amygdala contacts together did not survive correction for multiple comparisons although analysis of single contacts showed a single contact displaying significance between 100 and 200ms in the beta band (see Figure 7).

Discussion

The current study examined the extent to which the information-theoretic predictability of a musical note modulates the oscillatory activity in both a) lateral frontal and temporal areas involved in auditory sequence processing and b) amodal limbic areas involved in salience detection and physiological arousal. As we were particularly interested in the time-course of prediction error signaling in these regions, we used information content as a measure of expectation violation (or prediction error) to predict oscillatory activity. Contrary to our hypotheses, neither the observed modulation of IFG pars orbitalis nor the observed modulation of left amygdala activity survived correction for multiple comparisons across time-windows and frequency bands. However, we were able to provide robust evidence of parametric modulation of oscillatory activity by expectation violations in the STG, the right MTG, the ACG (primarily in ventral regions) and left insula. Presence and timing of the observed oscillatory activity, notably in the gamma frequency band, showed congruence with patterns found in predictive processing in other domains. Thus, our findings emphasized the utility of musical stimuli in revealing the wide-reaching neural consequences of predictive processing in general.

The role of the lateral temporal lobe in melodic expectancy and predictive processing.

We predicted that STG would show robust modulation by IC in lower (theta to beta) and higher (gamma) frequency bands, with the latter, in particular, reflecting auditory prediction error signaling (Arnal et al., 2011). Accordingly, our results showed a broadband increase in bilateral STG oscillatory activity as a function of increasing IC that was most extensive in the theta (0 to 400ms)
Predictive processing of music in frontal and temporal cortices and most short-lived in the beta band (0 to 200ms). Further, and most critically, our results in the STG confirmed robust gamma modulation by IC, which compared to lower frequency bands, had the longest latency: emerging at approximately 200 ms and then again at 500 ms post note onset.

On the one hand, the pattern of oscillatory activity observed here is easily reconcilable with a large number of studies that link theta-alpha and beta-gamma activity to auditory change detection and music syntax processing more specifically (e.g. Choi, Lee, Ko, Lee, Jung & Kim, 2013; El Karouï et al., 2015; Fuentemilla et al., 2008; Ko et al., 2012). Greater theta, alpha and beta power has previously been reported for higher IC musical events (Carrus, Pearce, & Bhattacharya, 2013; Pearce et al., 2010) while beta power in the STG, in general, is often associated with novelty processing in auditory mismatch negativity paradigms (Haenschel, Baldeweg, Croft, Whittington, & Gruzelier, 2000; Todorovic, Schoffelen, van Ede, Maris, & de Lange, 2015). On the other hand, the findings are also particularly important in corroborating a pattern of activity that has often been described in association with the predictive coding framework (Arnal et al., 2011). Specifically, here we show IC modulation of activity in the gamma band, which has been argued to propagate prediction errors following the comparison of predictions with the incoming sensory events. That this gamma activity modulation is limited to STG in the temporal lobe, and specifically to posterior STG contacts, is consistent with the notion that such bottom-up information originates from lower-level sensory processing areas. Further, while its latency may seem to be longer than in previous studies (e.g. Durschmid et al., 2016) it is important to note that our use of 100ms time windows limits the precision with which the onset of this gamma sensitivity can be reported and, indeed, it can be seen that the sensitivity does seem to emerge as early as half way into the preceding time-window (100 to 200ms). Finally, that beta activity, which has been argued to both carry top-down predictions (Arnal et al., 2011) as well as index expectation violations (Todorovic et al., 2015) also showed robust modulation by IC here, emphasizes this frequency band as playing a critical role in the predictive processing in the brain.
Predictive processing of music in frontal and temporal cortices

While STG houses the primary auditory areas and may thus be expected to play the most critical role, an extensive literature, documenting studies carried out with patients and healthy participants, demonstrates a more general role of the right lateral temporal lobe in melodic pitch processing (e.g. Johnsrude, Penhune & Zatorre, 2000; Patterson, Uppenkamp, Johnsrude, & Griffiths, 2002; Warren & Griffiths, 2003; Warrier & Zatorre; 2004). In addition to the IFG and the STG, the MTG has previously been implicated in music syntactic (Sammler et al., 2013) and syntax processing more generally (Brennan, Nir, Hasson, Malach, Heeger, & Pylkkänen, 2012; Kaan, & Swaab, 2002). However, the timing and nature of oscillatory activity in response to predictive processing in this region remained unclear. Here, we anticipated we would observe lower (theta-alpha) modulation in MTG as a function of IC. Interestingly, we observed that parametric modulation of oscillatory activity in this region was primarily in the right hemisphere, and particularly in the beta range. Here it important to note that while previous studies have shown beta power to be modulated by musical note IC (Carrus et al., 2013), ours is the first study to show that such beta oscillations may emanate not just from the bilateral STG but also from the right MTG. In other words, we show for the first time a specific involvement of right MTG beta power in the processing of melodic violations.

In addition to these beta band effects in the right MTG, we also observed a trend towards reduced MTG gamma power (between 300 and 600ms) for high IC events that was corroborated by the single contact analyses. Whilst it is necessary to be cautious with these findings, it is interesting to note that they resonate with that from a scalp EEG study that also reported decreases in gamma power in response to irregular musical chords (Herrojo-Ruiz et al., 2009). Those authors interpreted their results within the framework of the match-and-utilization model (MUM) and under the assumption that tonic chords constitute a match. However, the limited spatial resolution afforded by that scalp EEG study would have masked the MTG as the source of this reduced gamma signal. Critically, it would also have masked the presence and source of an effect that we see here and which is more in line with the predictive coding framework – namely an increase in gamma activity reflecting
Predictive processing of music in frontal and temporal cortices
prediction error signaling. Thus, taken together, previous non-invasive EEG methods have
contributed greatly to our understanding of the neural correlates of predictive processing. However,
by providing here, electrophysiological evidence of differing patterns of oscillatory activity in close-
lying lateral temporal areas (the STG and MTG), our study highlights the utility of intracranial EEG,
combined with computational modelling of music, in this field of inquiry.

The role of ventral ACG and insula in melodic expectancy and predictive processing.
In addition to aiming to throw light on the timing of predictive error signaling in lateral cortical
areas, a key aim of the current research was to examine whether parallels may be seen across cortical
and subcortical areas. Our motivation to investigate this emerged from the observation that areas like
the insula and ACG, are not only involved with salience detection, attention orientation and musical
syntax processing, but also in the mediation of physiological arousal. Here we provide valuable
confirmation of a role in music predictive processing, of key subcortical structures that are difficult
to isolate using non-invasive EEG (Deouell, 2007). Specifically, in addition to showing modulation
of STG and right MTG oscillatory activity by IC, our results also showed systematic IC modulation
in ventral regions of the ACG and insula, whereby increasingly high IC events resulted in a sustained
increase of oscillatory activity.

Modulation of ventral ACG oscillatory activity by IC was observed first in the theta (within 200 ms
of note onset) and then in the alpha band. (within 100 ms). However, interestingly and mirroring
STG activity to a degree not fully anticipated, ACG modulation was also observable later (from 400
to 500ms) in the gamma frequency band. With regard to what the activity in this region may mediate
during the prediction error signaling process, one possibility is that it is related to the greater
attentional resources that are likely being allocated to increasingly high IC events (Bush, Luu &
Posner, 2000; Carter, Braver, Barch, Botvinick, Noll, & Cohen, 1998). However, a further
compelling explanation for this observed modulation is that it reflects the role of the ventral ACG in
Predictive processing of music in frontal and temporal cortices

the processing and integration of emotional information (Mayberg, 1997) and in mediating an
interplay between cognitive processing and sympathetic activity (Critchley et al., 2001, Critchley et
al., 2003; Beissner, Meissner, Bär, & Napadow, 2013). Initial evidence that high IC notes in a
melody can lead to modulation of the autonomic response was provided by Egermann and colleagues
(2013), who demonstrated that unpredictable passages characterized by high IC led to greater
physiological arousal. Electrical stimulation of both ventral and dorsal regions of the ACG have long
been shown to result in the modulation of heart rate and blood pressure in animals (Ward, 1948;
Chefer et al., 1997) whilst in more recent work on humans, sympathetic activity has also been
observed to be related to ACG activity (Critchley et al., 2001, Critchley et al., 2003; Beissner,
Meissner, Bär, & Napadow, 2013). Therefore, one possibility is that, despite using isochronous
melodies only, the observed ventral ACG modulation reflects a mechanism by which violations of
melodic expectation can induce changes in emotional arousal (Meyer, 1956). Such a potential role is
supported by our findings of a similar modulation (albeit in delta and gamma power) in the insula
which, in addition to being implicated in deviance-detection (Hsu, et al., 2014) and musical violation
processing (Bianco et al., 2016; Cheung et al., 2018; Lappe et al., 2013), has also been associated
with mediating physiological arousal (Singer, Critchley, & Preuschoff, 2009).

Finally, given our intention to examine consistencies across cortical and subcortical areas, it is
interesting to note that the patterns of low and high frequency activity we observed in ventral ACG
and insula were highly consistent in terms of time course (from low to high frequency bands) with
those found in the STG and MTG. No study to date has shown a sensitivity to the strength of violated
expectations of gamma activity in the ventral ACG or insula, and we argue that one of the valuable
contributions of this study is showing this similar pattern of neuro-oscillatory responding across
cortical and subcortical areas. Without being able to estimate connectivity between these cortical and
subcortical areas (e.g., as in Omigie et al., 2015), we, unfortunately, cannot comment on the extent to
which the effects in these different regions are related. However, we suggest that this constitutes a
Predictive processing of music in frontal and temporal cortices
very interesting avenue of research that could be fruitfully pursued in future studies using the
intracranial recordings combined with computational modelling approach we demonstrate here.

IFG pars Orbitalis and amygdala during music listening.

The IFG is widely held to be involved in predictive processing more generally and in music syntax
processing specifically. However, our failure to show a strong role of pars orbitalis IFG in melodic
expectation processing here may not be considered so surprising given that previous studies have
mainly reported responses to music-syntactic violations in BA44 and BA45 subregions of the IFG
(Bianco et al., 2016; Cheung et al., 2018; Koelsch & Siebel, 2005). Despite the fact that BA47 has
not been as widely implicated in predictive processing as BA 44 and 45, our expectation of
modulation in this region was primarily based on findings that tension resulting from complex
musical structure modulates the OFC, and particularly the pars orbitalis of the IFG. Bearing that in
mind, our failure to show robust oscillatory activity modulation in IFG goes against the results of
Lehne and colleagues (2014), who showed left pars orbitalis IFG, as well as against those of Mikutta
and colleagues who showed left OFC activity to be involved in tension processing (Mikutta et al.,
2015). It is important to note that we did observe patterns somewhat in line with those studies.
Specifically, we observed a greater proportion of left than right contacts showing modulation by IC
and single contacts in the pars orbitalis of the IFG showed modulation from within 100ms to 600m
after sound onset although these effects failed to survive correction for multiple comparisons.
Further, while a lack of power cannot be excluded as a reason for the absent effect here, it is worth
noting that the hierarchy of stability may be less robust for individual tones than for chords, and
therefore the neural correlates of deviant processing in the IFG may not be as pronounced for
melodies (as employed here) as for harmony (Koelsch, 2012).

Lastly, since the amygdala is considered part of a network that, along with the OFC, modulates
attention to stimuli (Vuilleumier, 2005), we hypothesized a role for this region in responding to
Predictive processing of music in frontal and temporal cortices

expectation-violating melodic events (e.g., Zarcone, Van Schijndel, Vogels, & Demberg, 2016). However, our results showed little evidence of modulation of amygdala power by IC. Once again, it is possible that melodic structure does not modulate this region to the extent that manipulations of harmonic structure are able to. After all, here, even the highest IC notes modulating neural activity are not deviant per se, compared to the highly irregular events used in other studies (Koelsch et al., 2008a). Other possibilities, however, are that the right amygdala, which was not recorded from in this study, is more involved than the left amygdala in processing musically salient events (Lehne et al., 2014), or that, quite simply, the current stimuli used lack the complexity and richness necessary for recruiting emotion and reward areas during the cognitive processing of syntax.

Closing remarks

In sum, our findings throw light on the time-course of neural activation involved in music prediction error signaling, extending previous work on melodic and harmonic processing that has relied on non-invasive EEG/MEG and neuroimaging. Critically, our results emphasize the usefulness of a computational approach that, by allowing the estimation of conditional probabilities for individual events, offers a fine-grained quantification of the predictability of musical events that is difficult to obtain using purely music-theoretic analysis. Our use of a model based on domain-general principles of information theory yields observations about brain activity that are potentially generalizable across domains (Strange, Duggins, Penny, Dolan, & Friston, 2005), and may thus be profitably extended to provide further domain-general understanding of predictive processing in the brain.

Nevertheless, the current study has a number of shortcomings which, while shared with most iEEG studies (Mukamel & Fried, 2012), deserve elaboration. Firstly, the findings rely on a small number of contacts from a restricted number of patients, thus limiting the generalizability of significant findings and throwing doubt on some null results, which may be type II errors. Secondly, due to little overlap in contact distribution across patients, the current analysis had to focus on the effect of IC on
Predictive processing of music in frontal and temporal cortices different regions in isolation and could not investigate how these regions interact, as has proven highly enlightening in previous work (Omigie et al., 2015a). Thirdly, to minimize participation effort, our paradigm did not require participants to give feedback on subjective feelings of surprise or emotional arousal, which limits our capacity to draw strong conclusions about affective processing of the stimuli. Future studies in which data is collected from a greater number of patients, each implanted in different key ROIs simultaneously, would allow valuable insight into interaction within the network of brain areas mediating the processing of musical expectancy. Most importantly, however, studies that combine a) objective quantification using computational models of the kind employed here with b) stimuli with the capacity to induce greater affective responses and reward (Alluri et al., 2012), will throw more light-than was possible here- on the role that predictive processing plays in the aesthetic and emotional experiencing of music.

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Neuroscience 6, 16–23.


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Table 1: Demographic and background information on participating patients

<table>
<thead>
<tr>
<th>Patient</th>
<th>Date of recording</th>
<th>Age</th>
<th>Sex</th>
<th>Handedness</th>
<th>Seizure frequency (per week)</th>
<th>Age of epilepsy onset</th>
<th>Epileptic focus</th>
<th>Lesion</th>
<th>Cerebral localization of Implantation</th>
<th>Areas examined</th>
<th>Other regions</th>
<th>Medication</th>
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<tbody>
<tr>
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<td>03/06/14</td>
<td>62</td>
<td>F</td>
<td>Left</td>
<td>3</td>
<td>32</td>
<td>Bilateral anterior hippocampus</td>
<td>Right hippocampal sclerosis</td>
<td>Right and left temporal lobes</td>
<td>rMTG, lMTG, rSTG, ISTG</td>
<td>BA38, Hipp</td>
<td>Lamictal, Lysanxia</td>
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<td>01/07/14</td>
<td>30</td>
<td>F</td>
<td>Right</td>
<td>7</td>
<td>16</td>
<td>Superior frontal lobe (BA 6, 8)</td>
<td>Dilation of the right temporal horn</td>
<td>Right frontal lobe</td>
<td>rACG</td>
<td>BA 6, 8, 10</td>
<td>Keppra, Lamictal</td>
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<tr>
<td>pat_4</td>
<td>26/01/15</td>
<td>32</td>
<td>M</td>
<td>Right</td>
<td>0.25</td>
<td>24</td>
<td>Left temporopolar cortex</td>
<td>No lesion</td>
<td>Left temporal lobe</td>
<td>lAMYG, IMTG, ISTG</td>
<td>BA 20, 34, 36, 37, 38, Hipp</td>
<td>Keppra, Lamictal, Tegretol</td>
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<td>3</td>
<td>8</td>
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<td>No lesion</td>
<td>Right frontal lobe</td>
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<td>Dépakine, Lamictal</td>
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<td>Left temporal lobes</td>
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<td>Lamictal, Zonégran</td>
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<td>Left frontal and temporal lobes</td>
<td>IMTG, ISTG</td>
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<td>Vimpat, Diphantoin</td>
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<td>F</td>
<td>Right</td>
<td>-</td>
<td>-</td>
<td>Left temporopolar cortex, middle temporal cortex, BA, 20, 38</td>
<td>Right hippocampal sclerosis</td>
<td>Right frontal and left temporal lobe</td>
<td>rSTG, rMTG, rIFG, rIns, lAMYG</td>
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<td>BA 20, 38, 46, 47</td>
<td>Lamictal</td>
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38
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<th>Patient</th>
<th>Date</th>
<th>Age</th>
<th>Gender</th>
<th>Side</th>
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<th>Bilateral</th>
<th>Language</th>
<th>Medications</th>
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<td>M</td>
<td>Right</td>
<td>8</td>
<td>Hypersignal in Right Sub cortical paraventricular</td>
<td>IMTG, rMTG</td>
<td>Trileptal, Zonégran, Urbanyl, Vimpat, Lamictal,</td>
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### Table 2: Descriptive statistics of notes in the 10 bins of increasing IC.

<table>
<thead>
<tr>
<th>Bin</th>
<th>Information Mean(SD)</th>
<th>Size of preceding interval Mean(SD)</th>
<th>Tonal stability Mean(SD)</th>
<th>Pitch Mean(SD)</th>
<th>Position Mean(SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bin 1</td>
<td>0.67(0.22)</td>
<td>1.57(0.74)</td>
<td>4.77(0.98)</td>
<td>68.66(3.51)</td>
<td>21.42(11.99)</td>
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<tr>
<td>Bin 2</td>
<td>1.19(0.09)</td>
<td>1.43(0.67)</td>
<td>4.90(1.20)</td>
<td>67.88(3.24)</td>
<td>20.48(11.10)</td>
</tr>
<tr>
<td>Bin 3</td>
<td>1.50(0.10)</td>
<td>1.39(0.89)</td>
<td>4.79(1.10)</td>
<td>68.47(3.27)</td>
<td>19.68(11.98)</td>
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<td>Bin 4</td>
<td>1.83(0.09)</td>
<td>1.08(1.15)</td>
<td>4.72(1.09)</td>
<td>68.19(3.26)</td>
<td>21.08(12.56)</td>
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<tr>
<td>Bin 5</td>
<td>2.13(0.08)</td>
<td>1.20(1.26)</td>
<td>4.54(1.09)</td>
<td>68.37(3.56)</td>
<td>20.90(11.52)</td>
</tr>
<tr>
<td>Bin 6</td>
<td>2.44(0.10)</td>
<td>1.37(1.35)</td>
<td>4.45(1.18)</td>
<td>68.37(3.26)</td>
<td>19.20(11.43)</td>
</tr>
<tr>
<td>Bin 7</td>
<td>2.77(0.10)</td>
<td>1.50(1.47)</td>
<td>4.42(1.17)</td>
<td>69.19(3.33)</td>
<td>19.51(11.70)</td>
</tr>
<tr>
<td>Bin 8</td>
<td>3.23(0.17)</td>
<td>2.44(1.58)</td>
<td>4.28(1.07)</td>
<td>69.56(3.13)</td>
<td>18.72(11.43)</td>
</tr>
<tr>
<td>Bin 9</td>
<td>4.10(0.36)</td>
<td>3.59(1.98)</td>
<td>4.27(1.21)</td>
<td>68.85(3.67)</td>
<td>19.49(9.81)</td>
</tr>
<tr>
<td>Bin 10</td>
<td>6.09(1.21)</td>
<td>5.42(2.62)</td>
<td>4.11(1.33)</td>
<td>69.17(4.02)</td>
<td>19.05(10.66)</td>
</tr>
</tbody>
</table>
Table 3: Distribution of contacts in the different regions of interest.

<table>
<thead>
<tr>
<th>ROI</th>
<th>Number of participants</th>
<th>Number of contacts in ROI</th>
<th>Total number of contacts in the ROI</th>
<th>MNI coordinates of contacts</th>
</tr>
</thead>
<tbody>
<tr>
<td>STG</td>
<td>6</td>
<td>21</td>
<td>[50.9 - 8.7 - 7.5], [48.1 -13.5 - 9.7], [54.5 - 10.9 - 8.1], [-46.7 - 18.8 -7.5], [-52.3 - 19.9 - 4.5], [-48.6 -33.4 8.9], [-51.5 -35.1 12.3], [-55.4 -36.0 16.5], [-59.3 -38.0 19.6], [-47.9 0.6 -17.3], [-51.2 0.7 -13.9], [-59.0 0.9 -8.1], [48.7 -32.0 6.2], [51.8 -33.4 9.4], [57.0 -36.0 12.6], [60.2 -37.5 15.6], [64.5 -39.8 18.9], [-49.5 -17.9 -7.5], [-55.3 -18.0 -4.0], [-63.9 -23.7 -0.3]</td>
<td></td>
</tr>
<tr>
<td>MTG</td>
<td>7</td>
<td>33</td>
<td>[45.9 -10.6 -10.9], [44.1 2.2 -29.9], [50.7 3.1 -30.3], [56.2 3.6 -28.5], [52.0 4.7 -25.0], [-52.1 -2.8 -18.4], [-56.8 -0.8 -18.1], [-62.7 0.3 -17.8], [-55.6 -8.0 -22.0], [-60.3 -45.1 -11.7], [-64.1 -46.1 -8.6], [-57.8 -25.3 -12.3], [-64.7 -26.6 -11.0], [-71.6 -28.6 -8.1], [-45.1 -11.9 -13.4], [-55.5 -9.2 -19.1], [-52.8 -26.3 -15.1], [-57.6 -28.7 -13.7], [-62.2 -30.6 -13.2], [-67.0 -32.7 -13.0], [40.2 -1.4 -32.1], [47.5 0 -30.8], [-56.3 -14.8 -21.9], [-54.8 -2.0 -21.3], [-54.1 8.0 -30.3], [-58.3 5.9 -29.1], [-53.2 -41.5 -6.7], [-58.3 -43.3 -7.0], [-62.5 -45.2 -5.8], [-52.4 -19.8 -15.6], [-59.7 -19.7 -14.3], [-52.4 -18.8 -18.0], [58.8 -19.3 -17.9]</td>
<td></td>
</tr>
<tr>
<td>IFG</td>
<td>pars</td>
<td>13</td>
<td>[37.7 29.5 -3.4], [41.9 31.8 -3.0], [46.1 34.3 -2.6], [49.3 37.0 -2.1], [-29.1 25.9 -12.7], [-44.1 36.5 -4.9], [22.1 15.4 -23.8], [26.7 19.6 -20.5], [29.9 23.3 -18.4], [34.2 26.0 -16.1], [37.3 28.8 -13.9], [41.6 33.1 -10.7], [45.9 35.6 -8.8]</td>
<td></td>
</tr>
<tr>
<td>Orbitalis</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACG</td>
<td>3</td>
<td>5</td>
<td>[9.9 30.6 30.0], [8.8 46.0 -6.9], [9.6 17.7 34.5], [18.5 22.3 31.0], [-15.9 48.4 -11.4]</td>
<td></td>
</tr>
<tr>
<td>Left AMYG</td>
<td>4</td>
<td>4</td>
<td>[-19.4 -5.1 -21.3], [-26.7 -3.4 -22.5], [-13.7 -4.2 -25.7], [-19.8 -1.4 -27.7]</td>
<td></td>
</tr>
<tr>
<td>INS</td>
<td>2</td>
<td>5</td>
<td>[33.5 26.9 -3.7], [42.8 15.7 8.8], [47.0 17.0 10.9], [-39.8 8.5 8.3], [-42.5 10.7 11.1]</td>
<td></td>
</tr>
</tbody>
</table>
Predictive processing of music in frontal and temporal cortices

Figure Captions

Figure 1: A schematic of a unit of the prediction cycle with hypothesized effects.

Figure 2: The IC profile of a single sample melody (A), the distribution of IC across all notes in the corpus (B), and the mean IC (and SD) of notes in the 10 IC bins (C).

Figure 3: Distribution of contacts in cortical (small circles) and subcortical (large circles) regions of interest. STG: superior temporal gyrus (blue), MTG: middle temporal gyrus (black); pars Orbitalis of IFG: inferior frontal gyrus (red); ACG: anterior cingulate gyrus (Pink); AMYG: amygdala (Yellow), INS: insula (Green). Note that right hemisphere data have been projected onto the left hemisphere.

Figure 4: The time-course of spectral power changes in the different frequency bands in bilateral STG, MTG and orbital IFG as a function of IC bin. Blue lines show time-courses for the three lowest IC bins, green for the four middle bins and red for the three highest bins. Grey shading indicates the time windows showing significant modulation by IC bin.

Figure 5: The distribution of contacts in the STG, MTG and orbital IFG showing modulation by information content significant at p<.01. STG: superior temporal gyrus (blue), MTG: middle temporal gyrus (black); pars orbitalis of the IFG: Inferior frontal gyrus (BA47) (red). Note that right hemisphere data have been projected onto the left hemisphere.

Figure 6: The time-course of spectral power changes in the different frequency bands in bilateral ACG, left amygdala, and bilateral insula as a function of IC bin. Blue lines show time-courses for the three lowest IC bins, green for the four middle bins, and red for the three highest bins. Grey shading indicates the time windows showing significant modulation by IC bin.

Figure 7: The distribution of contacts in the ACG, Insula and Amygdala showing modulation by information content significant at p<.01. ACG: anterior cingulate gyrus (Pink); AMYG: amygdala (Yellow), INS: insula (Green). Note that right hemisphere data have been projected onto the left hemisphere.