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Cortical network dynamics of perceptual decision-making in the human brain

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Goal-directed behavior requires the flexible transformation of sensory evidence about our environment into motor actions. Studies of perceptual decision-making have shown that this transformation is distributed across several widely separated brain regions. Yet, little is known about how decision-making emerges from the dynamic interactions among these regions. Here, we review a series of studies, in which we characterized the cortical network interactions underlying a perceptual decision process in the human brain. We used magnetoencephalography to measure the large-scale cortical population dynamics underlying each of the sub-processes involved in this decision: the encoding of sensory evidence and action plan, the mapping between the two, and the attentional selection of task-relevant evidence. We found that these sub-processes are mediated by neuronal oscillations within specific frequency ranges. Localized gamma-band oscillations in sensory and motor cortices reflect the encoding of the sensory evidence and motor plan. Large-scale oscillations across widespread cortical networks mediate the integrative processes connecting these local networks: Gamma- and beta-band oscillations across frontal, parietal, and sensory cortices serve the selection of relevant sensory evidence and its flexible mapping onto action plans. In sum, our results suggest that perceptual decisions are mediated by oscillatory interactions within overlapping local and large-scale cortical networks.

Keywords: perceptual decision-making, neuronal oscillations, neuronal synchronization, magnetoencephalography, sensorimotor integration, attention, gamma-band, beta-band

We often need to select our actions based on perceptual interpretations of noisy, incomplete, or ambiguous sensory information about our environment. Imagine driving on the highway on a rainy night. To discern if the cars in front of you slowed down, you try to judge whether their brake lights are looming. After accumulating the sparse sensory information for a few moments you eventually decide to hit the brake.

During sensory-guided behaviors such as this one, perceptual states are flexibly mapped onto our motor actions, a process referred to as perceptual decision-making (Figure 1; Gold and Shadlen, 2001, 2007; Usher and McClelland, 2001; Heekeren et al., 2008; Tosoni et al., 2008). Dynamic interactions between multiple processing stages of the brain’s sensorimotor pathways lie at the heart of such decision processes. Invasive recordings in monkeys performing simple sensorimotor tasks have characterized the detailed functional properties of individual neurons in these processing stages (Schall, 2001; Glimcher, 2003; Romo and Salinas, 2003; Gold and Shadlen, 2007; Kable and Glimcher, 2009). These studies have shown that neurons in sensory, parietal, and frontal cortices, are involved in encoding the sensory evidence, accumulating this evidence over time, and planning the ensuing motor action. These results have inspired circuit models of the large-scale interactions between these processing stages (Wang, 2002, 2008; Mazurek et al., 2003; Lo and Wang, 2006; Gold and Shadlen, 2007). However, until recently, these large-scale interactions have only rarely been characterized directly (Heekeren et al., 2004; Pesaran et al., 2008), but mainly been inferred (Wang, 2002, 2008; Mazurek et al., 2003; Lo and Wang, 2006; Gold and Shadlen, 2007).

A hallmark of the neuronal interactions underlying perceptual decisions is their flexibility. The mapping of sensory evidence onto motor actions is not a stereotyped reflex but depends strongly on the decision-maker’s current behavioral goals. In different contexts, the exact same sensory input can be mapped onto different actions. Similarly, the same action can be selected in the face of different sensory inputs. To account for this flexibility, theories of cognitive control postulate “top-down” signals that selectively bias the flow of information in the brain’s sensorimotor pathways, such that only the sensory and motor neuronal populations relevant for the goal at hand communicate with one another, whereas goal-irrelevant pathways are shut down (Miller and Cohen, 2001). These top-down signals are thought to originate from prefrontal and posterior parietal association cortices (Miller and Cohen, 2001; Corbetta and Shulman, 2002).

Most current models of perceptual decision-making conceptually align the decision formation as a feedforward integration process from sensation to action (Usher and McClelland, 2001; Mazurek et al., 2003; Smith and Ratcliff, 2004). However, the actual cortical circuits implementing this process likely engage in highly recurrent interactions (Lamme and Roelfsema, 2000; Wang, 2008). Such recurrent interactions are mediated by the abundant bi-directional
cortico-cortical connections (Felleman and Van Essen, 1991). For example, sensorimotor association regions of the parietal and frontal cortex commonly display highly similar response profiles during decision formation (Kim and Shadlen, 1999; Shadlen and Newsome, 2001; Pesaran et al., 2008). Furthermore, neuronal activity even at the earliest sensory processing stages (such as visual cortical areas V1 and V2) does not merely reflect the sensory evidence but also the result of the perceptual decision process, likely via feedback from downstream areas (Donner et al., 2008a,b; Nienborg and Cumming, 2009). In sum, even simple perceptual decisions emerge from recurrent and flexible interactions between widely distributed regions of the cerebral cortex. How can we get a glimpse on these interactions?

**NEURONAL OSCILLATIONS: A WINDOW ON THE DYNAMICS OF DECISION-MAKING**

Magnetoencephalography (MEG) combined with source reconstruction techniques is ideally suited to simultaneously monitor the dynamics of neuronal population activity across widely separated regions of the human cerebral cortex and to characterize the dynamic interactions between these regions. Moreover, the exquisite temporal resolution of MEG allows for characterizing the fine temporal structure of cortical population activity. A pervasive feature of this activity is that it displays oscillations across a broad range of frequencies and spatial scales (Buzsaki and Draguhn, 2004).

A growing body of evidence suggests that such neuronal oscillations play an important functional role in cortical information processing. Coherent oscillations, within and between cortical regions may flexibly regulate the interactions among distributed neuronal populations (Engel et al., 2001; Salinas and Sejnowski, 2001; Varela et al., 2001; Fries, 2005; Haider and McCormick, 2009). On the one hand, synchronization of presynaptic spikes within a local cortical region enhances their functional impact on postsynaptic neurons in a super-additive fashion (Alonso et al., 1996; König et al., 1996; Azouz and Gray, 2003). On the other hand, the phase-alignment of presynaptic spikes to postsynaptic oscillations modulates their efficiency in driving postsynaptic spikes (Fries, 2005; Somers et al., 2007; Haider and McCormick, 2009). Both types of synchrony may serve as flexible mechanisms to control the gain of local and long-range neuronal communication.

Furthermore, neuronal oscillations at different frequencies may provide valuable mechanistic information about the interactions between groups of neurons (Siegel and Donner, 2010). First, they may index interactions within specific neural circuits. Local cortical oscillations in the gamma-band (30–80 Hz) provide an intriguing example for a spectral signature for which the detailed circuit-level mechanisms are becoming increasingly clear. Second, more generally, neuronal oscillations may reflect different types of neural interactions. In particular, they may reflect reverberating activity within local as well as large-scale cortical networks (Wang, 2003). Such reverberation may mediate persistent cortical activity and recurrent cortical processing (Siegel et al., 2000, 2009; Tallon-Baudry et al., 2001; Pesaran et al., 2002; Van Der Werf et al., 2008), which, in turn, may underlie the protracted accumulation of sensory evidence during decision-making. Third, neuromodulators such as norepinephrine or acetylcholine profoundly shape the synaptic interactions between cortical neurons. Thus, the spectral signature of neuronal oscillations may also reflect the effects of neuromodulators (Steriade, 2000; Rodriguez et al., 2004), which, in turn, play important roles in decision-making (Aston-Jones and Cohen, 2005; Yu and Dayan, 2005). In sum, the spectral signature of cortical population activity may provide rich information about the detailed neuronal circuit dynamics underlying decision-making. Such information might only be weakly expressed in the spiking activity of individual neurons.

**DISSECTING A PERCEPTUAL DECISION PROCESS**

How does the flexible mapping of sensory “evidence” onto action plans emerge from these neuronal circuit dynamics? We addressed this question in a series of MEG experiments in humans, dissecting a single perceptual decision into its component sub-processes. In all experiments subjects judged visual motion signals contained in dynamic random-dot patterns and reported their perceptual interpretation of these stimuli by button-press. By adjusting the fraction of coherently moving dots in the random-dot patterns, we could precisely and specifically control the strength of the sensory feature of interest, i.e., visual motion (Figure 2A; Scase et al., 1996). Moreover, this enabled us to titrate the sensory evidence to the subjects’ individual detection threshold. Near psychophysical threshold, perceptual choices vary form trial to trial in the face of identical sensory inputs (Green and Swets, 1966). We could exploit these intrinsic fluctuations to isolate the neuronal mechanisms underlying the varying choices without confounding variations of the sensory input. This task also allowed us to focus our analyses on well-characterized cortical regions, which had been identified in previous single-unit recordings in monkeys (Newsome et al., 1989; Britten et al., 1993; Kim and Shadlen, 1999; Shadlen and Newsome, 2001; Roitman and Shadlen, 2002; Gold and Shadlen, 2007) and fMRI studies in humans (Rees et al., 2000; Shulman et al., 2001; Serences and Boynton, 2007). We used an adaptive spatial filtering technique (linear beamforming; Van Veen et al., 1997; Gross et al., 2001) to estimate the dynamics of neuronal population activity within and among these cortical processing stages.

**GAMMA-BAND ACTIVITY IN VISUAL CORTEX REFLECTS SENSORY EVIDENCE**

We set out by investigating the input stage of the decision process, i.e., the encoding of sensory evidence for coherent motion (Siegel et al., 2007). Subjects were presented with dynamic random-dot
patterns at various different levels of motion coherence (Figure 2A) and, for each stimulus, reported the net motion direction (up vs. down) by button-press.

The moving stimuli induced robust population responses with a characteristic dynamic signature at sensors overlying visual cortex (Figure 2B). Following a transient low-frequency response (reflecting the visual evoked field), the MEG power increased in a broad high-frequency range (about 40–150 Hz), accompanied by a decrease in the low-frequency range below 40 Hz. Both of these response components were sustained as long as the stimuli were on the screen. The response in a narrower gamma-band from about 60 to 100 Hz increased approximately linearly with motion coherence. The solid line indicates the significant linear effect ($p < 0.01$; $n = 7$ subjects). (D) Cortical distribution of the average gamma-band response (62–102 Hz) across all levels of motion coherence (red shading) and of the linear increase of gamma-band activity with motion coherence (blue shading). Data is shown for one representative subject. See Siegel et al. (2007) for full details and data from all individual subjects (IPS, intraparietal sulcus; MT, middle temporal area). Reprinted and modified with permission from Siegel et al. (2007).

FIGURE 2 | Gamma-band activity in visual cortex reflects the encoding of sensory evidence. (A) Schematic illustration of dynamic random-dot patterns at three levels of motion coherence. The percentage of coherently moving dots controls the strength of the motion signal and thus the strength of sensory evidence for the motion-discrimination task. (B) Time–frequency representation of neural responses to dynamic random-dot patterns recorded at MEG sensors overlying visual cortex. Responses were averaged across six logarithmically scaled levels of motion coherence ranging from 0 to 100% (see panel C) and across seven subjects. (C) Gamma-band activity (62–102 Hz) over visual cortex during stimulus viewing (100–500 ms post stimulus onset) increases approximately linearly with motion coherence. The solid line indicates the significant linear effect ($p < 0.01$; $n = 7$ subjects). (D) Cortical distribution of the average gamma-band response (62–102 Hz) across all levels of motion coherence (red shading) and of the linear increase of gamma-band activity with motion coherence (blue shading). Data is shown for one representative subject. See Siegel et al. (2007) for full details and data from all individual subjects (IPS, intraparietal sulcus; MT, middle temporal area). Reprinted and modified with permission from Siegel et al. (2007).
**BUILD-UP OF CORTICAL POPULATION ACTIVITY IN MOTOR CORTEX REFLECTS THE EVOLVING ACTION PLAN**

The spectral signature of sensory encoding was highly consistent with the signature of action encoding at the cortical output stage of the decision process. In another experiment, the dynamic random-dot patterns shown on each trial contained either pure dynamic noise, or a weak motion signal titrated to each subject’s individual detection threshold (Donner et al., 2009). Subjects formed a decision about the presence or absence of this target motion signal. Stimuli were presented for a fixed duration of 2 s and followed by a blank delay, after which subjects reported their choice (Figure 3A). In this task, psychophysical coherence detection thresholds decrease over stimulus durations of more than 3 s in humans. Thus, presenting motion signals at the subjects’ individual detection threshold ensured that each subject accumulated the evidence for motion until the end of the stimulus interval.

To track the subjects’ evolving plan to report either “target present” or “target absent” within the brain, we mapped these two perceptual choices onto button-presses with different hands. This allowed us to exploit the contralateral bias of the cortical motor system (i.e., stronger activity in the hemisphere contralateral to the motor movement). Just before the execution of the motor response, gamma-band activity (about 60–100 Hz) in primary motor (M1) and premotor (PMd) cortices was selectively enhanced contralateral to the button-press (Figures 3A,B). As for the stimulus-driven responses in visual cortex discussed above, this gamma-band enhancement was accompanied by a decrease of low-frequency activity (about 10–35 Hz) with two distinct peaks in the alpha- and beta-bands. This antagonistic modulation of gamma-band and low-frequency activity in motor and premotor cortex is in line with previous intracranial and MEG studies (Crone et al., 1998a,b; Pfurtscheller et al., 2003; Cheyne et al., 2008).

Crucially, both components of the effector-selective activity were not only expressed after the completion of the decision process (i.e., after stimulus offset), but they built up gradually during stimulus viewing, while subjects were forming their decision (Figure 3C). Both components reliably predicted subjects’ final choice on single trials seconds before its execution. Thus, local gamma-band activity and an accompanying suppression of low-frequency activity in motor cortex reflect the evolving action selection in this protracted decision process.

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**FIGURE 3 | Oscillatory population activity in motor cortex reflects the evolving action plan.** (A) Time–frequency representation of lateralized neural activity recorded by MEG sensors overlying motor cortex during the motion-detection task. The lateralization displays the difference in activity contralateral minus ipsilateral to the side of the button-press. (B) Cortical distribution of the lateralized activity (left minus right-hand button-press) in the beta- and gamma-band during the 0.5 s before button-press. This effector-selective activity is localized to the motor cortex. (C) Time courses of choice-predictive lateralized beta- and gamma-band activity (contralateral minus ipsilateral to “yes” button) in M1, expressed as z-scores for the four trial categories. The shaded area marks the time of stimulus presentation. The mapping of perceptual choices (“yes”/“no”) onto hand movements (left/right) was counterbalanced across subjects. Dashed lines indicate significance levels (p = 0.05 by two-sided t-test). (D) Linear regression between the temporal integral of gamma-band activity in M1 across the stimulus interval (64–100 Hz) and the lateralized beta- and gamma-band activity in M1 (contralateral minus ipsilateral to “yes” button) at the end of the stimulus interval. Responses were binned (100 trials per bin) by the MT response integral and z-transformed. Reprinted and modified with permission from Donner et al. (2009).
We interpret the gamma-band enhancement as a more direct marker of the local neuronal representation than the suppression of low-frequency activity, for a number of reasons. First, for both, sensory and motor responses, gamma-band activity is positively correlated with local processing while low-frequency activity shows the opposite relationship. Second, intracranial recordings from motor cortex show that gamma-band modulations are typically anatomically more focused than the low-frequency suppressions (Crone et al., 1998a, b; Pfurtscheller et al., 2003). This also explains why these premotor gamma-band modulations are more difficult to detect in extracranial signals like EEG and MEG than in direct invasive recordings from the cortical surface. Third, gamma-band activity in motor cortex typically shows a stronger effector-selectivity (Crone et al., 1998a, b; Pfurtscheller et al., 2003; Rickert et al., 2005; Cheyne et al., 2008). Fourth, in our own data, the premotor gamma-band modulations were more closely linked to the ongoing decision process, as discussed in the next section.

**CHOICE-PREDICTIVE ACTIVITY IS DRIVEN BY INTEGRATED SENSORY EVIDENCE**

The results discussed so far established frequency-specific neuronal signatures for the task-relevant sensory evidence (visual motion: gamma-band activity in MT) and the evolving motor plan (button-press: gamma-band activity in motor cortex). A large body of evidence suggests that the evidence encoded in sensory cortex is integrated across time during the decision process (Kim and Shadlen, 1999; Shadlen and Newsome, 2001; Usher and McClelland, 2001; Roitman and Shadlen, 2002; Smith and Ratcliff, 2004; Gold and Shadlen, 2007). The simultaneous monitoring of sensory evidence and motor plan with MEG provides an ideal opportunity to test this model. If the choice-predictive build-up of activity in motor cortex is driven by the temporal integral of evidence for motion delivered by MT, then the trial-to-trial fluctuations of this motor activity should be predicted by the trial-to-trial fluctuations of the integrated MT activity.

This is what we found (Figure 3D; Donner et al., 2009). The temporal integral of gamma-band activity in MT predicted the choice-predictive motor activity in the gamma- and beta-bands. The opposite sign of correlation reflected the opposite sign of movement-selective activity in these bands (Figures 3A–C). This correlation was stronger for premotor activity in the gamma-band, suggesting that the local gamma-band activity is more tightly linked to the local neuronal representation than low-frequency activity (see above). Notably, the correlation between choice-predictive motor activity and gamma-band activity in MT was much weaker if not the integral of MT activity but only its instantaneous activity was taken into account. Further, the correlation between MT-integral and motor activity did not merely emerge at the end of the decision process (the stimulus offset), but was continuously expressed during stimulus viewing, irrespective of whether the decision-process resulted in a “target present” or a “absent” choice. These observations establish a first direct link between the neuronal dynamics of sensory evidence and motor plan during perceptual decision-making and provide strong evidence for the temporal accumulation model.

Importantly, our results do not imply that the temporal integration process takes place in the motor cortex. Any processing stage between MT and the motor cortex may be involved in accumulating the sensory evidence delivered by MT. Posterior parietal and prefrontal association cortex have long been known to exhibit persistent neuronal activity (Miller and Cohen, 2001; Wang, 2001; Pessoa et al., 2002; Machens et al., 2005). More recently, these regions have also been implicated in the temporal integration of sensory evidence (Wang, 2002; Heekeren et al., 2004, 2008; Gold and Shadlen, 2007; Kaiser et al., 2007).

The results discussed in this section show that the outcome of the temporal integration process is continuously routed through all the way even to primary motor cortex. This conclusion is consistent with a previous microstimulation study in the monkey’s oculomotor system (Gold and Shadlen, 2000) and suggests a highly continuous flow of information in the sensorimotor pathways (Kim and Shadlen, 1999; Shadlen and Newsome, 2001; Horwitz et al., 2004; Gold and Shadlen, 2007). This contradicts the traditional notion of a “central executive” that first completes the decision process before sending its end result to the output stage.

**FRONTAL–PARietAL BETA-BAND OSCILLATIONS PREDICT THE ACCURACY OF EVOLVING DECISION**

How can one obtain a more direct handle on the neuronal mechanism of the evidence–action mapping per se? A powerful strategy is to exploit the fact that, at low levels of sensory evidence, subjects often make incorrect choices. A substantial fraction of these performance fluctuations is thought to be driven by variability at the decision stage (Shadlen et al., 1996). Specifically, we reasoned that the more efficiently the evidence–action mapping mechanism operates on any given trial, the more likely subjects will make a correct choice on that trial. Thus, we compared neuronal activity in the motion-detection task between correct and incorrect trials (Donner et al., 2007). Enhanced activity in a narrow “low beta” frequency range (about 12–24 Hz) predicted the correctness of the subjects’ upcoming choice (Figure 4A). This effect specifically emerged during the stimulus interval, that is, while subjects were accumulating the sensory evidence and while activity was building up in their motor cortex. This performance-predictive beta-band activity was expressed in a widespread network across the cerebral cortex, predominantly in posterior parietal and dorsolateral prefrontal cortex and, less strongly, in visual (e.g., area MT) and motor cortex (Figure 4B).

Beta-band activity in these regions was higher before “hits” than “misses” on target present trials and higher before “correct rejects” than “false alarms” on target absent trials (Figure 4C). Thus, this beta-band activity predicted the accuracy (correct vs. error) but not the content (target present vs. target absent) of the upcoming choice. This suggests that the widespread, performance-predictive beta-band oscillations reflect the computations governing the decision process rather than neuronal representations involved at its different stages (deCharms and Zador, 2000). In other words, the trial-to-trial fluctuations of beta-band activity do not reflect fluctuations of the neuronal representations of sensory evidence, decision variables, or action plans, but rather fluctuations of the flexible mapping between these representations.

What is the specific functional role of the performance-predictive beta-band oscillations? Several recent studies reported analogous beta-band oscillations across frontal and parietal cortices during visuomotor tasks in humans (Gross et al., 2004; Hipp et al., 2011)
Top-down attention is mediated by local and large-scale synchronization during decision formation

Perceptual decision-making is not a passive response, but an active and flexible process. The evidence–action mapping can be flexibly adapted to our current behavioral goal. For example, this mapping bears functional similarity with attention. It reflects a mechanism that, similar to attention, governs the accuracy of a flexible sensorimotor transformation. The beta-band oscillations may reflect reverberant activity within and among visual, frontoparietal, and motor cortices (Wang, 2002; Engel and Fries, 2010). This large-scale network reverberation may help maintain and accumulate sensory evidence during the decision formation (Donner et al., 2007). Such activity may also reflect the maintenance of the sensorimotor mapping rule between accumulated sensory evidence and action. Further experiments are required to assess these not mutually exclusive scenarios.

and monkeys (Pesaran et al., 2002, 2008; Buschman and Miller, 2007, 2009). The performance-predictive frontoparietal beta-band activity in our task likely does not reflect trial-to-trial fluctuations of the subjects’ level of arousal or top-down selective attention. Fluctuations of arousal are typically slow, spanning several trials. Thus, such fluctuations should be expressed during the pre-stimulus baseline interval. Also fluctuations of selective attention are often evident during the pre-stimulus baseline (Ress et al., 2000). By contrast, there was no evidence for performance-predictive beta-band activity before stimulus onset, but this activity only occurred during the decision period of the trial. Furthermore, several studies discussed in the next section have demonstrated very different spectral profiles for top-down selective attention in similar cortical regions.

Nevertheless, the performance-predictive beta-band activity is often selective for only those sources of sensory evidence that are relevant for achieving our goal. Think again about driving your car at night. You may not only decide to hit the brake when traffic slows down but also when you want to exit the highway. In the former situation you pay attention to the cars in front of you. By contrast, in the latter case your motor response is based on other parts of the available visual information such as the road signs signaling your exit.

The top-down selection of sensory evidence is commonly referred to as “selective attention.” Oscillatory synchronization of neuronal activity has frequently been proposed as a candidate mechanism of attention (Gruber et al., 1999; Worden et al., 2000; Engel et al., 2001; Fries et al., 2001; Bichot et al., 2006; Saalmann et al., 2007; Wyart and Tallon-Baudry, 2008). In another MEG experiment (Siegel et al., 2008), we thus investigated how selective attention modulated cortical population dynamics along the different stages of the sensorimotor pathway. On each trial, subjects were visually cued to covertly shift their focus of attention to either the left or right visual hemifield, where two random-dot patterns were simultaneously presented after a delay (Figure 5A). Subjects then judged the net-direction of motion of the cued stimulus while ignoring the uncued stimulus. Again, we adjusted the levels of motion coherence to the subjects’ individual psychophysical thresholds. This ensured that the task was attentionally demanding. To isolate the neuronal correlates of spatially selective attention, we contrasted modulations of neuronal activity between the hemispheres contralateral and ipsilateral to the attended hemifield.

Visual–spatial attention modulated local oscillatory population activity throughout the visuomotor pathway in a spatially selective fashion (Figure 5B). We observed attentional modulations of local synchrony in visual cortical areas consistent with previous studies (Fries et al., 2001; Bichot et al., 2006; Saalmann et al., 2007; Wyart and Tallon-Baudry, 2008). Reprinted and modified with permission from Donner et al. (2007).
relations differed markedly between temporal intervals with more prominent low-frequency suppressions before stimulus onset and enhanced high-frequency activity during stimulation. Thus, the large-scale investigation of attentional modulations throughout the visuomotor pathway revealed a remarkable regional and temporal specificity of these modulations.

Having established that attention modulates oscillatory population activity within visual cortex and attentional control regions, we investigated if the attentional selection of the behaviorally relevant stimulus may be mediated by long-range synchronization between these processing stages. Membrane-potential oscillations establish periodic windows of enhanced excitability (Haider and McCormick, 2009). Attention may thus involve the phase-alignment of presynaptic spikes to postsynaptic oscillations to dynamically

Figure 5 | Local and large-scale synchrony reflects the attentional selection of sensory evidence. (A) Schematic illustration of the stimulus display during the spatially cued motion-discrimination task. (B) Attentional modulation of local oscillatory population activity in four regions of interest (marked on the inflated hemispheres to the left). Attentional modulations are displayed as predictive indices that quantify the probability with which an ideal observer can predict the direction of attention (left vs. right) from lateralization of neural responses between the left and right hemispheres on a single trial level. Predictive indices larger or smaller than chance level (0.5) correspond to an attentional enhancement or suppression of activity in the hemisphere contralateral to the attended hemifield, respectively. (C) Attentional modulation of frequency-specific long-range synchronization. Panels in the upper row display between which cortical regions spatial attention significantly enhanced synchronization in the hemisphere contralateral as compared to ipsilateral to the attended visual hemifield. Panels in the lower row depict corresponding attentional reductions of synchronization. Colors indicate the frequency bands of significantly modulated long-range synchrony. All attentional modulations are displayed separately for the delay interval (750–0 ms before stimulus onset) and stimulus interval (100–500 ms after stimulus onset). (*p < 0.05, **p < 0.01, ***p < 0.001 corrected; FEF, frontal eye field; PIPS, posterior intraparietal sulcus; MT, middle temporal area; V1/V2, pericalcarine cortex). Reprinted and modified with permission from Siegel et al. (2008).
regulate the information flow between areas (Engel et al., 2001; Fries, 2005). Consistent with this hypothesis, we found that attention modulated long-range oscillatory synchronization between frontoparietal control regions (IPS and FEF) and visual cortex (MT; Figure 5C). In contrast to the effects on local population activity, attentional modulations of long-range synchrony were comparatively stimulus independent (compare Figures 5C,B). Before and during stimulus presentation, attention selectively enhanced long-range synchronization in the gamma-band (35–100 Hz) and suppressed low-frequency synchronization in the alpha- and beta-band (5–35 Hz) in the hemisphere processing the attended stimulus. Thus, visuospatial attention established a spatially selective pattern of long-range synchronization between frontoparietal and early sensory processing stages.

This modulation of synchrony may allow specific neuronal populations within control regions to communicate top-down signals to specific populations in visual cortex without the control regions being necessarily involved in the sensorimotor transformation per se. Alternatively, phase coherence throughout the sensorimotor pathway may enhance the feedforward routing of selected information to the motor stage. Recent results on synchrony between FEF and V4 in monkeys (Gregoriou et al., 2009) suggest that both mechanisms are involved. During sustained attention, synchrony primarily mediates feedforward interactions, while feedback interactions are dominating during shifts of attention. In summary, our results suggest that long-range oscillatory synchronization between frontoparietal control regions and early sensory processing stages mediates the selective routing of sensory information required for the perceptual decision at hand.

**LOCAL AND LARGE-SCALE OSCILLATORY NETWORK INTERACTIONS MEDIATE PERCEPTUAL DECISION-MAKING**

The experiments described above provide a first comprehensive characterization of the cortical dynamics underlying a perceptual decision. We delineated the spectral signatures and cortical distributions of its component sub-processes. This revealed multiple local and large-scale oscillatory networks mediating the decision (Figure 6).

At the peripheral stages of the decision process (visual and motor cortex), local gamma-band activity consistently reflected the encoding of sensory evidence and the evolving motor plan (Figure 6A). Both, sensory and motor-related gamma-band activity was accompanied by a suppression of local low-frequency activity. Converging evidence suggest that local cortical gamma-band oscillations are generated by locally recurrent interactions between excitatory neurons and GABAergic interneurons (Siegel et al., 2000; Hasenstaub et al., 2005; Bartos et al., 2007; Gieselmann and Thiele, 2008; Cardin et al., 2009). These excitatory–inhibitory feedback loops become engaged during enhanced processing within local cortical networks and play an important role in shaping the structure of local neuronal representations (Heeger et al., 1996; Shapley et al., 2003). Thus, gamma-band activity within visual and motor cortex indexes the local circuit interactions underlying the encoding of sensory evidence and motor plans. Furthermore, local excitatory–inhibitory interactions, and thus local gamma-band activity, may not only be driven by feedforward inputs to the local network (e.g., visual input in visual cortex) but also by feedback signals originating from higher cortical regions (e.g., attentional signals in visual cortex; Fries et al., 2001; Siegel et al., 2008).

![Local Gamma Networks Evidence & Action Encoding](image1)

**FIGURE 6 | Oscillatory interactions in local and large-scale cortical networks mediate perceptual decision-making.** (A) Local gamma-band activity in visual (MT) and motor cortex (M1) reflects the encoding of sensory evidence and motor plan, respectively. Top-down processes modulate this local activity. (B) Large-scale beta-band activity in a network centered on prefrontal (dIPFC) and parietal cortex (IPS) is involved in mapping the sensory evidence onto motor plans. (C) Large-scale gamma-band synchrony in a network across prefrontal (FEF), parietal (IPS), and motor cortex (MT) mediates the attentional selection of task-relevant sensory evidence (M1, primary motor cortex; MT, middle temporal area; dIPFC, dorsolateral prefrontal cortex; IPS, intraparietal sulcus; FEF, frontal eye field).
The whole-brain coverage of MEG allowed us to also identify oscillatory interactions in two large-scale cortical networks as central components of the decision process. First, beta-band activity in a widespread network centered on prefrontal and parietal association cortices seems to be instrumental in linking sensory evidence to motor plans (Figure 6B). Specifically, this large-scale beta-band activity may mediate the active maintenance of past evidence during its accumulation and/or its flexible routing to motor plans. Second, top-down attentional selection of particular sources of sensory evidence was associated with enhanced gamma-band synchronization in another large-scale network, spanning frontoparietal attentional control regions and extrastriate visual areas (Figure 6C). Gamma-band synchronization within this network may mediate the attentional selection of the behaviorally relevant visual inputs.

Remarkably, the large-scale beta- and gamma-band networks partially overlapped in the posterior parietal cortex. This suggests that overlapping or even identical groups of neurons within this region may simultaneously participate in these two distinct large-scale networks. The posterior parietal cortex may thus serve as a hub integrating the large-scale processes mediated by these networks. Thus, shifting the perspective from the responses of individual cortical neurons onto their embedding into large-scale functional networks may help understand the functional role of posterior parietal cortical neurons onto their embedding into large-scale functional networks (Kopell et al., 2000). Furthermore, the spectral signatures of large-scale interactions may also depend on local oscillatory mechanisms underlying these oscillations. As outlined above, the detailed mechanisms underlying local gamma-band activity are becoming increasingly clear. By contrast, comparatively little is known about the mechanisms shaping the spectral signatures of oscillatory interactions within large-scale cortical networks. These signatures likely reflect global network properties such as the number of involved regions, synaptic distances, and conduction delays as well as local cellular properties such as the time-constants of various membrane conductances (Kopell et al., 2000). Furthermore, the spectral signatures of large-scale interactions may also depend on local oscillatory networks embedded within these larger networks. Locally generated oscillations such as local gamma-band activity may provide the basic temporal scaffolding necessary to synchronize activity across larger networks. Large-scale spectral signatures may thus reflect global network properties, cellular properties, as well as the spectral signatures of embedded local oscillations. In sum, the spectral signatures of local and large-scale neuronal oscillations provide valuable information about the specific circuit-level interactions underlying decision-making.

CONCLUSIONS AND FUTURE DIRECTIONS

Taken together, our results begin to unravel how neuronal interactions within overlapping cortical networks form flexible interfaces between perception and action during simple, well-controlled cognitive tasks. On the one hand, fast oscillations within local networks subserve the representation of sensory evidence and motor plans at the periphery of the decision process. On the other hand, coherent oscillations across large-scale cortical networks mediate the integrative processes connecting these local representations, such as the selection of relevant sensory evidence and its flexible mapping onto action plans.

We are still far from understanding the mechanisms that underlie these oscillatory interactions during decision formation. We propose three directions of future research that may be crucial for unraveling these mechanisms. A first fundamental question is if the large-scale oscillatory interactions between distant brain regions are regulated by a superordinate “control region,” or if they primarily emerge from distributed network interactions. Evidence suggests that the prefrontal cortex may flexibly boost or block such interregional couplings, as necessary for the task at hand (Miller and Cohen, 2001). Indeed, our results revealed the FEF, a prefrontal region, as an integral part of the large-scale network controlling the flexible evidence selection. However, these interregional couplings may also emerge in a decentralized, self-organizing fashion. Future studies should distinguish between these alternatives.

Second, it remains to be determined how neuronal processing is integrated among distinct large-scale oscillatory networks. The partially overlapping beta- and gamma-band networks discussed above provide an intriguing example. Cross-frequency interactions, such as phase–amplitude and phase–phase coupling between different rhythms, may play an important role for linking different oscillatory processes. Indeed, a growing body of evidence demonstrates different types of cross-frequency interactions in cortical population activity (Schack et al., 2002; Lakatos et al., 2005; Palva et al., 2005; Canolty et al., 2006; Jensen and Colgin, 2007; Siegel et al., 2009). Future studies should investigate if these cross-frequency couplings mediate interactions between different oscillatory networks and if these interactions between networks are modulated in a task-dependent fashion.

Third, the cortical systems involved in decision-making are known to be under profound influence of ascending neuromodulatory systems (Montague et al., 2004; Aston-Jones and Cohen, 2005; Yu and Dayan, 2005). These neuromodulatory systems are thus in an ideal position to dynamically shape these large-scale cortical interactions. Indeed evidence suggests that neuromodulatory systems play an important role in orchestrating oscillatory cortical activity (Rodriguez et al., 2004; Goard and Dan, 2009). Studying the impact of these neuromodulators on the patterns of large-scale cortical activity during decision-making will open up a new dimension in this field of research.

The results described here show that frequency-specific cortical population dynamics, as measured with MEG, provide a powerful window into the distributed dynamics of decision-making in the working human brain. Future studies should exploit this window to further our understanding of what it means “to decide.”

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