Behavioral/Cognitive

Cortical Oscillatory Mechanisms Supporting the Control of Human Social–Emotional Actions

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The human anterior prefrontal cortex (aPFC) is involved in regulating social–emotional behavior, presumably by modulating effective connectivity with downstream parietal, limbic, and motor cortices. Regulating that connectivity might rely on theta-band oscillations (4 – 8 Hz), a brain rhythm known to create overlapping periods of excitability between distant regions by temporally releasing neurons from inhibition. Here, we used MEG to understand how aPFC theta-band oscillations implement control over prepotent social–emotional behaviors; that is, the control over automatically elicited approach and avoidance actions. Forty human male participants performed a social approach–avoidance task in which they approached or avoided visually displayed emotional faces (happy or angry) by pulling or pushing a joystick. Approaching angry and avoiding happy faces (incongruent condition) requires rapid application of cognitive control to override prepotent habitual action tendencies to approach appetitive and to avoid aversive situations. In the time window before response delivery, trial-by-trial variations in aPFC theta-band power (6 Hz) predicted reaction time increases during emotional control and were inversely related to beta-band power (14 – 22 Hz) over parietofrontal cortex. In sensorimotor areas contralateral to the moving hand, premovement gamma-band rhythms (60 – 90 Hz) were stronger during incongruent than congruent trials, with power increases phase locked to peaks of the aPFC theta-band oscillations. These findings define a mechanistic relation between cortical areas involved in implementing rapid control over human social–emotional behavior. The aPFC may bias neural processing toward rule-driven actions and away from automatic emotional tendencies by coordinating tonic disinhibition and phasic enhancement of parietofrontal circuits involved in action selection.

Key words: approach–avoidance; frontal pole; MEG; phase-amplitude coupling; social–emotional control; theta-band oscillations

Introduction

Human cooperative social environment relies on our ability to control social–emotional behavior (Hare, 2017). The importance of this ability is illustrated by conditions in which this control fails. For instance, in social anxiety disorder, persistent avoidance of social interactions causes impairment in social and occupational functioning (Clark and Wells, 1995; Craske and Stein, 2016). Social–emotional disorders also illustrate how regulating social–emotional behavior requires more than suppressing automatic emotional action tendencies. In fact, social–emotional regulation is an action selection feat involving the selection of ade-
quate behaviors from numerous potential strategies while anticipating the consequences of those behaviors in other agents. For example, in a job interview, applicants might make it more likely to get the job when they overcome their tendency to avoid the test and dare to approach when asked who will present first. These emotionally laden counterfactual computations are thought to be implemented by the anterior prefrontal cortex (aPFC) (Daw et al., 2006; Boorman et al., 2009; Volman et al., 2013), a region also known to influence downstream neural activity in the amygdala and posterior parietal cortex (PPC) during emotional action control (Mars et al., 2011; Volman et al., 2011a, 2013, 2016). However, the neural mechanisms that support social–emotional regulation across this network remain unclear. In this study, we use neural oscillations as a metric for understanding how aPFC rapidly selects and implements alternative courses of action over prepotent habitual social–emotional behaviors.

Recent studies have shown that implementation of prefrontal control is often rhythmic, transferring information between regions through entrainment of neural oscillations (Helfrich and Knight, 2016). aPFC neurons tend to phase lock their spiking and gamma-band firing to the theta–band rhythm (4–8 Hz) (Ardid et al., 2015; Voytek et al., 2015). Frontal theta–band oscillations have consistently been involved in control over motivational behavior (Cavanagh and Frank, 2014; Cooper et al., 2015), including control over state-induced affective behavioral biases (Cavanagh et al., 2013) and feedback–related control in exploration behavior (Cavanagh et al., 2012). Theta-band oscillations might support emotional control by temporally orchestrating release of neurons from inhibition, thus creating overlapping periods of excitability across the cerebral network involved in social–emotional regulation (Colgin, 2013; Lisman and Jensen, 2013; Volman et al., 2013). Human aPFC has extensive anatomical connections with portions of the PPC (Mars et al., 2011; Neubert et al., 2014) and aPFC–PPC effective connectivity relies on theta–band rhythm modulations (Phillips et al., 2014; Karalis et al., 2016). Here, we test whether social–emotional action tendencies are controlled through theta–band oscillations. We explore the dynamics of interactions between theta–band oscillations in aPFC and the beta/gamma rhythms produced by posterior areas during action selection.

We measured neural activity in 40 human participants using MEG while they performed a social approach–avoidance (AA) task. In this task, participants approach or avoid visually displayed emotional faces by pulling or pushing a joystick, respectively. Approaching angry and avoiding happy faces requires implementing rapid control to override automatic tendencies to approach appetitive and avoid aversive situations. This complex form of control operates on the interaction between the emotional valence of percepts and actions and it is modulated by social psychopathologies and socially relevant hormones (Heuer et al., 2007; Louise von Borries et al., 2012; Radke et al., 2013, 2017; Enter et al., 2016).

Previous fMRI studies have shown that, during affect-incongruent trials, aPFC and PPC activity is increased and amygdala activity decreased, suggesting that the aPFC exerts control over social–emotional actions, possibly by interacting with PPC to assist in action selection (Rowe et al., 2008; Volman et al., 2011a). Building on those fMRI and electrophysiological observations, we predicted that the emotional control evoked during incongruent trials will result in stronger theta–band power in aPFC. We also explore whether theta–band oscillations over aPFC are coupled to rhythms involved in action selection; that is, beta- and gamma-band rhythms over parietofrontal areas (de Lange et al., 2008; Jenkinson and Brown, 2011; Brinkman et al., 2014; Voytek et al., 2015).

Materials and Methods

Participants. Forty-five undergraduate students from Radboud University took part in the experiment. Five participants did not complete the experiment, with three failing to follow task instructions (>30% error trials) and two showing substantial noise in the MEG data. Forty participants were considered for analyses (mean age: 23.5; SD: 2.8; range: 18–33 years). All participants signed an informed consent before the study and were compensated with monetary reward or research credits. Before inclusion, participants were screened for epilepsy and history of mental illness. All participants were males, right handed, and had normal or corrected-to-normal vision. The study was approved by the local ethical committee (CMO: 2014/288).

Materials and apparatus. The task was programmed using Presentation software version 16.4 (www.neurobs.com; RRID:SCR_002521). Stimuli were presented using a PROPixx beamer with a refresh rate of 120 Hz and a resolution of 1920:1080. MEG was acquired using a whole-brain CTF-275 system with axial gradiometers. Data were sampled at 1200 Hz after a 300 Hz low-pass filter was applied. Four sensors (MLF62, MLC32, MLC11, and MRF66) were permanently disabled due to high noise. The helmet was set at 15° and the participants were seated 80 cm from the screen. Head location was measured using localization coils in both ear canals and on the nasion and was monitored continuously using online head localization software (Stolk et al., 2013). In case of large deviations from the initial head position (+5 mm), we paused the experiment and instructed the subject to move back to the original position. During the task, participants responded using a customized Fiber Optic Joystick (fORP design) calibrated for each participant before the experiment. A displacement of 20% away from the center in the sagittal plane was taken as a response.

Participants performed an AA task that has been used in several previous studies (Volman et al., 2011a,b) and was adapted for MEG (Fig. 1A). Trials started with the presentation of a white fixation cross presented in the center of a black screen for 1000 ms. After fixation, a face was presented for 100 ms, after which the subject had 2000 ms to respond by pushing the joystick away or pulling it toward themselves. Participants received written instructions on screen before each block of 12 trials, in which they were instructed to push the joystick toward or away from themselves for happy or angry faces, respectively (congruent block), or
toward/away from themselves for angry/happy faces (incongruent block). These response rules alternated each block. The task consisted of 16 blocks, yielding 196 trials in total. Stimuli consisted of equiluminant faces that were presented at the center of the screen at a visual angle of 4.3 × 6.4 degrees. In contrast to cognitive control tasks involving conflict on the stimulus level, such as emotional Stroop tasks, which typically implicate the anterior cingulate cortex, conflict between emotional percepts and emotional actions such as mapped by the AA task is typically processed by higher-order brain regions such as the aPFC (Volman et al., 2011a).

High-resolution anatomical MRI images were acquired with a single-shot MPRAGE sequence (acceleration factor 2 with GRAPPA method, TR 2400 ms/TE2.13 ms, effective voxel size 1 × 1 × 1 mm, 176 sagittal slices, distance factor 50%, flip angle 8 degrees, orientation A ⇒ P, FoV 256 mm). To align structural MRI to MEG, we provided the participants with vitamin E capsules in the ears on the same locations as the localizer coils in the MEG system.

Procedure. Upon arrival, participants received verbal task instructions before changing into nonmagnetic clothing. Before the experiment, participants provided a saliva sample, enabling the quantification of hormone levels for other research purposes. Once in the scanner, participants performed a practice session containing four blocks of eight trials each using faces of different identity from those displayed in the main experiment. After the main session of the AA task, participants provided a second saliva sample and completed the State Trait Anxiety Inventory (Spilberger et al., 1970). Structural MRI was acquired in a separate session in which participants also performed the AA task using fMRI. fMRI and hormonal data are not reported here.

Behavioral analysis. Reaction time analyses were performed on correct responses only. We removed trials in which the reaction time exceeded a threshold of 3 SDs above or below the mean reaction time of the subject for each condition separately (1.6% of trials). Congruency effects in reaction times and error rates were computed by subtracting congruent from incongruent trials.

MEG preprocessing. MEG analyses were performed using MATLAB2015a (The MathWorks; RRID:SCR_001622), Fieldtrip toolbox (Oostenveld et al., 2011), and custom-written analysis scripts. After epoching the data into trials ranging from −3 s before until 1 s after response, we removed the third order gradient. The trials were detrended and demeaned to remove slow drifts and nonzero DC offset and filtered using a discrete Fourier transform filter to remove the 50 Hz line noise and 100 and 150 Hz harmonics. Next, we performed manual trial rejection to remove trials with large deviations or artifacts. Independent component analysis (Makeig et al., 1996) was performed to remove components that contained sources of noise (e.g., heartbeat, eye blink, joystick artifacts). After this step, all trials were inspected visually to remove any trial still containing large amounts of noise. For the sensor-level analysis, we interpolated sensors that were missing due to noise removal using a weighted average of neighboring sensors; for some sensors (n = 8), this was not possible due to removal of too many neighbors. Sensor-level analyses were performed on 263 sensors.

Spectral analysis. To facilitate interpretation of the topographical distribution of signal resulting from the CTF axial gradiometers, we calculated spectral power for the horizontal and vertical components of the estimated planar gradient on each sensor location, which we then summed (Bastiaansen and Knösche, 2000). This representation of the data ensures that power of a source is strongest just above that source (Hämäläinen et al., 1993).

Time–frequency representations (TFR) of power were estimated in two steps. For frequencies <40 Hz, we used short-time Fourier transform with sliding windows of 500 ms, multiplied with a Hanning taper and moving in steps of 50 ms. The frequency resolution was 2 Hz. We presupposed 6 Hz as theta-band and 14–22 Hz as beta-band activity. Theta-band definition was based on a priori expectations (Phillips et al., 2014; Cavanagh and Frank, 2014; Cooper et al., 2015; Karalis et al., 2016). Beta-band definition was based on the congruency effect observed over all sensors. For frequencies >40 Hz (gamma-band, 40–130 Hz), we used three orthogonal Slepian tapers and sliding time windows of 200 ms (moving in steps of 50 ms), creating frequency smoothing of ~10 Hz (frequency resolution of 10 Hz; Percival and Walden, 1993). In the gamma-band analyses, we identified changes in power evoked before response (~1000 until 0 ms) compared with a baseline period occurring before stimulus presentation (average of ~800 until ~500 ms). This procedure increased statistical power by narrowing the frequency search space. The frequency band showing increased power (60–90 Hz) was used in further (orthogonal) comparisons between conditions. With the exception of single-trial analyses, all analyses were focused on the contrast between incongruent and congruent conditions.

Source analyses. Congruency effects were localized using DICS beamforming (Gross et al., 2001). We computed a single-shell head model (Nolte, 2003) for each subject using anatomical MRI. Next, we warped the individual MRI images to a template grid in MNI space (spatial resolution of 8 mm). To reconstruct activity in the interval 0.5 s before response delivery (period of action selection, see Results), we used a Hanning taper followed by a Fourier transform centered at 6 Hz and three orthogonal Slepian tapers for beta-band (center frequency 18 Hz) and gamma-band (center frequency 75 Hz). Slepian tapers result in more smoothing in the frequency domain and allowed us to reconstruct activity in a wider frequency range. A common spatial filter was created for each frequency based on all trials. This filter was consequently applied to congruent and incongruent trials separately. We computed relative change in power for incongruent versus congruent (incongruent − congruent)/(incongruent + congruent) to assess differences between conditions. Region labeling was done based on the Harvard–Oxford atlas implemented in FSL (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/; RRID:SCR_002823).

Spatial filters applied to ROIs. To assess whether gamma-band power is linked to theta-band phase, we constructed three spatial filters using linearly constrained minimum variance beamforming. Peak locations for the phase amplitude analyses were defined based on the peak grid-point resulting from the group level brain−behavior correlation analysis for the theta-band (aPFC; MNI: 40 48 −6) and the group-level gamma-band congruency effect [left postcentral gyrus (−28 −32 64) and right parietal superior lobe/postcentral gyrus (28 −42 70)]. These filters were applied to the MEG data, after which the dipole direction containing most variance was extracted.

Brain−behavior correlations. The source-reconstructed neural congruency effects in theta- and beta-band power were correlated with condition differences in reaction time (behavioral congruency effect). We calculated the correlation between the behavioral and the neural congruency effects for each grid point, after which we used Fisher’s z transform to convert Spearman’s r values into z values. Trial-by-trial estimates (<0.5 s until response) of aPFC theta-band power were obtained from aPFC peak grid point resulting from the group level brain−behavior correlation (centered at 40 48 −6). Trial-by-trial estimates of parietofrontal beta-band power were obtained from a precentral grid point (centered at 50 −10 36) resulting from the group-level brain−behavior correlation. For each subject, we set up a multiple regression model. First, we used trial-by-trial estimates of theta-band power (aPFC), beta-band power (precentral), and their interaction (thetaaPFC * betaprecentral) as predictors of the time series of reaction times separately for congruent and incongruent trials. The regressors were z-scored within each condition and regression parameters were estimated for each condition and participant separately using robust regression (MATLAB function robustfit). Standardized beta values for each subject were used in a second-level analysis testing whether trial-by-trial aPFC theta-power and precentral beta-power better accounted for reaction time variance during incongruent than during congruent trials (one-sided dependent-sample t tests in SPSS; RRID:SCR_002865) and whether those regression effects were different from zero (one-sample t tests). In this approach, single-trial estimates of power were regressed against single-trial reaction times separately for each condition, a procedure that is orthogonal to the group-level correlation between behavioral congruency and congruency in theta- and beta-band power.

Connectivity analysis. To assess whether aPFC influences downstream areas by influencing beta-band activity, we correlated condition differ-
ences in theta-band power extracted from aPFC (MNI: 40 48 — 6) with whole-brain condition differences in beta-band power. Spearman’s r values were transformed to z values using Fisher’s transform.

For theta-gamma coupling, we decomposed a spatial filter applied to the aPFC (MNI: 40 48 — 6) into 6 Hz complex signal using STFT with a 500 ms window tapered with a Hanning filter, moving in steps of 1 ms, after which the peak times between −0.5 s and response were extracted. Next, we time locked the remaining reconstructed time series (which were extracted from peak locations in the gamma-band condition difference) around each theta-band peak time point. From this (theta peak-locked) signal, we extracted gamma-band (40 — 130 Hz) power by performing STFT with a Hanning window containing six cycles per frequency band (df = 6/df) and moving in steps of 10 ms. We computed relative change in TFR between conditions for the phase-locked gamma-band power. We tested whether the gamma-band condition difference changed as a function of the phase of the theta-band extracted from aPFC, an approach that is orthogonal to testing for a difference between the experimental conditions. For interpretational purposes, we also computed the event-related field of the phase-locked signal from the aPFC to show the underlying theta-band waveform shape.

Statistical analyses. Statistical analyses on the sensor and source level consisted of cluster-based nonparametric permutation tests (Maris and Oostenveld, 2007). This procedure ensures correction for multiple comparisons over time of sensors (or grid points) and frequencies while also taking into account the dependency in the data by clustering neighboring points showing the same effect. As test statistics, we used t-values on the sensor-level data, z-values (fisher transformed r values) for brain–behavior correlations and theta-beta connectivity, and relative change (incongruent — congruent)/(congruent + incongruent) for the theta-gamma coupling. Unless stated otherwise, the reported p-values refer to cluster-corrected statistics.

Results
Behavioral costs of control over social–emotional action
Reaction times were longer for incongruent (M = 722 ms, SD = 176) compared with congruent trials [M = 680 ms, SD = 147; t(39) = −4.33, p < 0.001; Fig. 1B; effect size d = 0.26, calculated as (M1 — M2)/SDpooled]. In addition, we found higher accuracy levels for affect-congruent (M = 95.4%, SD = 4) versus incongruent trials (M = 92.3%, SD = 6.5) (t(39) = 3.07, p = 0.004, d = 0.57). These results illustrate that the task is effective in inducing behavioral costs when participants need to override their emotional action tendencies.

aPFC theta-band power increases during emotional action control
As detailed in the introduction, we hypothesized that frontal theta-band activity would be involved in control over social–emotional action tendencies. We found stronger theta-band power over anterior sensors before response during control of automatic emotional actions (i.e., incongruent vs congruent trials; center-frequency: 6 Hz, Fig. 2A, B). This congruency effect is statistically significant between 350 and 100 ms (maximum difference at 200 ms) before response initiation when controlling for multiple comparisons over sensors (N = 66; ROI on all anterior channels) and time points (p = 0.016; Fig. 2A, C). Reconstructing the source locations of this change showed increases of theta-band activity for incongruent trials with local maxima at right frontal pole (aPFC/superior frontal gyrus (20 40 50), frontal pole (26 70 8), and frontal orbital cortex (12 24 — 24) (Fig. 2D), with the strongest theta-band increases in the before response (20 40 50; Fig. 2E).

To localize theta-band activity that is relevant for behavior, we correlated condition differences in theta-band power — 0.5 s before until response in each grid point with reaction time effects over participants. ROI analysis on all frontal areas yielded a cluster of activity where the theta-band congruency-effect correlated positively with reaction time congruency effect (Fig. 2F, G, r(38) = 0.52, p = 0.0007 at peak grid point and r(38) = 0.44, p = 0.045 for the whole cluster, corrected for multiple comparisons over grid points). This cluster had two local maxima at MNI coordinates 40 48 — 6 and 28 30 40, which correspond to right frontal pole and middle frontal gyrus/superior frontal gyrus, respectively. This result indicates that those participants that show greater affect incongruency (as indexed by increased reaction times for incongruent vs congruent trials) also engage aPFC more strongly. Contra-lateral (left) frontal pole/inferior frontal gyrus showed similar correlations (maximum at 34 36 12), but this cluster remained below the cluster correction threshold. No clusters survived whole-brain thresholding.

Next, we implemented a post hoc exploratory analysis to determine whether the relation between increased theta-band power and emotional action control holds even on a trial-by-trial basis. Comparing single trial theta-band power derived from aPFC (MNI: 40 48 — 6) using one-sided t tests confirmed significant differences between conditions for theta-band power (t(39) = −1.77, p = 0.043). Theta-band power in the incongruent condition significantly correlated with behavioral performance in the same condition (t(39) = 2.69, p = 0.005). This was not the case for the congruent condition (t(39) = −0.29, p = 0.77). Together, these results confirm our hypothesis that theta-band power is a positive predictor of reaction time and show that aPFC is recruited more strongly with increased incongruence on a subject-by-subject as well as on a trial-by-trial basis.

Parietal and frontal beta-band activity decreases during emotional action control
Next, we tested whether an increase in theta-band power would be accompanied by parietal beta-band desynchronization. A large cluster of decreased alpha-/beta-band power in the incongruent versus the congruent condition resulted from the whole-brain sensor-level analysis (Fig. 3A, B), with significant condition differences between 600 ms before response until response onset (maximum difference at time 0, p = 0.0064; Fig. 3C, corrected for multiple comparisons over time points; sensors, N = 263) and frequencies. Differences were present over most sensors (Fig. 3B) and ranged from 8—26 Hz (Fig. 3A), with a peak between 12 and 18 Hz. Splitting the observed effect in alpha-band (8—12 Hz) and beta-band (14—26 Hz) (de Lange et al. (2008)) showed a significant decrease for beta-band activity (14—26 Hz, p = 0.0023; −600 until response), but not for alpha-band (8—12 Hz, p = 0.11). For source reconstruction of the beta-band activity, we used 18 Hz as center frequency with a frequency smoothing of 4 Hz (Fig. 3D). Time series reconstruction from right superior parietal lobule/supramarginal gyrus (44 — 40 56; Fig. 3D) illustrates that the beta-band desynchronization in this area peaks just before response and suggests a tight temporal relation with the frontal theta-band effect. These results, combined with the aforementioned theta-band results, indicate the involvement of aPFC theta-band and PPC beta-band activity during the control over social—emotional action tendencies, in close anatomical correspondence with previous studies on the control over emotional behavior (Volman et al., 2011a, 2013).
Correlating beta-band power and behavioral congruency differences revealed a large cluster ($r(38) = -0.68, p < 0.0001$ peak grid point and $r_{(38)} = -0.48, p < 0.0001$ whole cluster) over right precentral/postcentral gyrus with local maximum at 48 $-$ 2 36 (Fig. 3F). This correlation indicates that participants with a large behavioral congruency effect show larger suppression in beta-band power (Fig. 3G). Single trial beta-band power extracted from the peak location (48 $-$ 2 36) was significantly correlated with behavioral performance ($t_{(39)} = -3.06, p = 0.002$ for incongruent, but not for congruent trials; $t_{(39)} = -0.137, p = 0.45$) and also differed significantly between conditions ($t_{(39)} = 1.8, p = 0.04$). This indicates that beta-band desynchronization over parietofrontal regions is behaviorally relevant when control over social–emotional actions is implemented.

Effective connectivity between aPFC and parietofrontal areas during emotional action control
As a next step, we tested whether aPFC theta-band activity modulates rhythms involved in action selection. To assess connectivity between aPFC theta-band power and activity in anatomically downstream areas, we correlated the condition difference in aPFC theta-band power (MNI: 40 48 $-$ 6, local maximum of brain–behavior congruency effects (Fig. 2F,G) with the condition difference in beta-band power across each brain grid point on a subject-by-subject basis. Whole-brain analysis yielded a significant cluster ($r_{(38)} = -0.61, p < 0.0001$ peak grid point; $r_{(38)} = -0.46, p = 0.021$ whole cluster; corrected for multiple comparisons over grid points) over right precentral gyrus/postcentral...
gyrus with a local maximum at $50 - 10$ extending anteriorly into the middle frontal gyrus and posteriorly into the superior parietal lobule (Fig. 4A, B). Contralateral (left) precentral gyrus/postcentral gyrus showed similar connectivity below the whole-brain-corrected threshold. This effect indicates that participants with larger increases in aPFC theta-band power during control over social–emotional actions also show a larger decrease in beta-band power over parietofrontal areas. These findings suggest functional coupling between those two oscillatory phenomena in those two cortical regions. There were no significant connectivity effects after ROI analysis over the parietal cortex (Volman et al., 2011a), exploratory correlational analyses with the (theta-band) seed grid point at $28 - 30 - 40$; middle frontal gyrus/superior frontal gyrus; single-trial correlations between aPFC theta-band and beta-band activity from the connectivity peak location ($50 - 10$), or exploratory phase-amplitude coupling between aPFC theta-band phase and precentral beta-band power.

### Gamma-band activity increases during emotional action control

Given that the aPFC theta-band is thought to provide temporal organization to gamma-band activity (Lisman and Jensen, 2013; Voytek et al., 2015), we explored whether controlling emotional action tendencies resulted in changes in gamma-band power. To determine the frequency range of gamma-band activity evoked in these experimental settings, we first compared gamma-band power evoked before response with gamma-band power during baseline over all trials. This comparison showed an increase in

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**Figure 3.** Emotional control decreases beta-band power in parietal and frontal cortex. A, Time–frequency plot of between conditions power differences (congruency effect: incongruent - congruent/congruent + incongruent) averaged over sensors with a significant effect (see B). Time 0: response onset. The dashed box shows the time–frequency interval with a significant congruency effect ($-600$ to $0$ ms before response; $14 - 26$ Hz). B, Topographic distribution of sensors with a significant congruency effect at $14 - 26$ Hz (stars). C, changes over time in beta-band power ($14 - 26$ Hz) averaged across significant sensors (see B). The epoch with a significant difference between conditions is marked in gray. D, Cortical distribution of beta-band congruency effects (center frequency, $18$ Hz). E, Time series of $18$ Hz activity extracted from superior parietal lobule ($44 - 40 - 56$). F, Cortical distribution of correlations between beta-band ($18$ Hz) and behavioral congruency effects, with a significant cluster over right precentral gyrus (MNI coordinates of local maximum: $48 - 2 - 36$). G, Correlation between beta-band and behavioral congruency effects. Black dots represent measurements from each participant. Beta-band power changes are extracted from the right precentral maximum.
The increased theta-band power observed in this study could be an instance of low-frequency modulations of cortical ensembles (Jensen and Mazaheri, 2010); for example, the frequently observed theta-based coordination of mediofrontal neuronal ensembles during rule retrieval (Colgin, 2013; Harris and Gordon, 2015). However, the anatomical location and functional characteristics of the theta-band effect suggest a more specific mechanism: we show that this theta-band-related form of control emerged before response delivery from the aPFC rather than during feedback processing from medial frontal sources previously associated with inhibitory control and memory retrieval (Cavanagh et al., 2012; Colgin, 2013; Cavanagh and Frank, 2014). The aPFC, consistently shown to be involved in previous fMRI and transcranial magnetic stimulation studies of social–emotional action control (Kalisch, 2009; Volman et al., 2011a, 2013;
Morawetz et al., 2017), has been associated with the ability to control immediate action tendencies while implementing more abstract goals (Burgess et al., 2007; Badre and D’esposito, 2009; Koechlin, 2016; Mansouri et al., 2017), possibly by keeping online nonchosen response options (Boorman et al., 2009). Controlling emotional action tendencies, unlike emotional Stroop-like tasks involving stimulus-level conflict, requires considering the relative benefit of the unchosen behavioral strategy before a switch in response set is implemented (Boorman et al., 2009). The timing, anatomical location, and downstream effects of the current theta-band findings fit with the notion that control of emotional action tendencies involves maintenance of counterfactual choices in aPFC.

Parietalfrontal beta-band oscillations during control of social–emotional behavior

Emotional control evoked beta-band desynchronization localized to parietal cortex. Those features are not compatible with the beta-band synchronization elicited in inferior frontal gyrus during action inhibition (Swann et al., 2012; Aron et al., 2014; Bastin et al., 2014; Picazio et al., 2014). Here, beta-band desynchronization likely reflects increased engagement of areas involved in action selection (Brinkman et al., 2014) through release from tonic cortical inhibition (Khanna and Carmena, 2017) and increased corticospinal excitability (van Elskij, 2010). This release from inhibition is facilitated by decreases in GABA-ergic tone (Jensen et al., 2005; Yamawaki et al., 2008) and is accompanied by

Figure 5. Emotional control increases gamma-band power in parietal and frontal cortex during peaks of theta-band oscillations in aPFC. A, Time–frequency plot of between conditions power differences (congruency effect: incongruent — congruent/congruent + incongruent) averaged over sensors with a significant effect (see B). Time 0: response onset. The dashed box shows the time–frequency interval with a significant congruency effect (—350 to —50 ms before response; 60—90 Hz). B, Topographic distribution of sensors with a significant congruency effect at 60—90 Hz (stars). C, Changes over time in gamma-band power (60—90 Hz) averaged across significant sensors (see B). D, Cortical distribution of relative gamma-band congruency effects, with a significant cluster around the left central sulcus (—28 — 32 mm; 60—90 Hz activity extracted from left central sulcus (—28 — 32 mm). E, Time–frequency plot of gamma-band power congruency effects extracted from the local maximum in the left central sulcus (50—90 Hz) phase-locked to the aPFC theta-band signal before response. Contours are drawn around significant clusters where power is stronger in incongruent versus congruent trials. G, Event-related field of the theta-band signal extracted from aPFC.
increased gamma-band and spiking activity (Spinks et al., 2008; Donner et al., 2009). The parietofrontal reduction in beta-band power that we observed is inversely proportional to trial-by-trial slowing of responses during incongruent trials and significantly more so than during congruent trials, suggesting that this effect is not a trivial consequence of participants preparing a generic motor response, nor a systematic effect of task difficulty. In fact, the enhanced tonic beta-band desynchronization observed during incongruent trials might reflect stronger disinhibition of parietofrontal circuits when competition between multiple possible actions needs to be resolved and a larger neuronal search space needs to be considered (Cisek and Kalaska, 2010; Grent-’t-Jong et al., 2013; Brinkman et al., 2014).

Sensorimotor gamma-band oscillations during control over social–emotional behavior
Gamma-band synchronization in the motor system is linked to action preparation and movement selection (Donner et al., 2009; Schoffelen et al., 2011). When multiple response options are available during action preparation, such as during response conflict, increases in gamma-band oscillations are observed. This is often interpreted as simultaneous activation of multiple active response sets (Gaetz et al., 2013; Grent-’t-Jong et al., 2013), with an automatically triggered action competing with an alternative rule-based action. The increase in gamma-band power that we observed in the affect-incongruent condition might reflect enhanced coordination of local neuronal ensembles (Buzsáki and Wang, 2012) toward a state space suitable to initiate the correct, rule-based action (de Lange et al., 2008; Churchland et al., 2010; Kaufman et al., 2014) and away from the prepotent habitual action state. The phasic temporal relation between increases in central gamma-band power and peaks in aPFC theta-band oscillations indicates that selection of an alternative action could be implemented through interregional communication via phase-dependent modulations of gamma-band rhythms (Lisman and Jensen, 2013; Voytek et al., 2015).

Interpretational issues
It remains to be seen whether the relation between theta-band effects in aPFC and downstream beta- and gamma-band effects constitutes a functionally directional and monosynaptic interaction. It is known that aPFC sits at the top of the prefrontal hierarchy and projects to parietal and premotor areas (Ravnani and Owen, 2004; Miller and D’Esposito, 2005; Voytek et al., 2015; Koehler, 2016) and the timing of our effects suggests that aPFC might provide top-down regulation. However, given the lack of precise knowledge on the feedforward versus feedback connectivity of this circuit in humans (Neubert et al., 2014), the directionality of these effects remains elusive. The current findings do not exclude that other regions involved in emotional action selection could mediate interregional couplings between aPFC and parietofrontal cortex; for example, the pulvinar (Tyborowska et al., 2016) or the amygdala. The latter has been shown to be down-regulated by aPFC during social–emotional control (Volman et al., 2013) and is influenced directly by frontal theta-band oscillations during freezing (Karalis et al., 2016).

It could be argued that the theta-band effects that we report are generic byproducts of increased anxiety during incongruent trials. Previous studies have shown that theta-band activity in mPFC is involved in anxiety and fear behavior (Cavanagh and Shackman, 2015; Harris and Gordon, 2015), with increased theta-band connectivity between hippocampus and mPFC during fear-related inhibition of behavior (Adhkari et al., 2010; Khemka et al., 2017) and increased theta-band activity over mPFC in anxious individuals (Cavanagh and Shackman, 2015). However, the current changes in theta-band power were modulated by variations in performance on a trial-by-trial basis, an effect orthogonal to the systematic changes possibly related to state anxiety.

Source-level changes in aPFC theta-band and frontoparietal beta-band power arise in the right hemisphere. However, inspection of the data shows subthreshold effects in the corresponding contralateral regions, suggesting that the right hemispheric lateralization is a threshold effect of a bilateral process, which is consistent with previous fMRI reports (Volman et al., 2011b, 2013) and with the sensor-level scalp topographies. The anterior frontal location of the theta-band effect could be an artifact driven by task-related changes in head or eye position. This potential source of between-conditions differences is unlikely to account for the findings. There were identical stimuli and movements across conditions, head movements were monitored with millimeter precision during task performance (Stolk et al., 2013), and oculartifacts were aggressively removed with independent components analysis.

Phase–amplitude couplings can be inflated by nonsinusoidal oscillations (Lozano-Soldevilla et al., 2016; Cole et al., 2017), sharp transients in the data (Aru et al., 2015) or harmonics of lower frequencies (Jensen et al., 2016). In our case, theta-related harmonics and sharp transients are unlikely to play a role given that the theta-band and gamma-band signals originated from different cortical regions, with a clear sinusoidal theta-band signal in aPFC. Stronger theta-band power during incongruent trials might lead to more robust phase estimation, but this does not invalidate the presence of increased gamma-band power during peaks of the theta-band oscillation in aPFC.

Conclusion
This study defines neural responses to the problem of controlling human social–emotional behavior. Participants implement rapid changes in their predominant response set and select an alternative course of action by increasing theta-band power over aPFC, tonically decreasing beta-band power over parietofrontal cortex and transiently increasing gamma-band power over parietal and sensorimotor cortex through a mechanism phase locked to prefrontal theta oscillations. These findings provide clear mechanistic targets for interventional studies aimed at enhancing control over social–emotional behaviors in a number of psychopathologies.

References